Effect of Sequence Padding on the Performance of Protein-Based Deep Learning Models

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Abstract

Background: The use of raw amino acid sequences as input for protein-based deep learning models has gained popularity in recent years. This scheme obliges to manage proteins with different lengths, while deep learning models require same-shape input. To accomplish this, zeros are usually added to each sequence up to a established common length in a process called zero-padding. However, the effect of different padding strategies on model performance and data structure is yet unknown.

Results: We analysed the impact of different ways of padding the amino acid sequences in a hierarchical Enzyme Commission number prediction problem. Our results show that padding has an effect on model performance even when there are convolutional layers implied. We propose and implement four novel types of padding the amino acid sequences.

Conclusions: The present study highlights the relevance of the step of padding the one-hot encoded amino acid sequences when building deep learning-based models for Enzyme Commission number prediction. The fact that this has an effect on model performance should raise awareness on the need of justifying the details of this step on future works. The code of this analysis is available at https://github.com/b2slab/padding_benchmark.

Keywords: padding; amino acid encoding; protein encoding; deep learning

Background

Since the breakthrough of deep learning (DL) [1], deep neural networks are being successfully applied in computational biology [2, 3]. This is due to their capacity for automatically extracting meaningful features from raw data [4]. Specifically, DL is useful in the context of biological sequences, such as proteins or RNA, because it can learn directly from the sequence and hence, capture nonlinear dependencies and interaction effects. Some examples of applications of DL on biological sequences include prediction of specificities of DNA and RNA binding proteins [5], DNA function quantification [6], de novo peptide design [7], detection of conserved DNA fragments [8], prediction of protein associated GO terms [9] or quantification of the impact of genetic variation on gene regulatory mechanisms [3]. The specific DL architectures able to leverage the inner structure of sequential biological data are Convolutional Neural Networks (CNN) and Recurrent Neural Networks (RNN). CNNs imply translational invariance [10] and can be used to find relevant patterns with biological meaning [8, 5, 11, 12]. For their part, bidirectional RNNs (and the derived Long Short-Term Memory and Gated Recurrent Units) are appropriate for modelling biological sequences since they are suited for data with a sequential but
non-causal structure, variable length, and long-range dependencies [13, 14, 15, 16]. Both architectures are usually combined, as in DEEPre [17], where a CNN-RNN model performs a hierarchical classification of enzymes.

Proteins are long linear sequences constituted by amino acid residues attached covalently. These amino acid residues are represented by letters that cannot be directly processed by the mathematical operations used by DL models. Choosing how to digitally encode amino acids is a crucial step in this context, since it can affect to the overall performance of the models [18]. A comprehensive review and assessment on different amino acid encoding methods [19] shows that position specific scoring matrix (PSSM), an evolution-based position dependent methodology, achieves the best performance on protein secondary structure prediction and protein fold recognition tasks. However, this type of encoding is very consuming computationally [20] and its applicability is limited to proteins with known homologous sequences [19], which could highly decrease the generalisation capabilities of the predictor for non evolutionary related proteins. Traditionally, proteins have also been encoded into feature vectors [21, 22]. These encoding features are generally aggregative and not bijective, such as signatures, physicochemical properties or amino acid composition. From the aggregative features, the original sequence cannot be recovered, resulting in the loss of protein information.

The analogy between text and proteins, understood as sequences of characters with a meaning, motivates the application of Natural Language Processing (NLP) techniques to amino acid sequences. Along these lines, machine-learning derived embeddings [23, 24, 25, 26] and one-hot encoding [14, 9, 27, 12, 17, 7] have become very popular. Specifically, the latter method has been widely used in protein-based DL models since neural networks are able to extract features from raw data. An schematic explanation of one-hot encoding is shown in Figure 1B. Every amino acid of a protein sequence is represented by a binary vector of length \( n + 1 \), being \( n \) the number of different amino acids and placeholders. In this vector, all but the corresponding entry for that amino acid is set to zero. As a result, a protein of length \( L \) is represented by a \( (n + 1) \times L \) binary matrix.

