Amino acid pairing at the N- and C-termini of helical segments in proteins

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INTRODUCTION

Helices are the most common secondary structural motif observed in folded proteins. To understand completely the helix formation and stability, the contributing factors have to be assessed thoroughly. One of those relevant factors is the formation of intramolecular hydrogen bondings between main-chain C=O and N–H groups, specially belonging to residues spaced i, i + 4 apart. As a consequence, the N-terminus has unsatisfied hydrogen-bond donors, whereas the C-terminus has unsatisfied hydrogen-bond acceptors. Another important characteristic is the presence of a net dipole moment along the helix axis with the N-terminus polarized positively and the C-terminus polarized negatively. These two characteristics are already enough to induce different amino acid occurrences in the helix. In fact, the different tendency of the amino acids to occur in α-helices is known for some decades. 1–4 Moreover, it is reasonable to accept that each individual helical position has its own role in helix stabilization and, thus, show distinct amino acid distributions. Factors such as side chain–main chain hydrogen bonds, solvent exposure, conformational entropy, and side chain–side chain interactions contribute to the diversity of positional preferences.

It has been shown that the helical occurrence of the 20 type of residues is highly dependent on the position, with a clear distinction between N-terminal, C-terminal, and interior positions. 5–16 It has also been remarked the importance of the first nonhelical positions at both termini, namely, C-cap and, especially, N-cap. 17–26

More recent studies have shown that short sequences of residues may have an important role in protein folding and stability, namely, specific pairs found in parallel β-sheets, 27 loops 28 or inter-domain linkers, 29 helix-stabilizing i, i + 4 pairs, 30 and triplets of charged residues in helices showing a cooperative effect in their stabilization. 31,32

All of these studies suggest that different local propensities of the residues may be correlated with each other, and that their presence is important not only as individual residues but as part of small sequences (pairs, triplets, etc.) that induce and stabilize the helical structure. Statistics on known protein structures provide crucial information about that correlation, and can be very useful to reveal patterns of amino acid interactions. Moreover, the data of
known protein 3D structures are continuously increasing which makes obligatory the updating of the information provided by the statistical studies.

In this work, we present an exhaustive analysis of the occurrence of amino acid pairs in a recent and updated set of helical protein segments, taking into account a few factors such as the type of helix (α-helices or 3₁₀-helices), residue separation, position, and orientation in the helix. Based on the data collected we have also carried out a search for the most probable initial and final sequences in each type of helices. The main goal of this work was to extract information from a systematic survey of helices in proteins that support some empirical rules for their formation and stabilization. Moreover, this study also presents a web site (http://www.ncc.up.pt/~nf/rps) where the results are automatically and periodically updated, following the growth of the Protein Data Bank (PDB) database.

**MATERIALS AND METHODS**

The Pisces server was used to select a subset of protein chains from the PDB with structures solved at a resolution higher than 1.6 Å, with a R-factor lower than 25%, and showing a maximum of 20% homology. Secondary structure assignments were automatically done by PDB using the Kabsch and Sander algorithm. However, to avoid the use of incorrectly classified motif sequences, the HELIX and ATOM records in the pdb files were checked using our own programs, and those presenting inconsistencies were rejected. In addition, all the pdb entries that contain more than 10% of nonstandard or undefined residues were also discarded. Consequently, the original set of 1125 protein chains was reduced to 815 as the working set (available as Supplementary Material at www.ncc.up.pt/~nf/rps) which corresponds to a total number of 186,301 amino acid residues. Two subsets were then formed with resolution higher than 1.6 Å, with a protein chains was reduced to 815 as the working set containing at least seven residues.

