Evidence for the preferential reuse of sub-domain motifs in primordial protein folds

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Abstract
A comparison of protein backbones makes clear that not more than approximately 1400 different folds exist, each specifying the three-dimensional topology of a protein domain. Large proteins are composed of specific domain combinations and many domains can accommodate different functions. These findings confirm that the reuse of domains is key for the evolution of multi-domain proteins. If reuse was also the driving force for domain evolution, ancestral fragments of sub-domain size exist that are shared between domains possessing significantly different topologies. For the fully automated detection of putatively ancestral motifs, we developed the algorithm Fragstatt that compares proteins pairwise to identify fragments, that is, instantiations of the same motif. To reach maximal sensitivity, Fragstatt compares sequences by means of cascaded alignments of profile Hidden Markov Models. If the fragment sequences are sufficiently similar, the program determines and scores the structural concordance of the fragments. By analyzing a comprehensive set of proteins from the CATH database, Fragstatt identified 12,532 partially overlapping and structurally similar motifs that clustered to 134 unique motifs. The dissemination of these motifs is limited: We found only two domain topologies that contain two different motifs and generally, these motifs occur in not more than 18% of the CATH topologies. Interestingly, motifs are enriched in topologies that are considered ancestral. Thus, our findings suggest that the reuse of sub-domain sized fragments was relevant in early phases of protein evolution and became less important later on.

Keywords
ancient modules, domain evolution, fold space, protein evolution

1 | INTRODUCTION
Proteins are indispensable to maintain the complex processes constituting life. The relevance and functional scope of proteins is exemplified by the fact that the human proteins consist of appr. 70,000 splice variants and that enzymes catalyze about 4000 reactions. Some complex tasks like protein biosynthesis via ribosomes or the assistance of the assembly of macromolecular structures via chaperones require large multiprotein complexes. However, most functions are fulfilled by proteins that exist as monomers or belong to small heterooligomers, which requires the existence of a large number of highly specific proteins.

In contrast to the rich diversity of protein functions, the number of unique folds, that is, specific arrangements of major secondary elements with the same topological connections, is drastically smaller. The claim from 1992 by Cyrus Chothia that nature is restricted to
approximately 1000 folds\(^6\) has stood the test of time: The latest version (SCOP 2) of the SCOP database lists not more than 1388 different folds.\(^7\)

Due to their evolutionary relationships, protein sequences and structures resemble each other, if they share a common ancestor. Thus, a certain level of sequence\(^8\) or structure\(^7\) similarity signals homology of two proteins. Additionally, the composition of most proteins complies with a common architectural scheme, comprising one or a sequential series of several building blocks named domains. Domains are independent evolutionary and functional entities, typically consist of 100 to 250 residues and fold into a compact 3D-structure independently of neighboring elements.\(^10\) As evidenced by the Pfam database, more than 18 000 functionally different domains exist in nature\(^11\) and the combination of these basic building blocks most plausibly explains the evolution of larger proteins.\(^12\) Thus, the dominant processes that extend the repertoire of protein functions are duplication of sequences coding for one or several domains, their evolutionary modification, and for some of them their subsequent combination.\(^13\)

Whereas this model for the evolution of multi-domain proteins is generally accepted and experimentally proven,\(^14,15\) the evolution of domains is still a topic of recent studies. The probability that a functional protein can arise de novo has been considered unlikely,\(^16\) which suggests in analogy to the evolution of multi-domain proteins that a reuse of basic building blocks also drives the genesis of individual domains.

In a pioneering analysis, Eck & Dayhoff have identified in ferredoxin a short repeat element binding an iron-sulfur cluster and suggested the doubling of a shorter protein for the evolution of ferredoxin.\(^17\) By screening crystal structures for \(C_{\alpha} - C_{\alpha}\) contacts less than 10 Å apart, structurally heterogeneous loops with a length of 20 to 50 residues have been detected and proposed as being remnants of prebiotic ring-like elements, whose combination resulted in modern folds.\(^18\) Based on these findings, elementary functional loops (EFLs) have been characterized, these are structural-functional units that possess a closed loop structure and bear one or few residues that are involved in an elementary function.\(^19\) Utilizing position-specific scoring matrices, 525 profiles of EFLs have been specified that include 249 ones involved in binding nucleotide-containing ligands.\(^20\)

Alternatively, methods that compare structures and sequences irrespective of their functional role have been devised for the identification of protein segments of sub-domain size. The comparison of structures is often based on their superposition and the utilization of a similarity measure.\(^21\) An even stronger indicator signaling a common ancestry of proteins is significant sequence similarity.\(^8\) Due to their superior sensitivity, profile hidden Markov models (HMMs) are the method of choice for the detection of remote sequence homology.\(^22\) Although several studies\(^23-27\) used HMMs to account for sequence homology, the number of fragments that were found in similar sets of protein folds differ drastically. The database Fuzzle comprises more than 1000 fragments of various length,\(^26\) whereas not more than 40 fragments to be found in 118 folds have been reported by A. Lupas and coworkers.\(^23\)

We were interested in designing and utilizing an algorithm of highest sensitivity, whose application requires minimal user intervention to enable a comprehensive comparison of folds. This is why we implemented Fragstatt, which is based on cascaded HMMs. We concentrated on the analysis of clearly non-homologous proteins and assessed the detected motifs by means of statistical methods. After clustering of highly similar motifs, the large-scale application of Fragstatt identified only few fragments in a small number of putatively ancestral folds.