The main problem of one-hot encoding is that each protein has a different length, while all the input vectors should be of the same size to be fed into the model. To overcome this issue, sequence padding and truncation are usually applied [7, 12, 14, 9]. This means establishing a common length for all the proteins and then, truncating longer proteins to that length or filling shorter proteins with an “artificial” character up until that length (see Figure 1A). This process of completing a sequence is called padding and the character used for filling could be any that is not used on the sequences themselves, being the zero character (“0”) usually chosen. Padding zeros can be added at any point of the sequence, for example at the N- and C- terminals of the sequences [28]. In practice they are usually added at the end [7, 14]. However, details on the concrete steps when padding the sequences are often omitted as they are deemed of low importance for the results of the study [12, 9, 27, 17]. Even when this information is given, there is no proper justification on the padding choice [7, 14, 28]. Partly, this is due to the lack of exhaustive studies on the effect of padding the sequences. Up to our knowledge, the work of Reddy et al, 2019 [29] is the only study on the effect of padding sequences in deep
In this paper, we report a systematic analysis on how different types of padding affect to protein-based DL models performance. We evaluate this effect on three different DL architectures: only feed-forward neural networks (only\_denses), feed-forward neural networks coupled with a convolutional layer (1\_conv) and feed-forward neural networks coupled with a stack of convolutional layers (stack\_conv). We also introduce four novel padding types (mid-, strf-, rnd-, and zoom-) and we classify them along with the known types (pre-, post- and ext-) into dense and sparse paddings. Dense paddings are those keeping zeros together in a block (pre- at the beginning, post- at the end, mid- in the middle and ext- at both ends), while in sparse paddings, zeros are interspersed on the sequence (randomly the case of rnd- and uniformly for strf-) or amino acids are duplicated (zoom-). Finally we quantify the behavior for each type of padding. The task chosen for this study is a hierarchical classification of enzymes with two levels: the first is a binary classification of proteins into enzymes/non-enzymes, and the second is a multi-label prediction of the enzyme type.

## Results

### Performance metrics

A summary of the F1-score (macro average) on test for each architecture, each type of padding and each task is shown in Table 1. Since the trends observed for accuracy were analogous, we will focus on F1-score, but homologous tables for accuracy on test for both tasks and Area under the receiver operating characteristic curve (which will be referred as AUC from now) on test for task 1 can be found in Tables S1-S2.

### Table 1

| Padding type | Task 1 F1-score | Task 2 F1-score |
|--------------|----------------|----------------|
|              | 1\_conv        | only\_denses   | stack\_conv  |
| aug          | 0.756 ± 0.041  | 0.900 ± 0.011 | 0.790 ± 0.022 |
| ext          | 0.842 ± 0.022  | 0.896 ± 0.013 | 0.875 ± 0.010 |
| mid          | 0.868 ± 0.016  | 0.911 ± 0.018 | 0.874 ± 0.026 |
| post         | 0.873 ± 0.010  | 0.900 ± 0.014 | 0.879 ± 0.024 |
| pre          | 0.858 ± 0.011  | 0.899 ± 0.013 | 0.863 ± 0.028 |
| rnd          | 0.786 ± 0.014  | 0.896 ± 0.025 | 0.812 ± 0.011 |
| strf         | 0.867 ± 0.008  | 0.930 ± 0.011 | 0.851 ± 0.010 |
| zoom         | 0.868 ± 0.005  | 0.893 ± 0.021 | 0.862 ± 0.014 |
|              | 0.536 ± 0.025  | 0.531 ± 0.022 | 0.504 ± 0.026 |
|              | 0.554 ± 0.022  | 0.543 ± 0.034 | 0.540 ± 0.023 |
|              | 0.558 ± 0.021  | 0.542 ± 0.027 | 0.557 ± 0.024 |
|              | 0.550 ± 0.025  | 0.509 ± 0.075 | 0.554 ± 0.024 |
|              | 0.541 ± 0.020  | 0.541 ± 0.030 | 0.527 ± 0.028 |
|              | 0.452 ± 0.026  | 0.448 ± 0.034 | 0.455 ± 0.020 |
|              | 0.550 ± 0.024  | 0.548 ± 0.026 | 0.547 ± 0.024 |
|              | 0.543 ± 0.019  | 0.456 ± 0.063 | 0.515 ± 0.026 |

Table 1: Results are reported for all the different types of padding for both task 1 and task 2 in each one of the tested architectures. Mean ± standard deviation of the 10 folds.

learning models. It is applied on a NLP sentiment analysis task and only pre- and post- padding types are tested. Since it is a different application domain and the options they test are limited, a more comprehensive study for the case of biological sequences is needed. Likewise, alternative types of padding to those usually implemented (zeros at the end of the sequence, at the beginning or both) have not been yet explored.
of the Supporting Information. Figure 2 shows the F1-score on test for each type of padding in each of the tested architectures, both for task 1 and task 2. The same figure but for accuracy can be found in Figure S1 of the Supporting Information.

According to Table 1 and Figure 2, although the different architectures seem to have similar F1-score values, only denses architecture is the one that achieves the best performance for task 1 (paired Wilcoxon test, two-sided, \( p = 9 \times 10^{-15} \) vs 1_conv and \( p = 4 \times 10^{-13} \) vs stack_conv). Regarding task 2, we can see at Figure 2 that the trend is not as clear as in task 1. 1_conv has the best performance (paired Wilcoxon test, two-sided, \( p = 9 \times 10^{-4} \) vs only denses, \( p = 2 \times 10^{-5} \) vs stack_conv), while there are no statistical differences between stack_conv and only denses.