The relevant helical positions considered in this work are the first three residues and the first nonhelical residue at the N-terminus side of the helices N1, N2, N3, and N-cap, respectively, and the equivalent positions at the C-terminus side C3, C2, C1, and C-cap according to Richardson and Richardson’s notation. All other helical residues which evaluate the tendency of the particular pair of amino acids XY to integrate helices with X in position i and Y in position i+k (i = N-cap, N1, N2, N3, N-int, C3, C2, C1, C-cap; k = 1, 2, 3, 4). is the ratio of the relative frequency of the pair XY appearing at a particular position (i, i+k) in helices and the relative frequency of occurrence of that pair in all of the protein sequences. Therefore, the larger/smaller the value of the higher/lower the preference of the pair to occur in helical position i (a value close to 1 means no preference at all, reflecting similar distributions inside and outside the helices). On the other hand, the local propensity is defined in the subset of all the helical pairs by the ratio of the percentage of occurrence of the pair XY in a particular position i and the percentage of occurrence of that pair in the helices regardless of its position. This statistics is thus a measure of the preference of the pair for a particular position inside the helix. The term the number of occurrences of the particular pair XY found in the helices set with X in position i and Y in position i+k; in the summation is extended to all amino acids and it represents the total number of pairs in positions (i, i+k) found in helices; the number of occurrences of the particular pair XY observed in all the protein sequences and in all helices, respectively, where Y is always
found $k$ positions after $X$, independently of its position in the protein chain; in $\sum_{A,B} p_{AB}^{\text{hel}}$ and $\sum_{A,B} p_{AB}^{\text{fix}}$ the summation is extended to all amino acids, and it represents the total number of pairs observed in all the protein sequences and helices, respectively, where both amino acids are $k$ positions apart independently of the position in the protein chain. From the above definitions, the ratio of the global and local propensities gives the overall propensity of a pair to occur somewhere in a helix.

The tables with the global and local propensities calculated in our study are available in our web site at http://www.ncc.up.pt/~nlf/rps in two formats: as a Comma Separated Value (CSV) file and as a PDF file. We developed a program that automatically and periodically updates this information in three steps. In a first step, the program retrieves an updated list of protein chains with a maximum of 20% homology, whose structures were solved at a resolution higher than 1.6 Å and with a $R$-factor lower than 25%. That list may be obtained from the Dunbrack Lab website.33 In the second step, the retrieved list of protein chains is used to download the corresponding files from the PDB website.34,35 where each chain segment is identified by its secondary structure motif. Finally, the program uses each amino acid pattern occurrence frequency to compute the global and local propensities and to update the tables in our website.

## RESULTS AND DISCUSSION

### Individual propensities in $\alpha$-helices

The large and unbiased database used in this work, which includes almost 6000 helices selected with very tight criteria from the PDB,34,35 gives a strong statistical support to the observations. In general, there is good agreement between our findings and previously reported statistical analysis in protein $\alpha$-helices with smaller protein samples.5–12,14,37

Table I shows the global propensities of each individual amino acid for different positions calculated in our working data set in a way similar to Penel et al.11 At a first sight it seems that the amino acid distribution is highly dependent on the position in the helix. In fact, this was confirmed quantitatively for each of the helical positions by means of a $\chi^2$ test with 19 degrees of freedom and 0.5% level of significance. The $\chi^2$ values obtained were indeed highly significant at this level. Moreover, the $\chi^2$ values for the distribution of each amino acid over the eight extreme positions (N-cap, N1, N2, N3, C3, C2, C1, and C-cap) were also evaluated. Again, high significant $\chi^2$ values were obtained for the same level of significance, which confirm that, at this level, none of the amino acids have a uniform distribution over these eight positions, with the exception of Cys.

A close inspection of Table I shows that the results corroborate previous studies on this type of protein secondary structure.5–9,11,37

