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Local and macroscopic electrostatic interactions in single $\alpha$-helices

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Dedication: DNW would like to dedicate this paper to the memory of Prof. Dudley H. William FRS, an inspirational scientist, mentor and person.

The non-covalent forces that stabilise protein structures are not fully understood. One way to address this is to study equilibria between unfolded states and $\alpha$-helices in peptides. These, electrostatic forces are believed to contribute, including interactions between: side chains; the backbone and side chains; and side chains and the helix macrodipole. Here we probe these experimentally using designed peptides. We find that both terminal backbone-side chain and certain side chain-side chain interactions (i.e., local effects between proximal charges, or interatomic contacts) contribute much more to helix stability than side chain-helix macrodipole electrostatics, which are believed to operate at larger distances. This has implications for current descriptions of helix stability, understanding protein folding, and refinement of force fields for biomolecular modelling and simulations. In addition, it sheds light on the stability of rod-like structures formed by single $\alpha$-helices that are common in natural proteins including non-muscle myosins.

Folded proteins are stabilised by many weak intramolecular forces between thousands of atoms, including: hydrogen bonds, van der Waals' contacts, and Coulombic interactions between formal charges (i.e., salt bridges).¹ Because of the interdependence of non-covalent interactions, and the resulting cooperativity of protein folding, it is extremely difficult to disentangle contributions from these various components. For example, in terms of backbone interactions alone, CO$\rightarrow$NH$_{i+4}$ hydrogen bonds are regarded as the major factor stabilising $\alpha$-helices.² Whilst this remains unchallenged, recent studies have identified additional contributions from CO$\rightarrow$CO$_{i,n}$ n$\rightarrow$π* interactions,³ which can "compete" with traditional hydrogen bonding.⁴ Thus, it is important to develop continually, and to challenge our understanding of the fundamental forces underpinning protein folding, and of biomolecular interactions in water more generally. Improved understanding will lead to better modelling, prediction and design of biomolecular structures and assemblies ab initio, and with these many improved applications in the basic and applied biosciences will follow.

One way forward is to establish straightforward experimental models that isolate certain interactions, or reduce the complexity of their contexts. An example is the study of helix$\rightarrow$coil transitions in short peptides. This began with observations that certain peptide fragments of natural proteins exhibit considerable $\alpha$-helicity,⁵,⁶ and developed to the de novo design of model peptides that behave systematically with rational changes to sequence.⁷ The latter led to the dissection of various non-covalent interactions, and to sequence-to-structure relationships also apparent in fully folded natural proteins, including: $\alpha$-helical propensity scales;⁸-¹⁰ specific backbone-side chain interactions;¹¹ and pairwise and higher-order side-chain interactions.¹²-¹⁵

To balance the configurational entropy lost on folding into an $\alpha$-helix, a number of electrostatic forces probably contribute to favourable enthalpies of helix folding, including: backbone CO$\rightarrow$NH$_{i+4}$ hydrogen bonds; hydrogen bonds and salt bridges between side chains spaced i$\rightarrow$i+3 and i$\rightarrow$i+4;¹⁶ backbone-side chain interactions at both helical termini;¹⁷ and through-space electrostatic
interactions between charged side chains and the so-called helix macrodipole (Fig. 1). The latter is believed to arise from the alignment of individual dipoles of backbone peptide units parallel to the major axis of the α-helix. This has local consequences: it results in partial positive and negative charges at the N- and C-termini, respectively, which leads to side-chain preferences and ligand-binding phenomena at these sites. However, the evidence and roles for the macroscopic effects are less clear, and the energetic contributions to structure and stability by both surface-exposed salt bridges and the α-helix macrodipole remain unresolved.

Most recently, the field has come full circle with observations of natural long single α-helices that span 30 – 200 residues. These self-stabilising structures do not require tertiary or quaternary interactions to persist. Originally noted in the calmodulin-binding protein caldesmon and the non-muscle myosins VI, VII, and X, these single α-helices (SAHs) are predicted to occur widely. Their hallmark is alternating blocks of four negatively charged glutamic acid (Glu, E) and four positively charged arginine (Arg, R) or lysine (Lys, K) residues. They are also dubbed charged single α-helix (CSAH) domains, or E-R/K α-helical motifs and are proposed to form rigid rods or spacers to separate other protein domains.

Given this revived interest in single α-helices, but the incomplete understanding of the forces that stabilise them, we designed a series of peptides comprising predominantly Glu and Lys to prise apart different contributions to helix stability. Our first designs were consistent with preceding reports. However, further iterations conflicted with current thinking for the contributions from both salt-bridge and macrodipole interactions. Specifically, certain side-chain arrangements between Glu and Lys are favoured over others, and these dominate any manifestation of the macroscopic helix dipole; indeed, for these systems, there is no need to invoke macroscopic effects and local descriptions suffice.

RESULTS
Peptide design – iteration 1
We followed foregoing studies of de novo and natural SAHs to design two highly helical peptides, (E4K4)3 and (K4E4)3 (Table 1). In these, alternating blocks of four Glu and four Lys, which both have favourable α-helical propensities, were used to match the 3.6-residues per turn of the α-helix. This maximises potentially favourable i→i+4 E→K and K→E inter-side-chain salt bridges (Fig. 1), whilst minimising unfavourable inter-side-chain electrostatic interactions (Supplementary Results, Supplementary Table 1). N.B., in these two designs, both i→i+4 E→K and K→E interactions are possible, but one potentially outnumbers the other by 3:2 in each case. In addition, all four types of side-chain-terminus interaction are represented, e.g. blocks of four Glu or Lys at both the N- and C-termini of the peptides (Fig. 1). We used (E12K12)6 as a control peptide of the same length, which maximises repulsive i→i+4 E→E and K→K pairs (Supplementary Table 1).

Circular dichroism (CD) spectroscopy showed that (E4K4)3 was largely unfolded; and that both (E4K4)3 and (K4E4)3 had high mean residue ellipticities at 222 nm (MRE), which translated to 74% and 62% helix, respectively (Fig. 2a-c, Supplementary Fig. 28a&b and Table 1). (E4K4)3 was expected to be more helical than (K4E4)3 because it better matches the formal charges of the side chains and the partial charges of backbone in the N- and C-terminal turns (Fig. 1); and, the E→Ki→i arrangement counteracts the helix macrodipole better. This continued over a range of peptide lengths for (E4K4)n and (K4E4)n, where n = 1 – 4, with the former being the more helical in each case (Fig. 2c).