**Effect on input space**

We studied the distribution of the activations of the 1D Convolutional layer for the 1_conv model to analyse the effect of the padding type in the input space by means of a Principal Components Analysis (PCA). Figure 3 displays the density plot showing the principal components (PC) 1 and 2 of the activations from the 1D convolutional layer of the 1_conv architecture for each type of padding on each fold in task 2. In Figure S2 of the Supporting Information, the same representation for task 1 is shown. Focusing on Figure 3 for task 2, the distribution of the activations is very similar for dense types of padding (ext-, mid-, post- and pre-). These dense activations are grouped in two clusters separated along the PC1. Sparse paddings (rnd-, strf-, zoom-) activations have a distribution very different to that from dense paddings. In this case, activation points are condensed in one area, although each one of these types of padding has its own structure. Regarding enzyme types, according to the structure of the distributions, there seems to be two different groups: enzyme types 2,3,4 are very similar between them and in turn, different to types 1, 5, 6, 7. Table S9 of the Supporting Information quantifies the effect of the enzyme and the padding types on the PC1 of the activations using a linear model. All the terms of the model are significant.

**Explanatory models**

We used linear models to further explain the performance metrics and the effect of different variables (padding type, enzyme type, architecture) to the DL models behaviour. These explanatory models were also used to address specific questions regarding the effect of padding.

The full additive linear model in Equation 1 describes the F1-score values on test and it is shown on the Table S3 of the Supporting Information. It shows that some types of padding have indeed an effect on models performance, both for task 1 and 2. For example, for all the architectures (since it is an additive model) in task 1, aug-, pre-, ext- and rnd- have worse performance when compared to the reference padding type (post-) (\( p\)-value < 0.05). In the same setting but for task 2, rnd- and zoom have significantly worse performance than post- (\( p\)-value < 0.01).

Figure 4 and Tables S4-S8 gather the answers to our specific questions on the effect of padding on the different architectures and enzyme types. The colour represents in each case the difference between each category and the reference category of that factor. The sign of the corresponding estimate is represented if that difference is statistically significant. The constant term of a model (Intercept) shows the prediction when all the categorical variables have their reference values.
A. Does padding position affect performance?

Figure 4 A and Table S6 show that the classification performance of the baseline model (only_denses) for task 1 is the same for all the types of dense padding, except for strf-padding, which is better than post- (predicted F1-score of 0.926 vs 0.896). In the same way for task 2, only strf- significantly outperforms post-padding (0.472 vs 0.432), while zoom- has a worse performance (0.379).

B. What is the effect of switching between dense paddings?

Figure 4 B and Table S5 for task 1 show that for stack_conv, there are no differences in performance between dense paddings. Regarding task 2, only pre-padding is significantly worse than post-padding (0.457 vs 0.484) for stack_conv.

C. What is the effect of changing from the standard dense padding to sparse padding?

Figure 4 C and Table S6 show that for stack_conv in both tasks, sparse paddings have significantly worse performances than dense paddings. For task 1, post- significantly outperforms rnd-, strf- and zoom-padding (predicted F1-scores of 0.868 vs 0.801, 0.840 and 0.851, respectively). For task 2, post- also outperforms rnd- and zoom-padding (0.498 vs 0.400 and 0.460, respectively).

D. Is an ensemble of paddings beneficial?

We tested for the three architectures if data augmentation regarding padding (aug-padding) improved the performance when compared to post-padding (representing the dense paddings) and strf-padding (representing the sparse paddings). To quantify the changes, we used aug-padding as reference level in padding type. Figure 4 D and Table S7 show that both post- and strf- significantly outperform aug-padding both for task 1 and task 2.

For task 1, with the baseline architecture (stack_conv) aug-padding gets the worst predicted F1-score (0.786), while it is 0.875 and 0.847 for post- and strf- respectively. For the reference padding type (aug-), the stack_conv architecture performs worse than only_denses (0.785 vs 0.895) but better than 1_conv (0.785 vs 0.752). Interactions show that both strf- and post- have a more positive effect on performance with respect to the baseline (aug-) for 1_conv (0.863 and 0.868 vs 0.752 from aug-) than for stack_conv (0.847 and 0.875 vs 0.786). On the contrary, changing from aug- to strf- and post- have less performance improvement for only_denses (0.895 from aug- vs 0.926 and 0.896) than for stack_conv (0.786 vs 0.847 and 0.875), although even so only_denses still outperforms stack_conv.