| Ncap | N1 | N2 | N3 | Nint | C3 | C2 | C1 | Ccap |
|------|----|----|----|------|----|----|----|------|
| $\alpha$ | $3_{10}$ | $3_{10}$ | $3_{10}$ | $3_{10}$ | $3_{10}$ | $3_{10}$ | $3_{10}$ | $3_{10}$ |
| C | 0.9 | 0.5 | 1.1 | 0.5 | 0.0 | 0.5 | 0.5 | 0.9 | 0.9 | 0.9 | 0.9 | 0.0 | 0.8 | 1.8 | 1.0 | 0.9 | 0.7 | 1.4 |
| P | 1.0 | 1.2 | 1.3 | 3.7 | 2.6 | 4.0 | 1.1 | 1.2 | 0.1 | 0.4 | 0.0 | 1.2 | 0.0 | 0.0 | 0.1 | 1.9 | 1.3 |
| A | 0.9 | 1.0 | 0.5 | 0.6 | 1.1 | 0.8 | 1.2 | 1.2 | 0.8 | 0.8 | 0.9 | 0.8 | 0.8 | 0.7 | 0.5 | 0.9 | 1.0 | 0.8 | 0.8 |
| T | 1.0 | 1.2 | 2.0 | 1.0 | 0.8 | 0.8 | 0.9 | 0.9 | 0.8 | 0.8 | 0.7 | 0.5 | 1.5 | 1.4 | 1.5 | 0.5 | 1.2 | 0.6 | 0.7 |
| G | 1.0 | 1.2 | 1.2 | 0.9 | 0.7 | 1.0 | 0.7 | 0.8 | 0.7 | 0.7 | 0.2 | 0.5 | 0.3 | 0.6 | 2.2 | 0.8 | 2.7 | 2.1 |
| S | 0.9 | 1.6 | 2.5 | 1.1 | 0.9 | 2.0 | 1.0 | 1.4 | 0.7 | 0.7 | 0.7 | 0.7 | 1.1 | 1.0 | 1.0 | 0.7 | 1.0 | 1.0 |
| D | 1.0 | 0.8 | 2.4 | 2.0 | 0.9 | 0.9 | 1.6 | 1.0 | 0.8 | 0.8 | 0.7 | 1.0 | 0.6 | 0.9 | 0.7 | 0.6 | 1.0 | 0.9 |
| N | 0.8 | 1.1 | 2.1 | 1.6 | 0.5 | 0.7 | 0.8 | 1.6 | 0.7 | 0.9 | 0.7 | 0.7 | 1.1 | 2.1 | 1.7 | 0.6 | 1.3 | 1.0 |
| E | 0.9 | 0.7 | 0.6 | 0.7 | 1.4 | 1.3 | 2.6 | 2.1 | 1.3 | 1.0 | 1.5 | 1.0 | 1.5 | 0.9 | 0.8 | 0.7 | 0.8 |
| Q | 0.9 | 1.0 | 0.7 | 1.6 | 1.2 | 0.8 | 1.5 | 1.1 | 1.3 | 1.1 | 1.5 | 1.4 | 1.5 | 1.3 | 0.8 | 1.0 | 0.7 |
| K | 0.9 | 1.2 | 0.7 | 0.6 | 1.0 | 1.0 | 1.0 | 0.4 | 1.1 | 1.2 | 1.6 | 1.8 | 1.5 | 1.9 | 1.2 | 1.1 | 1.0 | 0.3 |
| R | 0.9 | 1.2 | 0.7 | 0.8 | 1.0 | 0.6 | 0.9 | 0.6 | 1.3 | 0.8 | 1.5 | 1.1 | 1.3 | 0.8 | 1.2 | 1.3 | 1.0 | 0.9 |
| Y | 1.1 | 0.5 | 0.7 | 0.5 | 0.9 | 0.2 | 0.7 | 0.9 | 1.0 | 1.2 | 0.8 | 1.2 | 1.0 | 2.4 | 1.0 | 2.1 | 0.6 | 0.5 |
| H | 1.1 | 0.5 | 1.0 | 2.1 | 0.8 | 0.8 | 1.1 | 0.3 | 0.8 | 0.8 | 1.2 | 1.3 | 0.8 | 1.3 | 1.3 | 0.9 | 2.4 |
| W | 0.9 | 0.0 | 0.7 | 0.4 | 1.2 | 2.3 | 1.0 | 3.1 | 1.0 | 2.4 | 0.9 | 0.8 | 0.5 | 2.3 | 0.5 | 2.3 | 0.6 | 0.4 |
| F | 0.9 | 1.2 | 0.5 | 0.5 | 0.9 | 0.3 | 0.7 | 1.1 | 1.1 | 0.8 | 0.7 | 1.4 | 1.0 | 1.5 | 0.9 | 2.0 | 0.6 | 0.8 |
| M | 1.5 | 1.3 | 0.6 | 1.0 | 0.7 | 0.3 | 0.6 | 0.0 | 1.4 | 1.6 | 1.5 | 0.6 | 1.2 | 1.0 | 0.9 | 1.9 | 0.6 | 0.3 |
| L | 1.0 | 0.8 | 0.4 | 0.4 | 1.0 | 0.9 | 0.6 | 1.2 | 1.5 | 1.8 | 1.6 | 1.4 | 1.5 | 1.2 | 1.0 | 1.5 | 0.7 | 1.0 |
| I | 1.2 | 1.0 | 0.3 | 1.0 | 0.8 | 0.6 | 0.5 | 0.4 | 1.2 | 0.9 | 1.2 | 0.8 | 0.6 | 0.3 | 0.5 | 1.5 | 0.6 | 0.7 |
| V | 1.1 | 0.6 | 0.3 | 0.5 | 0.8 | 0.7 | 0.6 | 0.5 | 1.0 | 0.7 | 0.8 | 0.4 | 0.7 | 0.5 | 0.5 | 1.0 | 0.6 | 0.7 |
Individual propensities in $3_{10}$-helices