As secondary structure can be stabilised by peptide association, we used sedimentation equilibrium experiments to determine peptide oligomeric states. These indicated that both (E4K4)3 and (K4E4)3 were monomeric in solution (Supplementary Fig. 30 and Supplementary Table 2).

Next, we used high-resolution 1H-NMR NOESY and TOCSY experiments to probe the structure in (E4K4)3 and (K4E4)3 in detail (Fig. 3 and Supplementary Figs. 31-34). Sequence-specific assignments were made for all backbone HN resonances, and the majority of the Hα resonances
for both peptides. From $\Delta \delta$ and NOE measurements, $(E_4K_4)_3$ was helical throughout with contiguous, sequential HN-HN$_{i+1}$ NOEs from Gly1 through Trp27 (Fig. 3g). Whereas, for $(K_4E_4)_3$ the N-terminal residues Lys2 – Lys4 appeared disordered (Fig. 3h); otherwise, the helical structure continued through the sequence from Lys5 to Trp27. This fraying in $(K_4E_4)_3$ seems best explained by the mismatched charge of the side chains of Lys2, 3 and 4 with the $\delta^+$ N-terminus; although the C-terminus is folded despite the negatively charged Glu residues at this $\delta^-$ end.

**Peptide design – iteration 2**

To reduce complications from end effects—i.e., local electrostatic interactions between charged side chains and the partially charged termini—we synthesised peptides in which $(E_4K_4)_3$ and $(K_4E_4)_3$ were flanked at one or both sides by tetra-alanine (Ala$_4$, Ala$_4$) blocks (Table 1). Alanine was chosen for its high helix propensity,$^{10}$ small size and charge neutrality. Although we anticipated some influence of these blocks on overall helicity of the peptides, we reasoned that these properties would best serve the purpose of homogenising end effects. The resulting six peptides, $A_4(E_4K_4)_3$, $(E_4K_4)_3A_4$, $A_4(E_4K_4)_3A_4$, $A_4(K_4E_4)_3$, $(K_4E_4)_3A_4$ and $A_4(K_4E_4)_3A_4$, showed the following responses to the flanks in CD spectroscopy (Fig. 2, and Supplementary Figs. 28c – f and 29c – f).

First, the percent helix in all of the peptides increased compared with the parents (Fig. 2d). This is almost certainly due in part to the increased lengths of the peptides and the choice of Ala as the flanking residues. It is also fully consistent with the Ala$_4$ flanks limiting the terminal fraying/unfolding of the peptides.$^{36-38}$ Second, changes in helicity were more pronounced for the $(K_4E_4)_3$-based peptides. This seems best explained simply by the removal of fraying due to the block of unfavourable positive charge from the helical N-termini. Consistent with this, within the $(K_4E_4)_3$ series the favourable impact of the flanks was largest at the N-terminus.

However, a final response was unexpected, and is less readily explained by current thinking. Comparing the overall helicities of the parent and doubly flanked peptides $(E_4K_4)_3 > (K_4E_4)_3$, but $A_4(K_4E_4)_3A_4 > A_4(E_4K_4)_3A_4$ (Table 1 and Fig. 2). This runs contrary to most reports of side chain-side chain interactions between Glu and Lys in the context of the helix macrodipole. Current understanding is that $E_i \rightarrow K_{i+4}$ should be more stabilising than the reverse combination because the side chains align energetically favourably with the helix macrodipole.$^{7,39}$ However, we find the opposite: $A_4(K_4E_4)_3A_4$, which has more potential $K_i \rightarrow E_{i+4}$ pairings than $E_i \rightarrow K_{i+4}$, is the more helical.

Further NMR measurements on $A_4(K_4E_4)_3A_4$ confirmed contiguous sequential HN-to-HN$_{i+1}$ NOEs from Ala3 through Trp35 (Supplementary Fig. 34). Thus, the entire central $(K_4E_4)_3$ block is $\alpha$-helical. Moreover, length-dependence experiments for the flanked peptides revealed that the $(K_4E_4)_3$ blocks were consistently more helical than $(E_4K_4)_3$ blocks (Fig. 2c). N.B., sedimentation-equilibrium studies confirmed that both $A_4(E_4K_4)_3A_4$ and $A_4(K_4E_4)_3A_4$ were monomeric in solution (Supplementary Fig. 30 and Supplementary Table 2).

$(K_4E_4)_3$ salt bridges are enthalpically better

The additional stabilising effect of $K_i \rightarrow E_{i+4}$ pairings compared with $E_i \rightarrow K_{i+4}$ was also evident in thermal unfolding followed by CD spectroscopy. The $E_4K_4$-based peptides showed broad and approximately linear transitions from predominantly helical to gradually less helical states, including the most folded $A_4(E_4K_4)_3A_4$ (Fig. 2b and Supplementary Figs. 29c&d). This is typical of short de novo $\alpha$-helical peptides.$^7$ By contrast, the $K_4E_4$-based peptides had sharper more-sigmoidal transitions (Fig. 2b and Supplementary Figs. 29d&f), and some had measurable midpoints ($T_m$) with $A_4(K_4E_4)_3A_4$ having the highest. Though there are examples of such behaviour,$^{40}$ it is unusual and suggests cooperativity in the transitions, and that more enthalpically favourable interactions are being made in the folded state of this peptide.

We fitted the thermal unfolding data to the Gibbs-Helmholtz equation (Fig. 2b and Supplementary Fig. 29). This assumed $\Delta C_p = 0$, and that helix-coil transitions are two-state processes. The latter is almost certainly incorrect as the "folded states" are likely ensembles of unfolded, folded and partly folded species; however, we found no evidence for alternative "unfolded" states by CD spectroscopy at least, i.e. no polyproline helices or $\beta$-structure. Caveats aside, the analysis is valuable because the gradients of the fits reflect the enthalpy change accompanying unfolding. For
our designed peptides, the enthalpies of folding were all exothermic (Supplementary Table 7). Moreover, the enthalpies for the \((K_4E_4)i\)-based peptides were higher than for the \((E_4K_4)i\), analogues. This suggests that \(K\rightarrow E_{i+4}\) salt bridges are somehow enthalpically better than \(E_i\rightarrow K_{i+4}\) interactions, and cooperate better to induce and maintain structure in \(A_4(K_4E_4)iA_4\).