Regarding task 2, with the baseline architecture aug-padding also gets the worst performance when compared to post- and strf-padding (0.421 vs 0.471 and 0.464). For baseline padding type, both 1_conv and only_denses significantly outperform stack_conv (0.453 and 0.449 vs 0.421). Interactions show that 1_conv reacts the same way to changes of padding type than stack_conv. But in the case of only_denses, changing from aug- to post-padding (0.449 vs 0.426) has less performance improvement than for stack_conv (0.421 vs 0.471), causing aug- to outperform post-.
E. Does the effect of padding type on performance depends on the enzyme type?

We checked for all the types of architectures and for dense and sparse paddings (represented by post- and strf-, respectively) the effect of enzyme type on model performance (only for task 2). Results ((Figure 4 E and Table S8) show that for both padding types, the performance for enzymes with the first EC number digit 2 (0.619), 3 (0.555), 5 (0.598), 6 (0.532) and 7 (0.597) is better than for digits 1 (0.457) and 4 (0.454). Interactions are not significant, meaning this trend applies to all the architectures. This is consistent with the results of the previous questions, where enzyme types 4, 1 and sometimes 6 are shown to decrease performance.

Discussion

Although it is not the aim of this paper to study differences on performance between architectures, in general terms only_denses has shown to achieve the best performance for task 1 while both convolutional architectures work better for task 2 (see Tables 1, S3-S6 and S8, and Figures 2-4). Quantitatively, for the full additive model (eq. 1) in task 1, only_denses get a predicted F1-score of 0.916 vs 0.864 of stack_conv and 0.853 of 1_conv for the baseline post-padding type. Using the same model for task 2, predicted F1-score of only_denses is 0.444 versus 0.454 of stack_conv and 0.464 of 1_conv. The reason why only_denses is the best architecture for task 1 could be that the task of classifying amino acid sequences between enzymes and non-enzymes is more related to the presence/absence or count of certain amino acids than to their position within the sequence. In other words, if we could consider them to be amino acid sets instead of sequences as it happens in other fields [30]. On the contrary, classifying enzymes into their types it is a more complex task that might imply amino acid patterns and position information, thus a convolutional architecture is a better choice.

Along the same lines, we have seen that task 1 has a better performance than task 2 for all the architectures (Table 1 and Figure 2). F1-score ranges from 0.756±0.041 to 0.930±0.011 for task 1, while for task 2 it is comprised between 0.448±0.034 and 0.558±0.021. Task 1 results are similar to those obtained by DEEPRe [17] for their equivalent Level 0 prediction, but results obtained for task 2 are worse than their report for Level 1. This was expected, since we use the same architecture for a simple binary classification and for a multi-class classification problem. A more complex, optimized model may improve the performance for the first digit prediction problem, but this was out of the scope of this study. We chose the architectures of both tasks to be as comparable as possible.

We have confirmed that padding type has an effect on model performance (see Tables S3-S8 and 1, Figure 4). In Figures 3 and S2 we could see that indeed, the models see differences for each type of padding in their input space.

In general, there are no differences between dense paddings (see Figure 4 A and B and Tables S4-S5), neither for convolutional or only_denses architectures. This applies for both task 1 and task 2, although for the latter pre-padding underperforms the rest of dense paddings (predicted F1-score 0.457 from pre- vs 0.484 from post-). Therefore, dense paddings are interchangeable for fully feed-forward and dense architectures and we could stick to the default option (post-padding).

There are differences between sparse paddings. For the baseline model (only_denses) in both tasks (Figure 4 A and Table S4), strf_padding has shown to outperform
the rest of the paddings: for task 1, strf- has a predicted F1-score of 0.926 vs 0.896 from post-; in task 2, strf- has an estimate of 0.472 while for post- it is 0.432; macro-average for the F1-score on test is 0.930 ± 0.011 for task 1 and 0.548 ± 0.034 for task 2 (Table 1). This might be because feed-forward neural networks are position-sensitive and moving a block of zeros along the sequence (as in different types of dense padding) can alter the way the networks process them. Strf- does not comprise a block of zeros, but they are spread uniformly along the sequence. This distribution seems to compensate this position sensitivity, by aligning certain relative positions of the protein where the model might be detecting abundance changes.

On the contrary, this improvement of performance caused by strf-padding does not apply for stack_conv architecture (Figure 4 C and Table S6). In this case, all sparse paddings perform worse than the baseline post-padding (except for strf- in task 2): for task 1, the predicted F1-score of post-padding is 0.868 vs 0.840, 0.801 and 0.851 from strf-, rnd- and zoom- respectively; for task 2, predicted F1-score is 0.498 for post- vs 0.400 and 0.459 for rnd- and zoom-. Thus, convolutional models works better with dense paddings than with sparse ones.