The size of the $3_{10}$-helices sample extracted from the original dataset is 168, which is substantially smaller than the data set of $\alpha$-helices (5388). However, after comparison of both cases, we were able to identify some common and distinct features.

A high dependence of the amino acid distribution on the position within this type of helices was also observed. In general, the distribution is similar but the particular cases of Cys and Pro seem to differ (see Table I). The former amino acid is far from having a uniform distribution, since some positions, such as C2, are preferred and others are either almost or completely avoided, namely, N2 and C3. As in the case of $\alpha$-helices, Pro shows a strong preference for N-terminus positions especially for N1 and N2, but not for N3. The case of Trp is noteworthy when compared with its occurrence in $\alpha$-helices. Its propensity is clearly lower at N1 and, remarkably, at N-cap, but higher at N2, N-int, C2, C1 and, especially, at N3.

In the $\alpha$-helices case the N-cap position was mainly occupied by big- and low-polarity amino acids ranging from Tyr to Val, whereas in $3_{10}$-helices there is a slight shift towards small side chains (from Pro to Asn). However, the case of the voluminous Trp is remarkable, because it seems to decrease in the terminal positions and to increase in the interior of the $3_{10}$-helices.

Residue pair propensities

The analysis of the 20 individual amino acid distributions in helices is not consistent enough to result in clear rules for the explanation and prediction of their formation and stabilization. That distribution may depend not only on the characteristics of the different positions but also on the interactions with neighbour residues. Obviously, pairs of amino acids are the next feature to be considered in a progressive complexity analysis. The formation of the helix within the living cell is controlled by signals, and we assume here that the N- and C-termini positions should present particular characteristics that may reflect the ability to start and to stop the folding process. The present study puts some emphasis on those helical and nonhelical positions as defined in Figure 1.

The systematic search of pairs in our database resulted in a huge amount of data which is collected in a set of tables, available as Supplementary Material at www.ncc.up.pt/~nf/rps/, from which a few relevant examples are presented here (see Tables II–V). Each table concerns either type of helical motifs ($\alpha$-helices or $3_{10}$-helices) and either type of propensity values (global or local). The first position in a pair is N-cap, N1, N2, N3, N-int, C3, C2, C1, or C-cap. The second position in a pair refers to residues at $i + 1$, $i + 2$, $i + 3$, and $i + 4$, where $i$ is the first position of the pair. In all the tables, the first residue in the pair is given by the row and the other one by the column. The 20 amino acids are sequenced from Cys to Val according to their size and polarity. The data are thus organized as a large number of $20 \times 20$ square matrices, which are periodically updated at our website as explained in the Materials and Methods section. The statistical significance of every propensity value was evaluated by calculating the associated error and $P$-value which are also available for download at the website. To make easier the reading of the propensity tables, a colour scale was introduced and those values with a $P$-value lower than 0.05 appear in bold and white.

An overview of all the matrices shows that they are not uniform neither symmetric. The discrepancy of the values suggest that there are, indeed, different preferences for the position of the pairs and that their orientation in the chain sequence is not meaningless, that is, in general, the frequency of occurrence of a particular pair XZ differs from that of ZX.