**Helical propensities and salt-bridges in folded proteins**

There have been extensive analyses of residue preferences and pairings in \(\alpha\)-helices\(^{12,41,42}\) but our results raised issues to be addressed afresh. Therefore, we analysed the conformations and interactions of Glu and Lys side chains in \(\alpha\)-helices from the PDB (Fig. 4).\(^{43}\)

First, we tested for specific side-chain-backbone interactions at the helical termini. Consistent with previous studies\(^9,36-38\) and our experiments, this analysis gave favourable propensities for Glu at helical N-termini and Lys at the C-termini of 1.24 and 1.26, respectively (Supplementary Table 8). However, using HBPLUS,\(^{44}\) we found that only small fractions (9% for Glu at N-termini, and 2.5% for Lys at C-termini) of these made side-chain-backbone hydrogen bonds (Supplementary Fig. 36). In the context of the many potential hydrogen bonds with water, this suggests strongly that the experimental and statistical preferences for Glu at N-termini are best attributed to proximal through-space electrostatic interactions between the \(\gamma\)-carboxylate and the \(\delta^+\) ends of nearby peptide-bond dipoles, rather than by specific atomic contacts; and the preferences for Lys at C-termini by similar interactions between the \(\varepsilon\)-amino moieties and the \(\delta^-\) charge.

Regarding potential \(E_i\rightarrow K_{i+4}\) and \(K_i\rightarrow E_{i+4}\) salt bridges, we found that more \(K_i\rightarrow E_{i+4}\) pairs (21%) formed salt bridges than \(E_i\rightarrow K_{i+4}\) (11%); and that the salt-bridged \(K_i\rightarrow E_{i+4}\) pairs were more structurally conserved, with a lower RMSD value of 1.08 \(\AA\), vs. 1.79 \(\AA\) for those made by \(E_i\rightarrow K_{i+4}\); calculated over all side-chain atoms, (Supplementary Fig. 37). To understand this, we compared the \(\chi_1,\chi_2\) distributions for Glu-Lys salt bridges (Fig. 4) and all possible \(\chi_1,\chi_2\) combinations for Glu and Lys in \(i\rightarrow i+3\) and \(i\rightarrow i+4\) pairings in \(\alpha\)-helices (Supplementary Table 9).

Glu has three dominant \(\chi_1,\chi_2\) combinations: gauche,trans \((gt, 36\%);\) trans,trans \((tt, 25\%);\) and gauche.gauche \((gg, 19\%);\) (Figs. 4a&d). However, when paired with Lys only certain combinations of these led to salt bridges: For \(E_i\rightarrow K_{i+4}\) pairs, salt bridges were only made with Glu in the second-favoured conformer \((tt)\) or a fourth minor conformation \((fg^*, 10\%);\) (Fig. 4a,e&f and Supplementary Table 9). Having to fix the side chain in these less-favoured rotamers incurs an entropic penalty. Nevertheless, the more-flexible side chain of Lys complemented these Glu conformers to account for 92% (23/25) of the \(E_i\rightarrow K_{i+4}\) salt-bridge interactions (Figs. 4b,e&f). By contrast, 75% of the salt-bridged \(K_i\rightarrow E_{i+4}\) pairs had the same paired conformation, with Glu in its preferred \(\chi_1,\chi_2\) conformation and Lys in its second-favoured conformer \((tt, 32\%);\) which was only marginally less preferred than \(gt\) \((39\%);\) (Figs. 4c,d,i). Hence, the entropic penalty of freezing out this conformer is near the minimum possible.

Thus, compared with \(E_i\rightarrow K_{i+4}\) salt bridges, the preferred \(K_i\rightarrow E_{i+4}\) salt bridges appear to sample fewer conformations and use more-favourable \(\chi_1,\chi_2\) pairs for each residue. We posit that this leads to \(K_i\rightarrow E_{i+4}\) interactions that are enthalpically better, because the side chains are predisposed in conformations to facilitate salt bridges. This is not to say that sequences dominated by \(E_i\rightarrow K_{i+4}\) arrangements will necessarily be less helical; from our data and the literature this is patently not the case. However, as discussed below, the mechanism of helix stability is subtly different.

**Uncoupling \((E_iK_4)\) and \((K_4E_4)\) blocks**

The peptide designs described above can make both \(E_i\rightarrow K_{i+4}\) and \(K_i\rightarrow E_{i+4}\) interactions. Thus, it is not possible to ascribe absolutely either the percent-helix or thermal unfolding properties to one \(i\rightarrow i+4\) interaction type. To address this, we designed further sequences with isolated \(E_iK_4\) or \(K_4E_4\) blocks flanked by \(A_4\) blocks; namely, \(A_4(E_iK_4)iA_4,\) \(A_4(K_iE_4)iA_4,\) \(A_4(E_iK_4)iA_4E_4K_4A_4\) and \(A_4(K_iE_4)iA_4E_4K_4A_4\) (Table 1). In these, only one type of \(i\rightarrow i+4\) \(E\rightarrow K\) or \(K\rightarrow E\) interaction can be made. The shorter peptides were surprisingly helical (Fig. 2c, Table 1, and Supplementary Fig. 28c&d), with \(A_4(K_iE_4)iA_4\) having the higher \(\alpha\)-helicity. Moreover, the thermal-unfolding curve for this peptide was sharper and more sigmoidal than for \(A_4(E_iK_4)iA_4\) (Supplementary Fig. 29c&d). The
longer peptides with the additional, but uncoupled, E$_4$K$_4$ or K$_4$E$_4$ blocks were highly helical at 83% and 97%, respectively. (N.B., some of the structure is likely due to the central Ala$_3$ block). Both peptides had sigmoidal thermal-denaturation curves with measurable midpoints, but the K$\rightarrow$E arrangement had the sharper transition and the higher $T_M$ (Table 1 and Supplementary Fig. 29g). This confirms that K$\rightarrow$E$_{i+4}$ potential salt bridges contribute more to helical stability than E$\rightarrow$K$_{i+4}$ pairs, despite the apparent misalignment with the helix macrodipole.