# METHODS

## 2.1 Design of Fragstatt

The algorithm Fragstatt was implemented in Python and consists of the four modules GraphCreator, PathFinder, PathAnalyzer, and MotifDetector; compare Figure 1. The input are two HHblits cascades \(c\)-tree\(P_i\) and \(c\)-tree\(P_j\) for the proteins \(P_i\) and \(P_j\) as roots. The non-root nodes of the trees represent HMMs and the edges local HHblits alignments of two HMMs. The first module, GraphCreator combines the two cascades to one graph by identifying common nodes that represent the same protein family. The second module, PathFinder identifies candidate paths connecting \(P_i\) and \(P_j\) by performing a depth-first search with a maximum search depth of four. The graph may contain a large number of parallel edges, which are caused by several HHblits hits that align various subsequences of \(P_i\) and \(P_j\). If a graph contains more than 10 000 edges, PathFinder randomly samples 10 000 of them. For each candidate path, PathFinder determines the overlap between the local alignments and discards all paths overlapping in less than five residues. As a first step, PathAnalyzer converts each of the paths identified by PathFinder to the multiple sequence alignment MSA\(_{ij}\) by joining the local pairwise alignments generated by HHblits. Subsequently, PathAnalyzer creates a matrix \(H_{ij}\), where each \(H_{ij}[k, l]\) indicates how often residues \(r_i^k\) and \(r_j^l\) from \(P_i\) and \(P_j\) were aligned in MSA\(_{ij}\). Based on these \(H_{ij}[k, l]\) values, the last module, MotifDetector, deduces an optimal traceback by using a modified Smith-Waterman approach.\(^28\) Begin and end of the traceback specify two fragments \(F^i_1 = P_i [b_{ij}, e_{ij}]\) and \(F^j_1 = P_j [b_{ij}, e_{ij}]\) that represent the common motif \(M_{ij}\). The two fragments are then superimposed by means of TM-align to determine their TM-score.\(^9\)

## 2.2 Determination of prototypical CA_motifs

CD-HIT\(^29\) with a cut-off of 90% sequence identity was applied to all entries of CA_motifs in order to reduce redundancy on the sequence level and the cluster centers and cluster members were stored. For all pairs of fragments, their TM-score was determined and a distance
matrix $D_{ij}$ was computed for all fragments specifying one of $n$ cluster centers according to

$$D_{ij} = \begin{cases} 
1 & \text{if } 1.0 - \text{TM-score} \left( F_i, F_j \right) > 0.45 \\
0 & \text{otherwise} 
\end{cases}$$

which implements a minimal TM-score of 0.55.

This distance matrix was used for a DBSCAN clustering. The parameter $minPts$, which defines the minimum number of neighbors a point must have to be considered as core point, was set to 2. The parameter $E$ defines the neighborhood radius and was chosen by optimizing the number of clusters and the average TM-score of all pairs within each cluster. Fifty-one percent of the initial motifs could be assigned to a cluster.

### 2.3 Searching multi-motif proteins

For each protein $P_i$, all fragments $F_i^j = P_i | [b_{ij}, e_{ij}]$ were deduced from CA motifs. To assess the number of positionally distinct sets of fragments, their Hausdorff distance was calculated pairwise according to:

$$d_{\text{Hausdorff}} \left( F_i^j, F_l^j \right) = \max \left( |b_{ij} - b_{lj}|, |e_{ij} - e_{lj}| \right)$$

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**FIGURE 1** Principles of motif detection by means of Fragstatt. The first module, GraphCreator takes as input two HMM cascades that is, c-tree($P_i$) and c-tree($P_j$) and builds a combined graph by combining corresponding nodes. The module PathFinder searches for paths linking $P_i$ and $P_j$. In this example, three paths were detected. PathAnalyzer combines the pairwise alignments of a path to a multiple sequence alignment (MSA) and determines a matrix $H_{ij}$, where $H_{ij}[k, l]$ indicates for each pair of residues $r^i_k$ and $r^j_l$ from $P_i$ and $P_j$ how often they were aligned. The last module, MotifDetector utilizes $H_{ij}$ to determine motifs by a traceback similar to a Smith-Waterman algorithm. The corresponding fragments are finally superposed to compute the TM-score via TM-align [Color figure can be viewed at wileyonlinelibrary.com]
The corresponding distance matrix $D_i$ was used for a DBSCAN clustering. The parameter $minPts$ was set to 2 and $E$ was set to 10. For each protein, the result of the clustering was evaluated and those proteins with at least two positionally distinct motifs were saved in a list of multi-motif proteins.

3 | RESULTS

3.1 | Design principles of Fragstatt and benchmarking

For the subsequent usage, we defined the terms motif and fragment as follows: A super-secondary structure, which can be found in at least two proteins will be called a motif. The manifestation of a certain motif in a protein will be called a fragment. For a protein chain $P_i$ with known sequence and structure, the manifestation $F_i^j$ of motif $M_k$ will be referred to as

$$F_i^j = M_k(P_i) = P_i[b_{ik}, e_{ik}]$$

(3)

where $b_{ik}$ and $e_{ik}$ are the begin and end position (respective residues) of fragment $F_i^j$ in $P_i$.

Fragstatt ( acronym for fragment instantiation) was customized to detect in non-homologous proteins relatively short, but similar subsequences that possess a matching 3D structure. In agreement with previous findings, the detected fragments must comprise at least 15 residues, like small super-secondary structure elements. As fragments are considered of sub-domain size, their maximal size was limited to 60 residues. The 3D structures of fragments were compared by means of TM-align; generally, domains with a TM-score $> 0.5$ can be assumed to possess the same fold. We regarded motifs as structurally equivalent, if the TM-score was $> 0.55$, because this cut-off corresponds to a $P$-value of 0.01 which we deduced from a null-model of fragment comparison; see below.

The standard principle for the detection of common motifs relies on a pairwise comparison: Two non-homologous proteins $P_i$ and $P_j$ distinguished by different folds, are scanned for local sequence similarities. If both the sub-sequences and the sub-structures related to two fragments $F_i^j$ and $F_j^i$ in $P_i$ and $P_j$ possess a certain similarity, the fragments are considered as representatives of the motif $M_k$ that corresponds to the mapping $M_k(P_i) \rightarrow M_k(P_j)$.