Pair propensities in $\alpha$-helices

Tables II–V concern the global propensities in $\alpha$-helices of the pairs ($i, i + 1$), ($i, i + 2$), ($i, i + 3$), ($i, i + 4$), respectively, with $i$ being the N-cap position, and they
### Table II
Global Propensities of $(i, i + 1)$ Pairs in $\alpha$-Helices ($i = N$-cap)

![Table II](image)

### Table III
Global Propensities of $(i, i + 2)$ Pairs in $\alpha$-Helices ($i = N$-cap)

![Table III](image)

### Table IV
Global Propensities of $(i, i + 3)$ Pairs in $\alpha$-Helices ($i = N$-cap)

![Table IV](image)

### Table V
Global Propensities of $(i, i + 4)$ Pairs in $\alpha$-Helices ($i = N$-cap)

![Table V](image)
are presented here as representative examples. A preference of small and polar residues such as Thr, Ser, Asp, and Asn for position N1 is noticeable regardless the residue at N-cap (see Table II). The Cys residue at N-cap is the only exception as it pairs almost exclusively with Asp ($P_{\text{HNcapPN2}} = 3.4$). In general, the values of Asn are compared with those of Thr, but Ser and Asp show higher propensities to occur at N1. On the other hand, that position seems to be avoided by big and hydrophobic residues, namely, Trp, Phe, Met, Leu, Ile, and Val.

These results have a reasonable agreement with a previous experimental work on polyalanine-based peptides but new features have to be referred. In fact, Thr is more favorable at position N1 than in the helix interior but not in N2; in this latter position the highest propensity for Thr occurs only when Met is in N-cap (see Table III).

When the second position in the pair is N2, then a clear preference for Pro is observed, except if His is in N-cap, $P_{\text{HNcapPN2}} = 0.9$ (see Table III). We are currently trying to find a chemical/structural explanation for the particular behavior of the H$_{\text{N-cap}}$P$_{\text{N2}}$ pair.

Some interactions between residues have already been recognized as important structural factors of helix formation and stabilization. A typical example is the capping box, where the side chain of a residue in N-cap makes a H-bond with the backbone amino group of the N3 and, reciprocally, the side chain of N3 establishes a H-bond with the backbone amino group of the N-cap. The pair S$_{\text{N-cap}}$P$_{\text{N3}}$ was identified in a previous analysis on a smaller dataset as particularly relevant to define a capping box motif, presenting a value of 13 for its global propensity. Concerning this particular point, the results that come out of our analysis are based on the data shown in Table IV. In fact, all types of residues show a high propensity to appear at N-cap associated with Glu at N3. However, the highest pair propensity does not appear with Ser ($P_{\text{HNcapPn3}} = 2.5$) but with Met ($P_{\text{HNcapPn3}} = 4.4$). These results, on one hand, are reflections of the high ability of Glu in N3 position to be involved in the formation of capping box motifs due to its long side-chain terminating in a carboxylate group. On the other hand, Asp, which has got a shorter side-chain, shows lower pair propensities for the same position as a consequence of its lower potential to form a capping box motif, as referred in a previous study.

Considering pairs starting at N1, it is found, as expected from the results above, predominance of small and polar residues in the first position (Asn, Asp, Ser, and Thr). They show a great variability in propensities, but the strongest couplings appear for Pro in position N2, and Glu in N3 and N4.

At the interior of the helices the pairs are mainly formed by large and polar residues and by Ala as well, but almost never by Pro. On the other hand, Gly appears almost exclusively in positions $i + 3$ and, especially, $i + 4$ of these inner pairs, and with different preferences to the residue in the first position $i$. Interesting is the high global propensities of the pairs ($C_{\text{Nint}}$, $G_{i+4}$), ($A_{\text{Nint}}$, $G_{i+4}$), ($M_{\text{Nint}}$, $G_{i+4}$), and ($L_{\text{Nint}}$, $G_{i+4}$) when compared with the corresponding low values for ($N$, $i + 3$) pairs.

The findings are based on the high statistical significance of the pair propensities as shown in the corresponding tables available in the Supplementary Material at www.ncc.up.pt/~nf/rps.

**Pair propensities in 310-helices**

Concerning the 310-helices, the dimension of the sample is substantially smaller (168 helical sequences) than the previous case (5388), which prevents us to present here an accurate statistical comparison. However, since the website is being periodically updated, the analysis will be done in the follow-up work of this study. Nevertheless, with the whole set of tables stored at present it is possible to compare both cases with some care (see Supplementary Data at www.ncc.up.pt/~nf/rps). A clear distinction from the previous case is easily perceived at first sight, which is the large number of cells with zero value. In the propensity tables, a value of zero means that the pair is observed nowhere in the chain but never at that particular position. Although this may be a consequence of the small dimension of the sample, the distribution of the propensity values among all the 400 possible pairs is very irregular, and consequently some preliminary conclusions may be inferred.