Consideration of E/K$_r$ to K/E$_{i+3}$ interactions

Our designs could also make i$\rightarrow$i+3 interactions (Supplementary Table 1). Therefore, we examined such pairs in $\alpha$-helices of the PDB. Interestingly, more E$\rightarrow$K$_{i+3}$ pairs (23.4%) made salt bridges than the K$\rightarrow$E$_{i+3}$ pairs (15.6%) (Supplementary Table 8b). Also, for the E$\rightarrow$K$_{i+3}$ pairs there was one dominant rotamer combination that accounted for 77% of E$\rightarrow$K$_{i+3}$ salt bridges made; whilst, for K$\rightarrow$E$_{i+3}$ pairs multiple conformers made salt bridges, with two accounting for 60% of the interactions (Supplementary Fig. 38). In addition, we found that E$\rightarrow$K$_{i+3}$ salt bridges were more structurally conserved, with lower a RMSD (1.09 Å) than for K$\rightarrow$E$_{i+3}$ salt bridges (2.27 Å) (Supplementary Fig. 37). These trends are the opposite of those observed for the i$\rightarrow$i+4 pairs, which suggest that E$\rightarrow$K$_{i+3}$ salt bridges might be more helix stabilising than K$\rightarrow$E$_{i+3}$.

To test this, we synthesised A$_4$(E$_3$K$_3$)$_3$A$_4$ and A$_4$(K$_3$E$_3$)$_3$A$_4$. Both peptides were highly helical and showed clear thermal unfolding transitions (Table 1 and Supplementary Figs. 39b&d). However, they were less helical than the E$_4$K$_4$ and K$_4$E$_4$ analogues, A$_4$(E$_4$K$_4$)$_3$A$_4$ and A$_4$(K$_4$E$_4$)$_3$A$_4$, illustrating that i$\rightarrow$i+3 interactions are less favourable than i$\rightarrow$i+4. In addition, and as predicted above, the trend was switched compared with E$_4$K$_4$ and K$_4$E$_4$, i.e., A$_4$(E$_3$K$_3$)$_3$A$_4$ was more helical and had a higher $T_M$ than A$_4$(K$_3$E$_3$)$_3$A$_4$ (Table 1). Nonetheless, and perhaps counter-intuitively, the folding of A$_4$(K$_3$E$_3$)$_3$A$_4$ was the more exothermic (Supplementary Table 7). We suggest that this is because A$_4$(K$_3$E$_3$)$_3$A$_4$ can make more of the higher-enthalpy K$\rightarrow$E i$\rightarrow$i+4 interactions (Supplementary Table 1).

The role of the helix macrodipole

Whilst K$\rightarrow$E$_{i+4}$ salt bridges make enthalpically better interactions and lead to more-folded $\alpha$-helical peptides than E$\rightarrow$K$_{i+4}$, this arrangement of positively charged lysine blocks N-terminal to negatively charged glutamate blocks is predicted to be unfavourable with respect to the helix macrodipole. Therefore, we calculated the electrostatic potential ($V_E$) of a model $\alpha$-helix in vacuum, and at varying distances (x) from the central helix axis (z) by approximating partial dipole charges as point charges (Fig. 5a and Supplementary Fig. 40). In this model, the electrostatic potential for an individual peptide dipole was $\approx \pm 0.7$ V (Fig. 5b). For $x = 0$ in an extended helix, i.e. directly along its axis, we found that the central peptide-bond dipoles cancelled, leaving a negligible macrodipole at the termini ($V_E \approx \pm 1.5$ V, Fig. 5c). At $x = 2.5$ Å, i.e. where the majority of backbone atoms lie, the residual macrodipole is overshadowed by local electrostatics from these atoms, with $V_E \approx \pm 10$ V (Fig. 5d). At $x = 5$ Å, i.e. approximately at the termini of side chains, the effect is similar to $x = 0$, but with the macrodipole diminished further ($V_E \approx \pm 0.6$ V, Fig. 5e). Indeed, at this distance the macrodipole is approximated by partial point charges at the termini of the helix (Fig. 5f). Thus, and even in the absence of solvation effects, the influence of the macrodipole on the stability of central K$\rightarrow$E interactions is predicted to be minimal.

Correlation with helix-prediction programs

Considerable effort has gone into capturing the data from helix$\leftrightarrow$coil transitions in computational methods to predict helix content from peptide sequence, with AGADIR probably being the most popular.45 We made AGADIR predictions for both the percentage helix and thermal unfolding behaviours of our designs, and compared these with our experimental data (Supplementary Figs. 41&42). At first sight, the predictions appeared good. However, there were some deviations from the observed values, particularly for the longer and Ala$_3$-flanked variants. Moreover, the thermal unfolding data were not modelled well by AGADIR, which tended to over predict stabilities. This paper provides new data that could be modelled to improve these predictions and to aid peptide designs.
Biological context
Our observations that certain side-chain arrangements are more stabilising than others, i.e. $K \rightarrow E_{i+4}$ > $E \rightarrow K_{i+4}$ and $E \rightarrow K_{i+3}$ > $K \rightarrow E_{i+3}$, may help understand the behaviour of SAHs. To examine this in the natural sequences, we counted all of the potential $D/E \rightarrow K/R$ and $K/R \rightarrow D/E$ pairs spaced $i \rightarrow i+3/4$ in sequences from non-homologous proteins assigned as SAHs from CD spectroscopy and/or electron microscopy; we then normalised the data with expected rates (Supplementary Fig. 43). Firstly, as anticipated, we found that aspartic acid (D) hardly featured in the sequences or pairings. Secondly, the normalised frequencies of oppositely charged pairs were much greater than those for like-charge pairs, by an average factor of 7 for $i \rightarrow i+4$, and 3 for $i \rightarrow i+3$. Third, except for the $E \rightarrow R$ combinations, which had similar observed/expected ratios, the propensities of oppositely charged $i \rightarrow i+4$ pairs were greater than those for the corresponding $i \rightarrow i+3$ pairs. Fourth, and consistent with our model-peptide studies, there were more $R/K \rightarrow E_{i+4}$ pairs than the reverse orientation. Fifth, the data for $i \rightarrow i+3$ pairings in natural SAHs were less clear-cut, although the pairs with overall highest propensity were $E \rightarrow R_{i+3}$, and $K \rightarrow E_{i+3}$. Finally, there was a consistent slight preference for Arg-based pairings over those involving Lys (apart from $K \rightarrow E_{i+3} > R \rightarrow E_{i+3}$), which may reflect arginine’s potential to participate simultaneously in two salt bridges.15