The differences in activations of the convolutional layers in Figure 3 further support the classification of paddings into dense and sparse and are in line with the results we have just reported. The activations for the dense paddings showed to be very similar between them. This is expected due to the translational invariance of the convolutional layers [10]: if zeros are kept together they should be processed in the same way by the convolutional layers, no matter where they are located. In turn, the activations of dense paddings are very different from the sparse ones (Figure 3). Sparse paddings have also a similar structure, where the activations are condensed in only one centered group.

We have also tested if data augmentation regarding padding (i.e. artificially increasing the size of a dataset by representing one protein by different possibilities of the padded one-hot encoded amino acid sequence) improved model performance as in image deep learning models [31]. Our results (Figure 4 D and Table S7 of the Supporting Information) have shown that aug-padding underperforms dense and sparse paddings both for fully dense and convolutional architectures for both tasks: for stack_conv task 1 aug- achieves a predicted F1-score of 0.786 vs 0.875 and 0.847 from post- and strf-, respectively; for task 2, 0.421 from aug- vs 0.470 from post- and 0.464 from strf-. In Figure 2 it also shows to have the worst performance in both tasks for convolutional architectures. Hence, an ensemble of mixed dense and sparse paddings does not improve the performance of the models in this case. Augmented data using only sparse paddings or only dense paddings might work better, because then sequences would be in similar activation spaces.

We observed that models underperformed in enzyme types 1 (oxidoreductases) and 4 (lyases) (Figure 4 and Table S8, the predicted F1-scores for enzymes 2, 3, 5, 6 and 7 are 0.619, 0.555, 0.598, 0.532 and 0.597, respectively, while for 1 and 4 it is 0.457 and 0.454). It does not seem to be related to the number of samples (Figure S3 of the Supporting Information), to sequence length (see Figure S4 of the Supporting Information) or to the distribution of the activations (Figure 3). Therefore, we assume that this is caused because these enzyme types are inherently more difficult
to classify, as it happens in [32]. EC number prediction can be challenging in some cases due to divergent evolution (two enzymes with a completely different EC may actually be very similar in sequence) [33] and parallel evolution of enzyme function (two completely unrelated enzymes catalyse the same reaction and thus, share EC number) [34].

In Figure 3 there also seems to be two groups of enzymes according to the distribution of the activations: 1, 5, 6 and 7 vs 2, 3 and 4. This could be partly related to the sequence length: Figure S4 of the Supporting Information show that enzyme types 2, 3 and 4 are shorter than 1, 5, 6 and 7 ($p = 9e^{-54}$ for Mann-Whitney-Wilcoxon test for independent samples, two-sided); moreover, these differences are not so visible for zoom-padding, for which models cannot count zeros. On the other hand, table S9 of the Supporting Information reports negative coefficients for enzyme types 2, 3 and 4, and positive coefficients for enzymes 5, 6 and 7 (enzyme type 1 is the reference) in the explanatory linear model for PC1, which further supports this grouping.

The results of this study have been obtained for amino acid sequences. It would be needed as a future work to check if this effect of padding on model performance can be translated to other biological sequences that are also one-hot encoded and padded, such as RNA [35, 36] and miRNA [37] or DNA sequences [6].

**Conclusion**

The effect of padding amino acid sequences when they are one-hot encoded had not been comprehensively addressed in the literature yet. The lack of this analysis has caused numerous studies to disregard this step, most of the times taking the “default” option and in some cases, even omitting the details around it. In this paper, we have shown that padding position has an effect on model performance.

We have tested seven types of padding using three different deep learning architectures in a hierarchical enzyme classification problem. It is the first study analysing the impact that padding one-hot encoded amino acid sequences has in the performance of the studied task.

Our results show that padding the amino acid sequence has an effect on the performance of the model. Therefore, more attention should be given to this often omitted step of data pre-processing when building deep learning models for one-hot encoded proteins.

We propose and analyze novel ways of padding proteins when one-hot encoding them for machine learning models (strf-, zoom-, rnd-, mid-). Up until our knowledge, these types neither have been previously seen in the literature nor is their implementation publicly available. We provide the code for their application (https://github.com/b2slab/padding_benchmark), since we have shown that some of them could be more suited for their specific task or architecture.