Fragstatt identifies local sequence similarities by means of HHblits, which is a state-of-the-art method for the comparison of HMMs and the detection of remote homology. In order to increase the sensitivity beyond previous studies, Fragstatt cascades several HMM searches, because this approach has proven enhanced sensitivity in remote homology detection.

Combining the outcome of cascading HMM alignments (compare Figure 1) related to a protein $P_i$, results in a tree-like structure c-tree ($P_i$), where $P_i$ is the root, and all internal nodes $f^i_j$ are intermediates sharing local alignments with the corresponding parent and the child nodes. For the final step of motif detection, the 3D structures of root proteins $P_i$ are required. Thus, these proteins were taken from PDB.

In order to achieve higher coverage, the HMMs specifying internal nodes were taken from Pfam, which is more comprehensive than PDB, as it also contains proteins whose structure is unknown. In a preprocessing step, for each protein $P_i$, a c-tree($P_i$) with a depth of three was computed and all hits with an E-value below 1.0 were accepted. A subsequent depth-first search determined for each pair $P_i$ and $P_j$ all paths $P_i \rightarrow f_i^j \rightarrow \ldots \rightarrow f_j^i \rightarrow P_j$ consisting of maximally four local HMM alignments. If these local alignments were consistently overlapping each other, they were merged to a multiple sequence alignment (MSA). A further processing of the MSA (see Methods [section 2]) led to the identification of fragments $F_i^j = P_i[b_{ik}, e_{ik}]$ and $F_j^i = P_j[b_{jk}, e_{jk}]$. Finally, the corresponding substructures were compared by means of TM-align$^9$ and if the TM-score was $> 0.55$, $F_i^j$ and $F_j^i$ were considered as instantiations of motif $M_k$.

In order to confirm that the results generated with this fully automated Fragstatt protocol are comparable to those that were manually curated by experts in their fields, we recapitulated two recently reported experiments; for details see Supporting Information (Appendix S1). To begin with, we confirmed that Fragstatt is able to detect a motif shared between the $(i)x_0$-barrel and the flavodoxin-like fold. Moreover, we tried to recapitulate a set of 40 putatively ancestral peptide-sized motifs (which we named AncPept) that was deduced from a comprehensive comparison of SCOP domains by means of HHsearch. Fragstatt recapitulated 85%, namely 34 of the 40 AncPept motifs and most plausibly, due to the manual curation and compilation of AncPept, these data were not fully reproducible with the automatic approach of Fragstatt. In summary, we concluded that the sensitivity of Fragstatt is comparable to alternative methods and that Fragstatt is capable of finding shared fragments in a fully automated manner, which was not feasible so far.

3.2 | A large-scale Fragstatt scan found motifs in 26 CATH architectures

As it was our aim to survey most of the “protein universe”, that is, a large number of folds observed in nature, we had to carefully select the protein chains to be analyzed. In order to compile a representative but redundancy-free dataset, we opted for an analysis of single-domain proteins and the hierarchical classification of their structures by means of CATH.$^{37}$ From top down, the first three CATH levels are class (derived from secondary structure content), architecture (describing the gross orientation of secondary structures, independent of connectivity) and topology (clusters structures into fold groups according to their topological connections and numbers of secondary elements).$^{38}$

The analysis of an initial dataset that was compiled on the lowest CATH level we considered, namely on topologies, gave rise to 77 million protein pairs to be analyzed and around 900 000 (1.2%) motifs that occurred in different CATH topologies were detected by MotifDetector. We named this set $CT\_motifs$; however, the visual inspection of several motifs made clear that the CATH level
“topology” is inappropriate for the automatic identification of motifs to be found in evolutionary unrelated folds: A large number of these motifs occurred in proteins that are evolutionary closely related although they belong to different CATH topologies. A typical example are beta-propeller proteins that consist of 4 to 12 radially arranged beta-blades.39 From an architectural point of view, the proteins share a common scheme of composition, namely the annular repetition of blades. It is assumed that beta-propeller proteins arose divergently and evolved by amplification and diversification after the formation of a prototypical blade motif.39-41 As we were interested to identify motifs shared between evolutionary unrelated folds, we had to choose the next level of the CATH hierarchy, namely the level of different architectures to select protein pairs \( P_i \) and \( P_j \). Unfortunately, this choice did not solve the problem completely, because beta-propellers are, depending on the number of blades, grouped into different CATH architectures. We could circumvent this issue by utilizing a “black list” containing combinations of architectures to be removed from the analysis. Note that this list does not exclude the comparison of, for example, propellers with other architectures.

Applying this criterion and accepting only fragments of length 15-60 residues, whose structural superposition resulted in a minimal TM-score of 0.55, Fragstatt found 12 532 motifs. This set, which we named \( CA_{\text{motifs}} \) because it arose by comparing different architectures, comprises 1.4% of the 900 000 \( CT_{\text{motifs}} \) Fragstatt found by comparing proteins with different CATH topologies. \( CA_{\text{motifs}} \) occur in 2870 different PDB chains that belong to not more than 26 CATH architectures and 245 CATH topologies. These low rates strongly suggest that motifs are rarely shared between proteins possessing different CATH architectures.