DISCUSSION
We have found that: (1) short peptides of ≈20 – 30 residues comprising alternating blocks of Glu and Lys, formed highly $\alpha$-helical structures; (2) flanking these with Ala blocks increased helical content and revealed context-dependent phenomena; for example, (3) Glu residues in the $N$-terminal turn strongly promoted $\alpha$-helix formation, and Lys in these positions disrupted it; whereas, (4) similar effects at the $C$-terminus were less pronounced; (5) central $K \rightarrow E_{i+4}$ arrangements made enthalpically better side-chain salt-bridge interactions than $E \rightarrow K_{i+4}$ pairs; as a result, (6) although central $E \rightarrow K_{i+4}$ and $K \rightarrow E_{i+4}$ arrangements both favoured helix formation, the resulting helical structures were different in nature, particularly in how they responded to temperature; and (7) $i \rightarrow i+3$ potential salt-bridge interactions were weaker than those made by $i \rightarrow i+4$ arrangements. We place these results in context as follows.

1. Local interactions between side-chain and backbone moieties at helical termini. It is well established, in peptides9,36-38 and proteins,11 that certain side chains interact favourably with polar backbone groups and/or partial charges at the $N$- and $C$-termini of $\alpha$-helices.25 Our first peptide designs confirmed this: peptides with $N$-terminal blocks of lysine were less folded and more frayed than those with blocks of glutamate at that end. Curiously, however, in protein structures of the PDB, we found very few examples of formal hydrogen-bonded or other atom-atom contacts between side chains and backbone atoms at the termini. This indicates that such interactions are ephemeral, if there at all, and that more-general electrostatic interactions are at play between side chains and the partial charge at the $N$-terminus. Moreover, for our de novo peptides, these effects were much more pronounced at the $N$-terminus than at the $C$-terminus. This is readily understood as side chains emerge from the helical backbone and point towards the $N$-terminus (Fig. 1).

2. Local interactions between side-chain groups in the central regions of helices. To remove the end effects, we flanked the Glu/Lys-based peptide sequences with tetra-alanine units. This led to a surprising discovery: $i \rightarrow i+4$ $K \rightarrow E$ pairs stabilised $\alpha$-helical structure more than the reversed $E \rightarrow K$ orientation. This result was consistent across all of our peptides, and was supported by the analysis of the PDB. The latter revealed that $K \rightarrow E_{i+4}$ pairs make more formal salt-bridges than $E \rightarrow K_{i+4}$ pairs. Moreover, the mechanisms through which such salt bridges are formed appears different in the two cases: whereas, those in $K \rightarrow E_{i+4}$ pairs employ a small number of sterically favoured side-chain conformations, salt-bridges made by $E \rightarrow K_{i+4}$ arrangements use many more and less-favoured conformations. We argue, that this leads to enthalpically better $K \rightarrow E_{i+4}$ salt bridges, and entropically favoured $E \rightarrow K_{i+4}$ interactions. This was manifest in the thermal unfolding transitions of different peptides: those with more $K \rightarrow E_{i+4}$ arrangements had sharper, and in some cases sigmoidal, thermal unfolding curves; whereas, peptides dominated by $E \rightarrow K_{i+4}$ pairs showed broader, less-defined transitions.
These findings are noteworthy as there is disagreement in the literature over which pairing is the more stabilising, with all possible stances being adopted: $E \rightarrow K_{i+4} > K_{i} \rightarrow E_{i+4}$, \textit{vice versa}.\textsuperscript{16} We believe that our combined experimental and bioinformatics studies help resolve this.

3. \textbf{Non-local interactions between side chains and the macroscopic helix dipole.} Our finding that $K_{i} \rightarrow E_{i+4}$ pairs are more stabilising than $E_{i} \rightarrow K_{i+4}$ arrangements has a further implication. This result is contrary to that expected from consideration of the macroscopic helix dipole alone.\textsuperscript{7} Organising $K_{i} \rightarrow E_{i+4}$ (+ve $\rightarrow$ –ve) pairs in an $\alpha$-helical conformation should require work against the helix macrodipole, which runs from a partially positively charged $N$-terminus to a partially negatively charged $C$-terminus; \textit{i.e.}, it will cost free energy and disfavour the helical state. In contrast, folding of $E_{i} \rightarrow K_{i+4}$ pairs should counter the dipole and therefore be favourable. That we do not see this effect—indeed, we observe the opposite—requires explaining.

The most straightforward explanation is that the macrodipole does not operate and is attenuated out where the charged moieties of the side chains reside. That is, even in the absence of solvation and dielectric effects, the charged moieties of side chains such as glutamic acid and lysine are not expected to make significant Coulombic interactions with the macrodipole. That said, they are expected to interact with the terminal partial charges arising from the half-dipoles organised at the $N$- and $C$-termini. Thus the local, terminal effects of the helix dipole are not in doubt; indeed, these are well established, and our results and conclusion in point 1 above support this. What we do question, however, is the need to invoke helix-dipole arguments beyond this, \textit{i.e.}, any macrodipole effect of the $\alpha$-helix; we argue that the local description of the helix dipole, and how this affects side-chain preferences, binding of co-factors in proteins and so on, should be sufficient for explanations of experimental data and theoretical treatments.

In addition to contributing to fundamental understanding of helix stability, we envisage that our systematic set of designed peptides, the inventory of potential interactions within these, and the experimental data that we provide on helix content and thermal stability, will provide a useful resource to computational chemists and biochemists developing empirical and theoretical methods for analysing and predicting helix-forming peptides and protein fragments, and force fields for protein modelling more generally. Finally, in the context of synthetic biology, the fully folded peptides that we present may find use as linkers and spacers of tuneable length and thermal unfolding behaviour to combine protein domains in defined ways.\textsuperscript{46,47}

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\textbf{Author contributions}
EGB, CFJF and DNW designed the research; EGB made the synthetic peptides and performed the CD spectroscopy; MPC and EGB collected and analysed the NMR data; GJB, EGB and DNW performed the bioinformatics; EGB and RBS constructed and analysed atomistic models for the peptides; NL performed and analysed the helix-dipole calculations; EGB, GJB and DNW wrote the paper. All authors reviewed and contributed to the manuscript.