Our results show that there are no differences between dense paddings. Thus, we can stick to the traditional post-padding, which has proved to outperform the other padding types for convolutional architectures. Regarding sparse paddings, our newly proposed strf-padding has shown to be the best choice for fully feed-forward neural networks, outperforming both dense paddings and the other types of sparse paddings. Lastly, data augmentation regarding the padding (aug-padding) does not
Table 2 Distribution is shown according to the enzyme type, which is determined by the first digit of the EC number. Entries without EC number are considered as non-enzymes.

| Enzyme type | Non-enzymes | Total enzymes |
|-------------|-------------|---------------|
| 1           | 11187       | 10872         |
| 2           | 3843        |               |
| 3           | 2123        |               |
| 4           | 1281        |               |
| 5           | 603         |               |
| 6           | 1715        |               |
| 7           | 120         |               |

improve performance. In contrast, it seems to add noise that causes performance to decrease.

This analysis has been applied to the specific task of EC number prediction. Although we cannot extend these results to other tasks or other deep learning architectures, this is an starting point that highlights the need to stop neglecting the padding step when one-hot encoding amino acid sequences, since we have shown that it has an effect on model performance.

### Material and methods

#### Material

Different types of padding were evaluated in the UniprotKB/Swiss-Prot database [38] (version 2019_05) protein entries for taxonomy Archaea. For computational reasons we established an upper threshold of 1000 amino acids for sequence length, leaving 19,385 proteins for training the models (more than 99% of the original entries). For the enzyme classification task performed for the padding analysis, Enzyme Commission number (EC number) annotation was used. EC number is a numeric classification schema for enzymes related to the chemical reactions they catalyze. Each EC number is constituted by 4 numbers separated by dots, being each one a progressively more specific classification. We only used the first digit of the EC number, which refers to the class of enzyme (1: oxidoreductases, 2: transferases, 3: hydrolases, 4: lyases, 5: isomerases, 6: ligases and 7: translocases) and considered the entries without EC number annotation as non-enzymes. 214 entries with more than one EC number were expanded as additional samples, having a total of 19,599 samples. Table shows the enzyme type distribution of the dataset and Figure S3 of Supporting Information represents this distribution. Data was divided 70/15/15% in training, validation and test sets. The training set was used to fit the model, the validation set was used to evaluate the model fit in each epoch and tune hyperparameters accordingly, and the test set was used to externally evaluate the final model fit. To check the consistency of the results, this splitting was randomly performed 10 times, so each model was trained and tested in each one of these data partitions.

#### Amino acids encoding and protein padding

Amino acids were represented by one-hot encoding. Seven different padding types were applied to those sequences shorter than 1000 amino acids (see Figure 5): (I) **post-padding**, adding zeros at the end of the sequences; (II) **pre-padding**, adding zeros at the beginning of the sequence; (III) **mid-padding** (middle), adding the
zeros in the middle of the sequence; (IV) strf-padding (stratified), distributing the zeros uniformly across the sequences; (V) ext-padding (extreme), adding zeros at both ends of the sequence in a balanced way (half of the padding pre- and half of the padding post-); (VI) rnd-padding (random), adding zeros at random positions of the sequence; (VII) zoom-padding, similar to stratified padding but instead of zeros, contiguous amino acids are repeated; this is the only padding type that "modifies" the sequence length. Additionally, (VIII) aug-padding (augmented) will assess the use of data augmentation regarding padding: each sequence will be represented by the seven different padding strategies.

We divided the types of padding in two groups: (1) dense paddings, those strategies that keep the sequence to a great extent, i.e. post-padding, pre-padding, ext-padding, mid-padding and (2) sparse paddings, which comprises those types of padding which repeatedly modify the structure of the sequence by inserting elements in between: strf-padding, rnd-padding and zoom-padding.

Classification task: hierarchical models
We tackled the enzyme classification task as a hierarchical problem with a level-by-level prediction strategy, as in [17] (see Figure 6), although we only approached the first two levels of the structure. This decision was taken due to the data imbalance (see Figure S3 on the Supporting Information and Table ) between non-enzymes and the less populated enzyme classes (e.g. class 7). We built two prediction models. Firstly, a binary classification model that, given a sequence, predicts if it is an enzyme or not. From now on, it will be referred as task 1. Secondly, a multilabel classification model with seven outputs that, given a sequence classified as enzyme by the first model, predicts the class of the enzyme (the first digit of the EC number). This will be referred as task 2.

Models architecture
We analysed the padding effect on three DL architectures: 1. a model with only feed-forward neural networks (it will be referred as only_denses), 2. a model with feed-forward neural networks and one 1D convolutional layer (1_conv) and 3. a model with feed-forward neural networks and five 1D convolutional layers stacked in parallel (stack_conv). The schematic representation of the four models can be found in Figures S5 and S6. CNNs from the second and third model are aimed to detect meaningful patterns in the amino acid sequence. In all cases, dropout is used to prevent overfitting [39]. The only_denses model was considered as baseline or reference model, the 1_conv model was chosen to study the effect of adding a convolution to the model and stack_conv was taken to check the effect on a convolutional architecture of relative complexity [14, 27]. Further details of the models and the corresponding hyperparameters can be found on the Supporting Information.