3.3 | \( CA_{\text{motifs}} \) are spread unevenly among CATH architectures

CATH architectures differ quite drastically in the number of assigned protein domains. Extreme examples are architectures 3.20 (alpha-beta barrel) which includes 16 668 domains and architecture 2.110 (4 propeller) which is comprised of not more than 55 domains. We wanted to confirm that our focusing on single-domain proteins did not introduce a bias and additionally assess the distribution of motifs identified by Fragstatt. This is why we determined for proteins chains their CATH classes and compared the frequency distributions of all CATH entries, single-domain proteins, and the proteins possessing \( CT_{\text{motifs}} \) or \( CA_{\text{motifs}} \). As Figure 2 confirms, the histograms of CATH classes resulting from the full content of the database and the single-domain proteins are nearly identical. The \( CT_{\text{motifs}} \) are over-represented in the class of mainly alpha and under-represented in the class of mainly beta proteins. In contrast, the distribution of the \( CA_{\text{motifs}} \) agrees well with the assignment of the full CATH set. Thus, we concluded that by focusing on single-domain proteins, we selected a sample that adequately represents the full set of protein structures. Moreover, we did not observe a striking enrichment of motifs in one of the CATH classes.

\[
E_i = \log \frac{f(CA_{\text{motifs}})}{f(CA_i)}
\]

where \( f(CA_i) \) is the frequency of all single-domain proteins belonging to a certain CATH architecture \( CA_i \).

In contrast, the analogous assessment of assigned proteins on the architecture level (compare Figure S3) revealed a severe bias for the full CATH content: The five most populated architectures are 3.40 (3-layer (ab) sandwich), 1.10 (orthogonal bundle), 3.30 (2-layer sandwich), 2.60 (sandwich), and 1.20 (up-down bundle). About 65% of all domains belong to one of these five architectures. Thus, in order to normalize the frequency \( f(CA_{\text{motifs}}) \) of a motif determined on the level of a CATH architecture, we calculated an enrichment factor \( E_i \) according to:

In Table 1, the architectures are sorted according to the enrichment factor \( E_i \). A positive number indicates that an architecture \( CA_i \) is overrepresented among the single-domain proteins contributing \( CA_{\text{motifs}} \). In contrast, a negative \( E_i \)-value signals that fewer than expected proteins with architecture \( CA_i \) possess a \( CA_{\text{motif}} \). The overrepresented architectures, ordered from high to low enrichment factor, are 2.140 (8 propeller), 3.20 (alpha-beta barrel), 1.25 (alpha horseshoe), 4.10 (irregular), 2.130 (7 propeller), 2.40 (beta barrel), 1.20 (up-down bundle), 1.10 (orthogonal bundle), and 2.70 (distorted sandwich). Except of architecture 2.140 (8 propeller, \( E_i = 1.62 \)), sparsely populated architectures are underrepresented and the two highly populated “mainly alpha” architectures 1.10 (orthogonal bundle, \( E_i = 0.29 \)) and 1.20 (up-down bundle, \( E_i = 0.30 \)) are enriched. Moreover, most of these overrepresented architectures either show an internal symmetry like alpha-beta barrels or propellers or are repetitive like alpha horseshoes and up-down bundles.
As Fragstatt deduced motifs from protein pairs ($P_i$, $P_j$) possessing different CATH architectures, we wanted to know which pairs of architectures were linked, and if some combinations were particularly abundant. We counted how many motifs were detected for each combination of CATH architectures, listed the results in Table S3, and visualized the numbers as a network in Figure 3. Relatively few combinations of architectures constitute the greatest part of all hits: Most abundant are the following five combinations of two CATH architectures: 3.20 (alpha-beta barrel) $\rightarrow$ 3.40 (3-layer(aba) sandwich), 1.10 (orthogonal bundle) $\rightarrow$ 3.40 (3-layer(aba) sandwich), 1.20 (up-down bundle) $\rightarrow$ 1.25 (alpha horseshoe), 1.20 (up-down bundle) $\rightarrow$ 1.10 (orthogonal bundle), and 1.10 (orthogonal bundle) $\rightarrow$ 3.30 (2-layer sandwich). This overrepresentation of few folds has a marked impact on the length distribution of the motifs. A strong bias originates from the comparison of CATH architectures 3.20 $\rightarrow$ 3.40, because the outcome dominates the motifs longer than 40 residues (compare Figure S4).

In sum, five combinations constitute 57% of all motifs. With a frequency of 25%, motifs that occur both in architectures 3.20 (alpha-beta barrel) and 3.40 (3-layer(aba) sandwich) are the most abundant ones and one representant is the $\beta\alpha_2$ motif identified earlier. Additional five of the most frequent combinations belong to the class "mainly alpha"; moreover, symmetrical architectures like 3.40 (alpha-beta barrel) or repetitive architectures like 1.20 (up-down bundle) and 1.25 (alpha horseshoe) are abundant. Of the alpha helical proteins, architecture 1.10 (orthogonal bundle) seems most central, showing strong connections to architectures from all classes; compare Figure 3. The mean number of edges is 8.5, and we assume that the