The authors declare no competing financial interests.
Figure 1: Electrostatic interactions in the α-helix. Cartoon representation of the α-helix showing the following: CO→NH-4 hydrogen bonds (green dashes); the peptide-bond dipole (thick sticks, centre), many copies of which align to give the helix macrodipole; clusters of backbone NH and CO groups, which give rise to local δ+ and δ− charges at the N- (blue) and C-termini (red), respectively; interactions between charged side chains and the helix terminal charges (arrows); and salt bridges between charged side chains spaced i→i+3 and i→i+4 apart (labelled spheres).

Figure 2: Helicities of the designed peptides in solution. (a) CD spectra at 5 °C of: (EiKzl)6 (black crosses), (EiKz)3 (red circles), (KiEz)3 (blue squares), A4(EiKzl)3A4 (red open circles), and A4(KiEz)3A4 (blue open squares). (b) Thermal unfolding curves followed by the change in MRE in for all except (EiKzl)6, with the symbols and colours from (a) preserved. Fits to the Gibbs-Helmholtz equation are shown by solid lines. (c) Fraction helix as a function of peptide length for the (EiKzl)n (red) and (KiEz)n (blue) peptides with (open symbols) and without (filled symbols) tetra-alanine flanks (data points are joined by lines to guide the eye). (d) Increase in fraction helicities at 5 °C of the alanine-flanked peptides A4(EiKzl)3, (EiKzl)4A4 and A4(KiEz)4A4 (red colours) relative to (EiKzl)4, and for A4(KiEzl)3, (KiEz)4A4 and A4(KiEz)4A4 (blue colours) relative to (KiEzl)4. Conditions: 100 µM peptide concentration in PBS 137 mM NaCl, pH 7.4.

Figure 3: Locating α-helical structure by NMR spectroscopy. 1H-NMR spectra for (EiKzl)3 (a&c) and (KiEzl)3 (b&d). (a&b) Overlaid fingerprint regions of the TOCSY (coloured contours): red for (EiKzl)3 and blue for (KiEzl)3 and NOESY (black contours) spectra. Note the larger range of HN shifts for (EiKzl)3. (b&d) NOESY spectra showing the amide regions and large number of sequential HN-HN, NOEs. Note the first NOE of this type for (KiEzl)3 is between Lys-5 and Glu-6 (KS/E6 circled in d). (e&f) δδ plots for the backbone Hν chemical shifts of (EiKzl)3 and (KiEzl)3, respectively. Key: Glu, red; Lys, blue; Gly and Trp, grey. δδ values are the difference in sequence-correlated random coil chemical shifts from those observed. δδ values are greater for (EiKzl)3 than (KiEzl)3 on average, consistent with a more folded α-helix. (g&h) Connectivity diagrams for (EiKzl)3 and (KiEzl)3, respectively. Note there are only a few identifiable Hδ-HN NOEs due to significant peak overlap in this region (a&b). Key: thick bars, medium intensity NOEs (1.8 – 3.5 Å); thin bars, weak NOEs (1.8 – 5.0 Å); grey, ambiguous; d, NOE hidden by diagonal; D, degenerate; O, overlapped. Conditions: 1 mM peptide concentration in PBS (137 mM NaCl) with 10% D2O, pH 7.4 at 5 °C; (EiKzl)3 800 MHz, and (KiEzl)3 900 MHz.

Figure 4: Side-chain interactions observed in α-helices from the Protein Data Bank. χ1, χ2 distributions and conformers for Ei→Kzl and K→Ezl pairs in high-resolution X-ray crystal structures. (a – d) Normalised frequencies of preferred χ1, χ2 angles for Glu (a&d) and Lys (b&c) residues in Ei→Kzl (a&b) and K→Ezl (c&d) pairs. Key: red for Glu, blue for Lys; white bars indicate the fraction of each rotamer combined in all α-helices; pale bars indicate pairs where no salt bridge is made; and dark bars indicate pairs where a salt bridge is formed. (e – f) Examples of salt-bridging pairs. For the Ei→Kzl pairs, there are two dominant rotamer combinations, with Glu (tt; tg2) plus Lys (gt; gtt) and gtt; ttt (68%) and 6 examples (24%), respectively, and two minor combinations (g&h, 1 example of each (8%)) with Glu:Lys (tt; tgg) and Glu:Lys (gt; ttt). Whereas for the K→Ezl pairs there is one clearly preferred conformation (Glu, gtt; Lys, ttt) (I, 39 examples (75%)). Only the gtt; ttt combination would be better, but inspection of molecular models revealed that with χ1 = g for Lys takes the ε-amino group too far from the γ-carboxylate of Glu to form a salt bridge. Examples were taken from the PDB as follows: (e) 1kcp A246-A250; (f) 1moq A481-A485; (g) 3n0u A177-A181; (h) 2r75 A141-A145; (i) 1egw A30-A34. Images were generated with PyMol (www.pymol.org).

Figure 5: Electrostatic potential of a model α-helix. (a) A 32-residue α-helix with its long axis aligned along z. (b) The electrostatic potential (Vs) of a single residue from the helix at x = 0 Å. (c-e) The electrostatic potential along the entire helix at varying distances (x) from the helix axis: (c) x = 0 Å; (d) x = 2.5 Å; and (e) x = 5.0 Å. Note, the increased limits for the Vs-axis in part (d). (f) The electrostatic potential arising from point charges of +0.5 and -0.5 at positions equivalent to the N- and C-terminal helical atoms on the z-axis, respectively. The electrostatic potentials in all parts (b-f) were calculated in vacuum.
Table 1: Peptide designs. Names, sequences, mean residue ellipticities at 222 nm (MRE_{222}) and 5 °C, corresponding percent helices, and midpoints of thermal denaturation curves (T_{m}) for the peptides used in this study. Conditions: phosphate buffered saline (PBS, 137 mM NaCl), pH 7.4, and 100 μM peptide concentration. The purity and identities of the peptides were confirmed by HPLC and MALDI-TOF mass spectrometry (Supplementary Figs 1–27).