The analysis of the global pair propensities revealed a preference for small amino acids to occur at N-terminus positions. The particular case of Cys seems to be distinct from the others as we already pointed out while analyzing the values of individual propensities. When Cys is at N-cap position there is a clear preference to couple with Cys in N1. Reciprocally, it appears at N1 position only when Cys, and also Lys and Phe but in less extent, is preceding it in N-cap.

In the previous section it was referred that Pro prefers the N2 position in α-helices, but in the case of the 310-helices it occurs almost exclusively at N1 and N2 with similar preferences. Big and aromatic amino acids show also a distinct behavior, but the particular case of Tryptophan is noteworthy. It never appears at N-cap position, and at N1 it only pairs with Val, Asp, Ile, and Lys at N2, N3, N4, and N5, respectively. On the other hand, at C-terminus, it appears at C-cap only paired with Asn, Lys, and Tyr at C1, C2, and C3, respectively.

**Relevant short amino acid sequences at N-terminus and C-terminus**

The positions at the N- and C-termini of the helices are nowadays recognized as to play an important role in
their stabilization. We assume here that the sequences of amino acids at both termini may be highly correlated with the start and stop processes of the helix formation, as well as with its thermodynamic stabilization. Therefore, we looked for relevant short sequences at both termini of the helices, including the first nonhelical positions N-cap and C-cap, from which a few representative examples are reported in Table VI. To save computation time, we did not consider all possible sequences but only those that had the potential to be more frequent based on the high propensity values of the pairs they include. Starting at a particular position, the methodology employed here was able to select the next amino acid in the sequence to which a high pair propensity is associated, and continued forward using the same criterion in order to establish a high probable pathway. Then, an automatic search of those selected sequences was carried out in all protein chains of the working sample.

Table VI shows that the sequence M N-capTN1EN2EN3 (MTEE--) never appears even though the partial pair propensities in α-helices are particularly high, namely, $P_{MN\text{cap}T_{N1}} = 3.1$, $P_{T_{N1}E_{N2}} = 4.0$, and $P_{E_{N2}E_{N3}} = 3.1$. The same happens to many other sequences (like the next four examples in the table) meaning that high propensities of the constituent pairs do not guarantee a high probable sequence.

Other sequences occur with reasonable frequency but rarely or never inside a helix, which is the case of V_PD-- at the N-terminus (in this notation the broken line indicates the direction of the remaining helical sequence and a dash means a position that may be occupied by any residue).

On the other hand, high propensity pairs may give a clue to find positive examples as, for instance, the sequences _SPE--, _TPE--, _DPE--, and _SRE--. These initial triplets of amino acids appear almost exclusively within helices and approximately half of the observations occur exactly at N1 despite the large number of other positions available in the helices. This is not surprising considering the known ability of Glu to be involved in the formation of a capping box motif at N3, as discussed earlier, but only these four triplets appear more frequently.

At the C-terminus we have found triplets with high incidence but low local preference within the helices, which are the cases of _ALA_, _EAA_, _LAK_, _IAR_, _ALM_, and _WLK_. There is another group of short sequences that present low overall occurrences but very high preferences to positions at the interior of helices or their termini. For example, _ENP appears eight times in α-helices, out of a total of 33, and exactly with Pro in C-cap.

The frequencies of occurrence of these short sequences in our protein working set suggest a probability function which is far from a uniform distribution because helical motifs represent only 36% of the total length of all the protein chains considered (66,340 amino acid residues out of a total of 186,301).

Another example that may illustrate very well the specific features of small sequences in helices is --EKYP. It is observed only five times in proteins, but four of them appear exactly with Pro at C-cap position of α-helices. This sequence adopts a typical conformational as shown in Figure 2 (corresponding to 1JR8 entry of PDB), and

| Sequence | α-Helix | $\alpha'$-Helix |
|----------|---------|----------------|
| In protein chains | In local | In helix | In local | In helix |
| Ncap | N1 | N2 | N3 | C3 | C2 | C1 | Ccap | Ncap | N1 | N2 | N3 | C3 | C2 | C1 | Ccap |
| _SPE_ | 51 | 24 | 29 | 0 | 0 |
| _TPE_ | 64 | 24 | 28 | 1 | 1 |
| _DPE_ | 60 | 23 | 24 | 1 | 1 |
| _SRE_ | 44 | 10 | 23 | 0 | 0 |

Table VI shows the number of occurrences of some short sequences selected from the analysis of the global pair propensities.
for which the corresponding data are summarized in Table VII. The structure shows an interesting interaction between the two distended side chains of Lys and Tyr, where the average distance $\langle \text{Lys}_N \cdot \text{O}_h(\text{Tyr}) \rangle$ is 3.7 Å.