| CATH architecture $CA_i$ | $E_i$ | $f(\text{CA_motifs})_i$ | $f(\text{CA})_i$ | $f(\text{CA_SD})_i$ | $f(\text{CT_motifs})_i$ |
|--------------------------|------|----------------------|----------------|----------------|------------------|
| 2.140 (8 propeller)     | 1.62 | 0.08                 | 0.04           | 0.02           | 0.15             |
| 3.20 (alpha-beta barrel)| 1.23 | 13.98                | 2.43           | 4.08           | 7.21             |
| 1.25 (alpha horseshoe)  | 1.13 | 7.63                 | 1.19           | 2.45           | 3.71             |
| 4.10 (irregular)        | 0.80 | 3.19                 | 0.70           | 0.43           | 1.56             |
| 2.130 (7 propeller)     | 0.78 | 1.26                 | 0.32           | 0.58           | 1.63             |
| 2.40 (beta barrel)      | 0.33 | 5.62                 | 4.99           | 4.03           | 2.79             |
| 1.20 (up-down bundle)   | 0.30 | 10.98                | 6.91           | 8.11           | 10.89            |
| 1.10 (orthogonal bundle)| 0.29 | 20.35                | 15.55          | 15.25          | 29.34            |
| 2.70 (distorted sandwich)| 0.07 | 0.65                 | 0.60           | 0.60           | 0.18             |
| 2.120 (6 propeller)     | −0.07| 0.33                 | 0.10           | 0.36           | 1.31             |
| 3.40 (3-layer(aba) sandwich)| −0.07| 18.78               | 22.10          | 20.05          | 17.42            |
| 3.80 (alpha-beta horseshoe)| −0.10| 0.47                 | 0.25           | 0.52           | 0.17             |
| 3.10 (roll)             | −0.52| 3.54                 | 3.28           | 5.93           | 3.91             |
| 3.90 (alpha-beta complex)| −0.58| 2.19                 | 5.13           | 3.93           | 1.87             |
| 3.30 (2-layer sandwich) | −0.69| 6.87                 | 17.51          | 13.66          | 10.08            |
| 2.102 (3-layer sandwich)| −0.75| 0.06                 | 0.12           | 0.12           | 0.61             |
| 2.20 (single sheet)     | −0.77| 0.28                 | 1.16           | 0.60           | 1.58             |
| 2.30 (roll)             | −0.81| 1.90                 | 3.10           | 4.26           | 1.66             |
| 2.10 (ribbon)           | −1.08| 0.37                 | 1.39           | 1.08           | 0.70             |
| 2.80 (trefoil)          | −1.12| 0.14                 | 0.24           | 0.43           | 0.03             |
| 3.50 (3-layer(bba) sandwich)| −1.76| 0.06                 | 1.31           | 0.37           | 1.33             |
| 2.60 (sandwich)         | −1.91| 1.24                 | 9.86           | 8.35           | 1.25             |
| 3.70 (box)              | −1.94| 0.02                 | 0.01           | 0.11           | 0.01             |
| 1.50 (alpha/alpha barrel)| −3.88| 0.01                 | 0.25           | 0.39           | 0.02             |
| 2.170 (beta complex)    | −4.81| < 0.01               | 0.40           | 0.49           | 0.22             |
| 2.160 (3 solenoid)      | −4.87| < 0.01               | 0.28           | 0.52           | < 0.01           |

Note: The first column denotes the number of a CATH architecture $CA_i$ and its name. $E_i$ is the enrichment factor determined as log-odds ratio according to Equation (2). A positive value indicates for architecture $CA_i$ an overrepresentation among the $CA_{\text{motifs}}$. The following two columns list the relative frequencies determining $E_i$. $f(\text{CA_motifs})_i$ is the frequency of single-domain proteins from $CA_i$ contributing a motif at the architecture level and $f(\text{CA})_i$ is the frequency of single-domain proteins belonging to $CA_i$. $f(\text{CA_SD})_i$ is the frequency of $CA_i$ among all single-domain proteins and $f(\text{CT_motifs})_i$ is the frequency of $CA_i$ among the single-domain proteins contributing a motif on the topology level. Rows are sorted according to the value of $E_i$. 

As Fragstatt deduced motifs from protein pairs ($P_i$, $P_j$) possessing different CATH architectures, we wanted to know which pairs of architectures were linked, and if some combinations were particularly abundant. We counted how many motifs were detected for each combination of CATH architectures, listed the results in Table S3, and visualized the numbers as a network in Figure 3. Relatively few combinations of architectures constitute the greatest part of all hits: Most abundant are the following five combinations of two CATH architectures: 3.20 (alpha-beta barrel) $\rightarrow$ 3.40 (3-layer(aba) sandwich), 1.10 (orthogonal bundle) $\rightarrow$ 3.40 (3-layer(aba) sandwich), 1.20 (up-down bundle) $\rightarrow$ 1.25 (alpha horseshoe), 1.20 (up-down bundle) $\rightarrow$ 1.10 (orthogonal bundle), and 1.10 (orthogonal bundle) $\rightarrow$ 3.30 (2-layer sandwich). This overrepresentation of few folds has a marked impact on the length distribution of the motifs. A strong bias originates from the comparison of CATH architectures 3.20 $\rightarrow$ 3.40, because the outcome dominates the motifs longer than 40 residues (compare Figure S4).

In sum, five combinations constitute 57% of all motifs. With a frequency of 25%, motifs that occur both in architectures 3.20 (alpha-beta barrel) and 3.40 (3-layer(aba) sandwich) are the most abundant ones and one representant is the $\beta\alpha_2$ motif identified earlier. Additionally, four of the five most frequent combinations belong to the class "mainly alpha"; moreover, symmetrical architectures like 3.40 (alpha-beta barrel) or repetitive architectures like 1.20 (up-down bundle) and 1.25 (alpha horseshoe) are abundant. Of the alpha helical proteins, architecture 1.10 (orthogonal bundle) seems most central, showing strong connections to architectures from all classes; compare Figure 3. The mean number of edges is 8.5, and we assume that the
relatively high number of edges of the architectures 2.40 (beta barrel, 18 edges), 3.10 (roll, 17 edges) and 1.20 (up-down bundle, 15 edges) is most likely due to their simple architecture and the existence of short repeating elements (beta strands and alpha helices) which can be found in many other architectures.

Next, we wanted to know whether there is a correlation of the age of folds and the abundance of CA_motifs. We used previously calculated values that are based on a maximum parsimony calculation and the NCBI taxonomy tree. These ages of SCOP folds lie in the range [0, 1]; folds assigned an age 1.0 have been estimated to have evolved before the divergence of the superkingdoms. We compiled the age distribution of CA_motifs by determining the CATH topology of the related fragments and the correspondence of CATH/SCOP folds by utilizing a Genome3D mapping. We related the age distribution of CA_motifs to the age distribution of our full dataset and determined enrichment factors as log-odds ratios of corresponding frequency values (compare Figure 4A,B). Compared to the CATH single-domain dataset, the estimated age of folds bearing CA_motifs is significantly higher (P-value 1.3E-74, Mann-Whitney U test). As indicated by the log-odds ratios, CA_motifs are overrepresented in ancestral folds and underrepresented in more recent ones. Most interestingly, 61% of the CA_motifs occur in those 52% of all folds that have an age of 1.0, that is, evolved most likely prior to the divergence of the three superkingdoms.