To remove complicating terminal formal charges, and to ameliorate end effects in the helices, the E/K regions in all of the peptides were flanked by glycine residues, and capped with acetyl (Ac) and amide groups (NH2) at their N- and C-termini, respectively. C-terminal tryptophan (W) or tyrosine (Y) residues were included for concentration determination.

**METHODS**

**General methods.** Rink amide ChemMatrix™ resin was purchased from PCAS Biomatrix Inc (St-Jean-sur-Richelieu, Canada). Fmoc protected amino acids, peptide grade DMF and HBTU (2-(1H-benzotriazol-1-yl)-1,1,3,3-tetramethyluronium hexafluorophosphate) were obtained from ACTG Bioproducts (Hesse, UK). All other reagents were acquired from Fisher Scientific (Loughborough, UK). Peptide concentrations were determined by absorbance at 280 nm using a NanoDrop 2000 spectrophotometer (Thermo Scientific) and employing Trp ε_{280} = 5690 M⁻¹ cm⁻¹ and Tyr ε_{280} = 1280 M⁻¹ cm⁻¹.

**Peptides synthesis.** Peptides were synthesised on a 0.1 mmole scale using a Liberty CEM microwave-assisted automated synthesiser (CEM corporation, Mathews, NC, U.S.A.) with standard Fmoc chemistry (deprotection of Fmoc using 20% piperidine in DMF) and HBTU coupling (0.45 M HBTU as activator, 1M DIPEA in DMF as activator base) on a rink amide ChemMatrix™ resin. Each coupling and deprotection step was performed twice to minimise amino-acid deletions from the sequence, in addition to extra washings between steps.
Acetylations of N-termini were performed in DMF using 0.25 mL acetic anhydride and 0.5 mL pyridine. The resin was washed with DCM before peptides were cleaved by treatment with TFA (trifluoroacetic acid) / triisopropylsilane / water (90:5:5 vol%) for 3 hours under agitation. The cleavage mixtures were extracted by filtration and the resin rinsed with a further 5 mL of TFA before the total volume obtained was reduced to <5 mL under a flow of nitrogen. Crude peptides were precipitated in cold diethyl ether (~50 mL) and centrifuged to produce a pellet, which was then dissolved in 50:50 (vol%) acetonitrile:water and freeze-dried to yield a fine white solid.

**Peptide purification.** Peptides were purified by reverse phase HPLC using a JASCO chromatography system with a KYA TECH HiQ C18 column (150 μm x 10 cm) and linear gradients of various buffers depending on the peptide sequence. \((E_4K_3)_{\text{A}}, \text{A}_{4}(E_4K_3)_{\text{A}}, \text{A}_{4}(K_4E_4)_{\text{A}}, \text{A}_{4}(E_4K_3)_{\text{A}}, \text{A}_{4}(K_4E_4)_{\text{A}}, \text{A}_{4}(E_4K_3)_{\text{A}}\) and \(\text{A}_{4}(K_4E_4)_{\text{A}}\) were purified using linear gradients of water (buffer A, 13 mM ammonium acetate) and 20:80 (vol%) water:acetonitrile (buffer B, 13 mM ammonium acetate). Suitable fractions were freeze-dried and re-purified using linear gradients of water (buffer A, 0.1 vol% TFA) and acetonitrile (buffer B, 0.1 vol% TFA). All other peptides were purified directly using linear gradients of water (buffer A, 0.1 vol% TFA) and acetonitrile (buffer B, 0.1 vol% acetonitrile).

**Peptide characterisation.** Peptide identities were confirmed by mass spectrometry (matrix: α-cyano-4-hydroxycinnamic acid) using an Applied Biosystems 4700 Proteomics Analyser MALDI-TOF instrument in reflector mode. Linear mode was used for \((K_4E_4)_{\text{A}}\). Peptide purity was confirmed using a JASCO chromatography system with analytical C18 (Phenomenex Kinetex. 5 μm, 100 x 4.6 mm) or C8 (GraceVydac 5 μm, 250 x 4.6 mm) HPLC columns to >95% purity for all peptides.

**Circular dichroism (CD) spectroscopy.** CD spectra and thermal denaturation curves were recorded at 100 μM peptide concentration, pH 7.4 in phosphate buffered saline (PBS), comprising Na₂HPO₄ (8.2 mM), KH₂PO₄ (1.8 mM), NaCl (137 mM) and KCl (2.7 mM). Data were recorded using a JASCO-815 spectropolarimeter fitted with a Peltier temperature controller and baseline corrected. Spectra were recorded at 5 °C from 260 – 190 nm, with a data pitch and bandwidth of 1 nm, and a scan speed of 100 nm/min. Fraction helix (%) = 100 / \([\theta]_{222} - [\theta]_{208}\) / \((-42,500(1-(3n) – [\theta]_{208}))\). Where \([\theta]_{208} = 640–45T = 415\) deg cm² dmol⁻¹ res⁻¹ at 5 °C, and n is the number of peptide bonds including the N-terminal acetyl.\(^{48,49}\) Thermal denaturation curves were obtained by heating from 0 – 85 °C at 40 °C/hr and monitoring the absorbance at 222 nm over 1 °C increments (16 sec delay, and 16 sec response times), with CD spectra being taken at 5 °C intervals. The midpoints of thermal transitions (T_m) were taken as the maximum of the first derivative of the thermal denaturation curves. Thermal unfolding curves were also fitted to the Gibbs-Helmholtz equation to give the calculated mid-point (T_m) and enthalpies (ΔH) of folding. These fits assumed ΔC_p = 0: and used the following equations in Sigma Plot to fit the observed MRE (MRE_ave) to the predicted MRE (MRE_pred) as a function of temperature:\(^{50}\) (1) \(T = T(C) + 273.15\) and \(T_m = T_m(C) + 273.15\), to convert temperatures to Kelvin; (2) \(K = \exp((-\Delta H/(1.987^\circ T)))\), to calculate the equilibrium constant \(K\) at any temperature; (3) \(F = K/(1+K)\), to calculate fractions folded \(F\) at any temperature; and (4) \(MRE_{pred} = ([MRE_{rel} - MRE_{untl}] \times F) + MRE_{untl}\), where \(MRE_{rel}\) = MRE of 100% helical peptide, and \(MRE_{untl}\) = MRE of unfolded peptide. Starting parameters were as follows: \(\Delta H = -20\) kcal mol⁻¹; \(T_m = 30^\circ\)C, \(MRE_{rel} = -32,000\) deg cm² dmol⁻¹ res⁻¹, and \(MRE_{untl} = -5000\) deg cm² dmol⁻¹ res⁻¹.