The single case where the sequence does not appear at N-cap position is observed in the 1MIN entry. The four-residue sequence defines a coil structure between two $\alpha$-helices, where Lys and Tyr adopt a conformation really distinct from that shown in Figure 2, which prevents the interaction between their side chains. To verify the specificity of the conformation, we surveyed the working sample of protein chains for all the pairs $\text{Lys}_i \cdot \text{Tyr}_{i+1}$ and $\text{Tyr}_i \cdot \text{Lys}_{i+1}$. We have found 659 examples of such pairs but only six different cases, not related to those mentioned previously, show geometries of interaction similar to that gathered in Table VII, namely, $d[(\text{Lys})_N \cdot \text{O}_h(\text{Tyr})] < 4$ Å, $-165^\circ < (\text{Lys})X_{1,4} < +165^\circ$ and $-60^\circ < (\text{Tyr})X_{1,2} < +100^\circ$. In addition, these six examples appear in helices or in turns. These results suggest that the sequence $\text{EKYP}$ with this particular side-chain interaction between Lys and Tyr has indeed a high specificity to C-terminus position.

### CONCLUDING REMARKS AND FUTURE DIRECTIONS

This work presents the most comprehensive analysis of helical motifs in proteins undertaken to date. An exhaustive study of the frequency of occurrence of individual amino acids and all possible pairs was carried out on a set of 5556 helices. Pairs of type $(i, i+1), (i, i+2), (i, i+3)$, and $(i, i+4)$ were considered starting at relevant positions near or within helices: N-cap, C-cap, N1, N2, N3, C3, C2, C1, N-int. The protein sample used in this work was sufficiently large and unbiased which gives confidence to the final results expressed in terms of global and local propensities. Some general features of residue pairs at both N- and C-termini were identified.

Some sequences, although occurring rarely, seem to play a very specific role because they are observed always at the same position of helices. This is the case, for instance, of the sequence $\text{EKYP}$ for which four out of five times it is observed at the same C-termini position of $\alpha$-helices.

In this work, a bioinformatics tool was developed, which can automatically follow the growth of the PDB database and update the propensities values in both $\alpha$-helices and $\beta$-helices. The results produced by the statistical analysis are periodically updated and stored as Supplementary Material in our website http://www.ncc.up.pt/~nil/rips.

The amount of information collected is huge and will need a further automatic analysis using, for instance, Inductive Logic Programming (ILP) algorithms, in order to obtain useful predictive rules. The physico-chemical characteristics of the 20 amino acids and the data concerning their individual and pair propensities generated in this work would be crucial to start the ILP studies.

### Table VII

**Main Characteristics of the $\text{---EKYP}$ Sequence**

| PDB entry | Sequence location | (Lys)$_N \cdot \text{O}_h(\text{Tyr})$ | $\chi_1$ | $\chi_2$ | $\chi_3$ | $\chi_4$ | $\chi_{1,2}$ |
|-----------|-------------------|---------------------------------|---------|---------|---------|---------|-------------|
| 1H16      | (A) 723–726       | 3.96                            | $-178.9$| $176.7$ | $174.2$ | $-177.3$| $-57.5$    |
| 1JR8      | (A) 66–89         | 3.39                            | $179.6$ | $174.5$ | $-177.8$| $-179.7$| $-55.9$    |
| 1T8U      | (A) 69–72         | 3.90                            | $-175.1$| $169.7$ | $176.6$ | $164.9$ | $-54.7$    |
| 1XY       | (A) 129–132       | 3.82                            | $-173.0$| $172.4$ | $-176.9$| $-175.2$| $-56.7$    |

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With this approach we aim to find some general rules that can be applied to any amino acid sequence in order to predict the stability of helical motifs.

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