A reason for the overrepresentation of CA_motifs in ancestral folds might be the higher flexibility of these folds in supporting demands imposed by different functions. More than 60% of all folds carry at most two enzymatic functions, but the (βα)_8-barrel and the Rossman fold allow hundreds. Thus, we estimated for the CATH single-domain dataset the designability of each fold, which depends on its stability and compatibility with a large number of sequences by counting the number of related superfamilies sfamfold. Using the age distribution introduced above, we created a histogram of mean values sfamfold; compare Figure 4C. The histogram indicates a strong bias in the fold-specific number of superfamilies and proposes higher flexibility and designability of ancestral folds. The highest sfamfold-value, namely 13.5, was observed for folds that have an age of 1.0.

In summary, these analyses show that the majority of CA_motifs was found in a small number of more ancient CATH architectures. Among these architectures, mainly alpha helical ones are overrepresented; moreover, the combination of CATH architectures 3.20 (alpha-beta barrel) and 3.40 (3-layer(aba) sandwich) contributes 25% to the CA_motifs. On the other hand, the many edges to be seen in Figure 3 indicate that less common motifs were detected in other CATH architectures.
3.4 Structural similarity of the CA_motifs

A critical parameter that affects the number of found motifs, is their structural similarity, which scholars consistently determine by computing the TM-score.\textsuperscript{25,26,34} The TM-score is normalized to the range 0 to 1.0, where 1.0 indicates a perfect match between the two structures. Domains with a TM-score $> 0.5$ can be generally assumed to possess the same fold.\textsuperscript{9,21} However, it was unclear to us whether this cut-off is adequate for the comparison of fragments of sub-domain size. More specifically, we wanted to ensure that the structural similarity of fragments that gave rise to CA_motifs is higher than that one to be expected by chance. Thus, we computed a “null model distribution” consisting of TM-scores resulting from a comparison of unrelated fragments. In order to create this fragment set, residue positions were randomly chosen in PDB entries contributing to CA_motifs. Subsequently, 12,532 fragments were excised with a length distribution following that of CA_motifs (compare Figure S4) and superposed to compute TM-scores.

Figure 5 shows the resulting distribution (named CA_rand) representing the TM-scores under the assumption that the null hypothesis (no structural similarity) is correct. We used CA_rand to fix the TM-score cut-off for a P-value of 0.01, which gave rise to the filter TM-score $\geq 0.55$ for motif selection.

Figure 5 also shows the TM-score distribution CA_all_motifs, which results, if Fragstatt is applied to all of our chosen representatives of CATH topologies without applying a TM-score cut-off for motif selection. The analysis of the two distributions makes clear that the comparison of randomly chosen fragments gives rise to a large number of TM-scores that are lower than those observed in CA_motifs, the mean values are 0.26 and 0.34, respectively. Both an unpaired t-test and a Wilcoxon signed-rank test gave highly significant results with a P-value $<1E-99$.

In order to estimate how the probability of Type I errors depends on the TM-score cutoff, we additionally determined the complementary cumulative distribution function $CCDF(CA_all_motifs)$ and plotted it in Figure 5. As can be seen, the fraction of false positive hits increases drastically for TM-score cut-offs below 0.5; the empirically determined P-values are 0.01, 0.016, and 0.31, if TM-score cut-offs of 0.55 (here), 0.50,\textsuperscript{23} and 0.30\textsuperscript{26} are chosen. In summary, these results
confirm that most of the motifs identified by means of Fragstatt possess a larger TM-score than that expected by chance.

3.5 Clustering results in 134 prototypical CA_motifs

CA_motifs are fragments of sub-domain size that were found by Fragstatt in two proteins with different CATH architectures. However, if the predecessors of the motifs emerged in the pre-LUCA era, it might be that the sequence similarity of modern manifestations of the same motif is too low to deduce homology, even with a highly sensitive HMM-based approach. Thus, it might be that the same or a highly similar motif occurs in several pairs of proteins. This is why we clustered all motifs $M_i$ from CA_motifs to identify prototypical motifs $M^*_i$; for details see Supporting Information.

In summary, 134 prototypical motifs $M^*_i$ were found, which are listed in Table S4. Three motifs ($M^*_1, M^*_2, M^*_3$) occur in more than four CATH architectures. Nine motifs occur in four ($M^*_4 - M^*_12$) architectures, 36 motifs ($M^*_13 - M^*_48$) in three architectures and the rest, summing up to 86 motifs ($M^*_49 - M^*_134$), only occur in two architectures. This means that over 90% of the motifs are present in not more than

![Examples of prototypical CA_motifs](wileyonlinelibrary.com)
three architectures, at the most. The motifs, which were found in a large number of architectures are rich of alpha helices ($M_1$ - $M_9$; compare Figure 6). In contrast, motifs with a more diverse secondary structure (like $M_{10}$) were found in fewer architectures. This was also observed for motifs rich of beta sheets (like $M_{10}$).

The five most abundant CATH architectures among the broadly represented motifs are 1.10 (orthogonal bundle), 3.40 (3-layer(aba) sandwich), 2.40 (beta barrel), 3.30 (2-layer sandwich), and 3.20 (alpha-beta barrel). This observation coincides with the frequencies of CATH architectures in Ca_motifs and substantiates the finding that most of the motifs were found in a small set of CATH architectures.