Implementation
Models were trained with an Adam optimizer [40] for 200 epochs, with a batch size of 54 (learning rate = 1E-4, $\beta_1 = 0.9$, $\beta_2 = 0.999$). Models were implemented in Python (Keras [41] 2.2.4 using as backend TensorFlow [42] 1.8.0) and run on the GPU NVIDIA TITAN Xp and NVIDIA GeForce GTX 1070.
Performance metrics
The final model was that those weights corresponded to the epoch with maximum validation accuracy in each case. We tested each selected model on the corresponding test set of that data partition. For evaluating and comparing the performance of the different padding types, accuracy, F1-score for each label and macro F1-score (computed by calculating metrics for each label and getting their unweighted mean) on the test set were used. AUC was also computed for task 1 since it is a binary classification problem. AUC was computed from raw predicted probabilities, while F1-score and accuracy were calculated from the binarized predictions at threshold 0.5. To statistically compare these metrics between architectures and types of padding, nonparametric two-sided Wilcoxon tests for paired samples were carried out [43].

Effect on input space
To analyse the effect of the padding type in the input space, we studied the distribution of the activations of the 1D Convolutional layer for the 1_conv model. This layer has sixty-four filters of size 5 (see Figure S5 of the Supporting Information).

We randomly selected seven proteins of each type (for task 1) and two proteins of each type (for task 2) from the test set. For the two tasks separately, for each type of padding (except aug_padding) and each fold, we used the final model to predict on those proteins. We extracted the activations of the 1D convolutional layer for each prediction and separated each one of the sixty-four filters as a different sample. This resulted in a matrix of dimensions $64 \times 1000$ representing the activations for each prediction. Stacking the activation data of all the predictions (10 folds $\times 7$ types of padding $\times 14$ enzymatic sequences = 980 activations matrices of size $64 \times 1000$) separatedly for each task, we performed a PCA to study and compare the distribution of these activations.

Explanatory models
The performance metrics were further described through linear models built upon different variables that could affect to the model behaviour. These explanatory models have already been used for similar purposes [44, 14] and provide a way of statistically quantifying and comparing the relevance of the considered variables on the models performance.

Differences in performance between the different types of padding for both tasks were explained and tested in terms of the following linear model:

$$F1 \sim \text{architecture} + \text{enzyme\_type} + \text{type\_padding}$$

This full additive model was used as a snapshot of the general contribution of each factor to the F1-score. Reference category for enzyme type was 0 (non-enzyme) for task 1 and enzyme type 1 for task 2, only_denses for architecture and post_padding for padding type. However, to answer more specific questions about the effect of padding on the different architectures and enzyme types, we built more precise, appropriate models in each case (Equations 3, 2 and 4). Some of them included
**Questions on the effect of padding addressed through the explanatory models.**

| Question                                                                 | Equation | Architecture | Padding type                  |
|--------------------------------------------------------------------------|----------|---------------|-------------------------------|
| A  Does padding position affect performance?                             | Eq. 2    | only_denses   | post-, pre-, mid-, ext-, strf-, md-, zoom-, aug- |
| B  What is the effect of switching between dense paddings?               | Eq. 2    | stack_conv    | post-, pre-, mid-, ext-       |
| C  What is the effect of changing from the standard dense padding to sparse padding? | Eq. 2    | stack_conv    | post-, strf-, md-, zoom-      |
| D  Is an ensemble of paddings beneficial?                                 | Eq. 3    | stack_conv    | aug-, post-, strf-            |
| E  Does the effect of padding type on performance depends on the enzyme type? | Eq. 4    | stack_conv    | post-, strf-                  |

Table 3 The explanatory model for each question is specified by the column Equation. The Architecture column and the Padding type column show the architectures and padding types included in each comparison, respectively. All the enzyme types are included for each question. Reference categories are indicated in bold.

an interaction term (represented by $\var{1} : \var{2}$) to check if the effect of $\var{1}$ on the F1-score depends on the value of $\var{2}$. For example, adding $\text{type_padding} : \text{architecture}$ would let us identify those cases where the effect of changing the type of padding is different between architectures. Reference category for enzyme type was still 0 for task 1 and enzyme type 1 for task 2, but for architecture and padding type it varies in each case. Table 3 summarizes the questions addressed through the explanatory models, their equations and the reference levels in each case. We considered only_denses as reference in Question A because it is the baseline model and we aimed to check if in this case, different paddings affect differently to model performance. In questions B-E, stack_conv is chosen as reference architecture since it is the more complex and thus, the closest to the state of the art. Question E was only applied to task 2 results because its aim is to check the effect of enzyme types.

\[
F1 \sim \text{enzyme_type} + \text{type_padding} \quad (2)
\]

\[
F1 \sim \text{architecture} + \text{enzyme_type} + \text{type_padding} + \\
+ \text{type_padding} : \text{architecture} \quad (3)
\]

\[
F1 \sim \text{architecture} + \text{enzyme_type} + \text{type_padding} + \\
\text{type_padding} : \text{enzyme_type} \quad (4)
\]

Linear models were built in the R statistical programming language [45]. P-values were adjusted for multiple testing by the False Discovery Rate (FDR) by Benjamini-Hochberg [46].