**Analytical Ultracentrifugation.** AUC sedimentation-equilibrium experiments were conducted for \((E_4K_3)_{\text{A}}, (K_4E_4)_{\text{A}}, \text{A}_{4}(E_4K_3)_{\text{A}},\) and \(\text{A}_{4}(K_4E_4)_{\text{A}}\) at 88 μM peptide concentration (110 μL) in PBS at 20 °C in a Beckman Optima XL-A analytical ultracentrifuge using an An-60 Ti rotor, with a cell comprising an aluminium centre piece and sapphire windows. The reference channel contained 120 μL of PBS buffer. The samples were centrifuged at speeds ranging 40 – 60 k rpm. and the absorbance recorded across the cell at a radial distance of 5.8 – 7.3 cm. The absorbance was recorded after 8 hrs at each speed, followed by another scan after 1 hr to check the sample had reached equilibrium before moving onto the next speed. Data were fitted to single-ideal species using Ultrascan\(^{51}\) and 99% confidence limits obtained by Monte Carlo analysis of the fits.
NMR spectroscopy. Each of (EiK)3, (KiEi)3, (KiEi)4, and A4(KiEi)3A4 were prepared at 1 mM peptide concentration in PBS, and the pH adjusted to pH 7.4 using 10 mM NaOH. Samples were then lyophilised and reconstituted in the appropriate volume of water/D2O (90:10 vol%) to give 1 mM of peptide at pH 7.4. 330 μL of each sample was measured in a D2O-matched Shigemi tube. High-resolution 2D NOESY and TOCSY spectra were recorded for (EiK)3, (KiEi)3, (KiEi)4, and A4(KiEi)3A4, with 250 ms mixing times at 600 MHz on a Varian VNMRS or INOVA spectrometer (University of Bristol) equipped with a triple resonance cryogenically cooled probe-head and room-temperature probe, respectively. Spectral widths of 7000 Hz, 8192 complex points in f2 and 800 complex points in f1 were used for both the NOESY and TOSCY spectra. Due to signal overlap at 600 MHz data for (KiEi)3, (KiEi)4, and A4(KiEi)3A4 spectra were also recorded at 900 MHz on a Varian INOVA spectrometer (University of Birmingham) equipped with a triple-resonance cryogenically cooled probe-head. Spectral widths of 8000 Hz, 8192 complex points in f2 and 1024 complex points in f1 were used for both the NOESY and TOCSY spectra.

Δδ values were calculated as the difference in the observed chemical shifts and the sequence-corrected random-coil chemical shifts. The latter were generated using the chemical shifts for intrinsically disordered proteins (http://www1.bio.ku.dk/english/research/pv/sbin_lab/staff/MAK/randomcoil/script/).52,53

Bioinformatics. Bioinformatics analyses were performed using in-house Perl scripts on α-helices from a non-redundant set (≤30% pairwise sequence identity) of 2775 sub-1.6 Å resolution X-ray crystal structures collated from the RCSB Protein Data Bank (PDB) using PISCES.54 Only helices of 12 amino acids or longer and as identified by Promotif,55 which uses a modified algorithm56 for protein secondary structure assignment, were considered.

The distribution of hydrogen bonds made by Glu, Lys, and Asn, were identified using Promotif55 at various positions within the α-helices. Hydrogen bonds were identified using HBPlus44 and categorised according to hydrogen bond donor type as follows: no hydrogen bond; ≥1 hydrogen bond to water; ≥1 hydrogen bond to a side chain atom; and ≥1 hydrogen bond to a main-chain atom.

Identification of interacting residue pairs within helices were identified as follows: The first four residues of each helix were classed as N-terminal; residues at least 4 positions in sequence away from the N- and C-termini were considered ‘central’; and the last four residues were classed as C-terminal. Numbers of attractive E→K→i+4, K→E→i+4, K→E→i+3, and K→E→i+3 and repulsive E→E→i+4, K→K→i+4, E→E→i+3, and K→K→i+3 pairs were identified. Expected numbers of pairs were estimated using the occurrence of each residue in the whole dataset. Salt bridges were considered to be formed if the distance between Lys Nζ and the centroid of (Glu Oα1, Oε2) was ≤ 4 Å. For the repulsive pairs, the mean shortest distance between side-chain atoms was recorded, along with the percentage of repulsive residue pairs with sub-4 Å distances between them.

χ1,χ2 Side-chain rotamer distributions of for both salt-bridging and non-salt-bridged central E/K→E/Ki+3/4 pairs were categorised as follows: t, χ > 120° or χ < -120°; g*, 0° < χ < 120°; g, -120° < χ < 0° for Glu and Lys residues in all α-helices. Theoretical rotamer combinations were modelled in PyMOL (www.pymol.org) and salt-bridge potential assigned if Glu O ε1/O ε2...Lys Nζ < 4 Å. Rotamer combinations were identified using Promotif.55

Calculating electrostatic potentials. A 32 residue polyalanine α-helix was aligned along its long axis, z, (x,y = 0,0), so that the N-terminal atom was at z = 0 (all distances are in Angstroms). The relative partial charges of each dipole were approximated by single point charges on atoms as follows: HN, -0.307; N, -0.621; C, +0.582; O, -0.574.57 Overall charge neutrality was preserved with the following point charges for side chain atoms: Cα, +0.250; Hα, +0.085; Cβ, -0.416; and Hβ, +0.129. The electrostatic potential in vacuum VE(x,0,z), (i.e. Coulomb’s potential) at a distance (x) from the helix axis (y = 0) was calculated in volts using Mathematica scripts to sum the electrostatic potentials arising from all of the atomically centred point charges.
References


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α-Helix Stability