3.6 Functional characterization of prototypical $M_i$-motifs

In order to characterize the functional role of the 134 prototypical motifs, we opted for the annotations of BioLiP (version Feb 05, 2021, 529 047 entries). We selected this database, because some molecules like ethylene glycol or malonic acid are used as additives to solve protein structures and are annotated as ligands. Thus, not all ligands present in the PDB database are biologically relevant and the curators of BioLiP verify the biological relevance of the reported ligands. The ligands bound by all $M_i$-motifs are listed in Table S5 and named according to the nomenclature of BioLiP.

BioLiP reports nucleotide binding for 11 (8%), and metal or ligand binding for 70 (52%) $M_i$-motifs. 30 (%) of the $M_i$-motifs bind at least two ligands and 64 (48%) motifs lack a reported binding function. The most abundant motif $M_1$, which occurs in 11 CATH architectures, is involved in binding nucleotides and HEM (heme). Motif $M_2$ occurs in four CATH architectures and binds nucleotides, FES (iron-sulfur cluster), and ZN (zinc). Motif $M_3$ binds NAD, NAP, and M7P (7-O-phospho-D-glycerol-alpha-D-manno-heptopyranose). The binding of five ligands is reported for motif $M_{69}$, namely GOL (glycerol), 3HC (3-hydroxybutyryl-coenzyme A), XE (xenon), FE (iron), and MN3 (manganese). Motifs $M_{100}$, binds four ligands, namely nucleotides, III (peptides), TFP (trifluoperazine), and CA (calcium). $M_{114}$ binds 4NB (4-nitrobenzoic acid), FEO (ferrioxoxiron), FE (iron), and ZN (zinc). To a great extent, these findings suggest for $M_i$-motifs a role in the coordination of cofactors, which is in agreement with the function deduced for AncPept motifs.

3.7 Identifying proteins possessing more than one motif

If ancestral motifs have been the building blocks for the evolution of proteins domains, it should be possible to find among the proteins contributing to Ca_motifs some that possess two or more motifs. After the elimination of motifs that correspond to overlapping fragments, we found 135 proteins that contained at least two non-overlapping fragments and all cases were inspected manually. Twenty-two proteins were discarded due to the low quality of the superpositions of the motifs (helix only or too short) leaving 113 cases, which belonged to 15 different CATH topologies; compare Figure 7. 90% of the cases belong to the three CATH topologies 3.20.20 (TIM barrel, 53%), 3.40.50 (Rossmann fold, 21%), and 1.25.40 (serine threonine protein phosphatase 5, 16%). We manually classified the 113 cases according to the three criteria repetitive, overlapping and fragmented; compare Figure S6. Repetitive indicates that the same motif was found multiple times at different positions in the same protein chain. Overlapping indicates that the positions of the motifs overlap and fragmented indicates that inside a larger motif smaller sub-fragments occurred that were interpreted as individual motif regions.

One hundred six of the 113 cases were repetitive and mainly based on motifs detected between TIM, that is, (ix)x$_3$-barrels and the Rossmann fold. Among the four cases that were non-repetitive and non-fragmented was the in silico designed protein OR258 (PDB-ID 4J29_A), which was discarded. The remaining three cases will be detailed in the following.

The transcriptional regulator PlcR (PDB-ID 2QFC_A) contains three distinct and separated motifs; compare Figure 8A. This protein belongs to the CATH superfamily 1.25.40.10 (tetratricopeptide repeat domain, TPR) and all superfamily members share the repetition of TPR motifs. PlcR contains two TPR motifs, which were also found by Fragstatt in (a) two DNA-binding proteins (PDB-ID 5K9B_B and 3EUS_B) from the CATH superfamily 1.10.260.40 (lambda repressor-like DNA-binding domains), (b) a transcription related protein (PDB-ID 3D3B_A), and (c) a lipid transport related protein (PDB-ID 2RK_L) from the CATH superfamily 1.10.940.10 (NusB-like) and the CATH superfamily 1.20.5.420 (immunoglobulin FC, subunit C), respectively. The third motif links PlcR to a mitochondrial protein (PDB-ID 1OM2_A), which belongs to the superfamily 1.20.960.10 (mitochondrial outer membrane translocase complex, subunit Tom20 domain). In PlcR, a long $\alpha$-helix interconnects motifs 1 and 2 with motif 3.

![FIGURE 7 CATH topologies of multi-motif proteins. 90% of the cases belong to the three CATH topologies 3.20.20 (TIM barrel, 53%), 3.40.50 (Rossmann fold, 21%) and 1.25.40 (serine threonine protein phosphatase 5, 16%). The histogram lists absolute numbers of multi-motif proteins [Color figure can be viewed at wileyonlinelibrary.com]](image-url)
Proteins possessing multiple motifs. A, The two repetitive TPR-like motifs of PlcR (PDB-ID 2QFC_A) are shown in green and orange. The manifestations of a third, distinct motif is shown in red. Multiple motifs found in the two homologs B, GmhA (PDB-ID 2XBL_B) and C, diaA (PDB-ID 2YVA_A). For both proteins the same two motifs were detected: A beta-alpha-beta motif (green) and an alpha-beta motif (red) [Color figure can be viewed at wileyonlinelibrary.com]

The isomerase GmhA (PDB-ID 2XBL_B) and the initiator-associating protein diaA (PDB-ID 2YVA_A) both possess a Rossmann fold and belong to the CATH superfamily 3.40.50.10 (α-6-phosphatase isomerase like protein; domain 1). In both cases (compare Figure 8B,C) a beta-alpha-beta motif is detected, linking GmhA and diaA to two TIM barrel proteins (PDB-ID 1JCN_A and PDB-ID 1I4N_A). Both TIM barrel proteins belong to the CATH superfamily 4.10.520.10 (IHF-like DNA-binding proteins). The second motif in the GmhA and diaA, an alpha-beta motif, links them to a DNA-binding protein (PDB-ID 2NDP_A), which belongs to the CATH superfamily 4.10.520.10 (IHF-like DNA-binding proteins).

In summary, by analyzing 309 proteins that contributed to CA_motifs, we found only two cases of folds, namely PlcR plus GmhA and diaA (the latter two are members of the same superfamily) that possess two clearly distinct motifs. In combination with our other results, this finding strongly argues against the frequent recombination of sub-domain sized motifs in the evolution of protein domains.

4 | DISCUSSION

4.1 | Several approaches provide no convincing evidence for the combination of motifs in non-repetitive folds

Since Dayhoff and Eck postulated the hypothesis that modern proteins emerged from the recombination of smaller recurring fragments,17 it has been shown that repetition plays a fundamental role in evolution.49 Gene duplication followed by fusion and diversification is a dominant evolutionary mechanism.50,51 A prototypical example is the βββ-barrel fold that most likely evolved via a two-step evolutionary pathway from a βαβ-motif constituting a quarter-barrel.52 Similar cases of a conserved internal symmetry can be observed in various protein folds, for example, beta-propellers,41 β-trefoils,53 or TPR repeat folds,54 which further substantiates the idea that protein domains can emerge from relatively small protein fragments. Based on these observations, the idea of proteins built from smaller modules has been developed; for a review see Söding & Lupas55.

AncPept,23 the bridging themes,27 Fuzzle,26 and our CA_motifs specify four sets of putatively ancestral protein motifs. AncPept, the bridging themes, and Fuzzle recruit motifs from pairwise alignments of HMMs representing SCOP domains. CA_motifs are based on HMM cascades and the CATH classification. We opted for this approach to improve detection sensitivity and to explore the effect of alternative structural classification schemes on the detection of putative ancestral protein motifs.

The AncPept motifs have been found in 118 (10%) of the 1194 considered SCOP folds.23 We concluded that this low abundance of primordial motifs is no convincing evidence for the existence of universally used building blocks. This insight has driven the sensitivity-aware design of Fragstatt and prompted us to utilize HMM cascades and low probability HHblits alignments to detect more motifs. We ensured that false positive alignments have no effect by deducing fragments via a traceback; compare the content of the matrix H shown in Figure S1. Despite these sensitivity-increasing efforts, the CA_motifs occur in not more than 245 (18%) of the 1391 considered CATH topologies, which is less than twice the coverage of the AncPept motifs and the utilization of all motifs is far from universal.

A critical parameter with a crucial influence on the number of proposed motifs is the cut-off chosen for the TM-score. For the determination of AncPept and CA_motifs, the cut-off values were 0.5 and 0.55, respectively. In the set of the bridging themes, 30% have a TM-score < 0.3 and only 25% a score > 0.5.27 For the identification of Fuzzle motifs, a TM-score threshold of 0.3 in combination with an RMSD threshold of 3 Å and no upper or lower bound for the length of the motifs has been used.26 As a consequence, the average length of Fuzzle motifs is 64 residues and the motifs were found in 519 (43%) of the 1221 considered SCOP folds. As Figure 5 demonstrates, a TM-score < 0.3 is close to the maximum of randomly sampled TM-scores, which is 0.2. Moreover, this cut-off corresponds to a significantly increased false positive rate indicated by a P-value of 0.31. Thus, these structural correspondences are from a twilight zone. Nevertheless, it is remarkable that motifs from the largest set, namely Fuzzle, have been detected in not more than half of the SCOP folds.

An indicator for the universality of a motif is the number of folds or topologies in which it was detected. The most universal AncPept motif has been found in 1423 and the most universal prototypical CA_motif in 13 folds or topologies. However, the majority of CA_motifs occur in only two, three or four folds or topologies. For motif
4.2 The usage of motifs seems restricted to early phases of fold evolution

F. Jacob’s phrases “Nature is a tinkerer, not an inventor” and “the probability that a functional protein would appear de novo by a random association of amino acids is practically zero” emphasize the role of gene duplication and disregard the possibility that new genes might arise de novo. However, the limited spread of motifs in a small number of folds does not provide convincing evidence for the modular structure of protein domains and the frequent reuse of motifs during the evolution of the full fold space.

On the other hand, former results and those presented here are compatible with F. Jacob’s idea and suggest that the reuse of short motifs played a role in the origin of protein folds that date back to the formation of RNA-motif complexes in early phases of protein genesis. A mapping of the five most ancient SCOP folds onto the CATH database showed that they belong to the CATH topologies Rossmann fold, TIM barrel, Trp operon repressor and alpha-beta plaits. These topologies are all observed among the CA motifs and similarly among the AncPept and Fuzzle motifs.

The earliest phases of protein evolution must have involved the ab initio invention of new folds, which were flexible enough to support a primordial form of life. The age-specific distribution of $\text{sham}_{\text{fold}}$ values, which reflects designability, supports this notion; compare Figure 4C. Generally, more than 60% of folds carry out one or two enzymatic functions, but some folds, like the βα8-barrel or the Rossmann fold, catalyze hundreds of reactions. Although these folds are considered ancestral, they contain the most reliably identified motifs. Thus, it is likely that the combination of motifs—often in a repetitive manner—was relevant for the creation of primordial proteins, perhaps by a combination of elementary functional loops. It is likely that structure and sequence similarity faded away for those copies of motifs that were less relevant for protein function or stability. Additionally, it might be that due to the evolution of a more sophisticated DNA replication and protein synthesis machinery, the combination of motifs became less likely for the genesis of more recent folds. Thus, in contrast to F. Jacob’s intuition, the de novo gene birth seems to continuously contribute to the recruitment of protein-coding sequences as evidenced by recent laboratory experiments.

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CONFLICT OF INTERESTS

The authors declare no potential conflict of interest.

DATA AVAILABILITY STATEMENT

The datasets supporting the conclusions of this article are available in the https://github.com/merkllab/motif_reuse repository. Program code is available from the corresponding author on reasonable request.

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**Supporting Information**

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