**Declarations**

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.
Availability of data and materials
The UniprotKB/Swiss-Prot database (version 2019_05) protein entries analysed during the current study can be accessed and downloaded through the following link: http://ftp.ebi.ac.uk/pub/databases/uniprot/previous_releases/release-2019_05/knowledgebase/uniprot_sprot-only2019_05.tar.gz. Since this data needs further filtering to get only taxonomy Archaea, we have also included data analysed in this article as the Additional File 2. The code is publicly available at https://github.com/b2slab/padding_benchmark.

Competing interests
The authors declare that they have no competing interests.

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Author’s contributions
RS conceived the presented idea. ALdR designed the models and the computational framework and analysed the data. RS and APL verified the analytical methods. MM supervised the findings of this work. ALdR wrote the original manuscript. All authors read and approved the final manuscript.

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References
1. Krizhevsky, A., Sutskever, I., Hinton, G.E.: ImageNet Classification with Deep Convolutional Neural Networks. In: Pereira, F., Burges, C.J.C., Bottou, L., Weinberger, K.Q. (eds.) Advances in Neural Information Processing Systems 25, pp. 1097–1105. Curran Associates, Inc., ??? (2012).
2. Angermueller, C., Parnamaa, T., Parts, L., Stegle, O.: Deep learning for computational biology. Mol Syst Biol 12 (2016). doi:10.15252/msb
3. Ersalan, G., Avcen, Z., Gagneur, J., Theis, F.J.: Deep learning: new computational modelling techniques for genomics. Nature Reviews Genetics 20(7), 389–403 (2019). doi:10.1038/s41576-019-0122-6
4. Lecun, Y., Bengio, Y., Hinton, G.: Deep Learning. Nature 521(7553), 436–444 (2015). doi:10.1038/nature14539. arXiv:1312.6144v5
5. Alipanahi, B., Delong, A., Weiarch, M.T., Frey, B.J.: Predicting the sequence specificities of DNA- and RNA-binding proteins by deep learning. Nature Biotechnology 33(8), 831–838 (2015). doi:10.1038/nbt.3100
6. Quang, D., Xie, X.: DanQ: a hybrid convolutional and recurrent deep neural network for quantifying the function of DNA sequences. Nucleic Acids Research 44(11), 107–107 (2016). doi:10.1093/nar/gkw226
7. Müller, A.T., Hiss, J.A., Schneider, G.: Recurrent Neural Network Model for Constructive Peptide Design. Journal of Chemical Information and Modeling. (2017). doi:10.1021/acs.jcim.7b00414
8. Li, Y., Quang, D., Xie, X.: Understanding Sequence Conservation With Deep Learning. bioRxiv, 400–406 (2017). doi:10.1145/3107411.3107425
9. Rifaioglu, A.S., Doğan, T., Martin, M.J., Cetin-Atalay, R., Atalay, V.: Deepred: Automated protein function prediction with multi-task feed-forward deep neural networks. Scientific reports 9(1), 7344 (2019)
10. Kauferm-Abrams, E.: Quantifying Translation-Invariance in Convolutional Neural Networks. arXiv.org, 1801–014501 (2017). 1801.01450
11. Evans, R., Jumper, J., Kirkpatrick, J., Sifre, L., Green, T.F.G., Qin, C., Zidek, A., Nelson, A., Bridgland, A., Penedones, H., Petersen, S., Simonyan, K., Crossan, S., Jones, D.T., Silver, D., Kavukcuoglu, K., Hassabis, D., Senior, A.W.: De novo structure prediction with deep learning based scoring. Thirteenth Critical Assessment of Techniques for Protein Structure Prediction (Abstracts) (2018).
12. ¨Ozt¨ urk, H.,¨Ozg¨ ur, A., Ozkirimli, E.: DeepDTA: Deep drug-target binding affinity prediction. Bioinformatics 34(17), 821–829 (2018). doi:10.1093/bioinformatics/bty953. 1801.10193
13. Almagro Armenteros, J.J., Sønderby, C.K., Sønderby, S.K., Nielsen, H., Winther, O.: DeepLoc: prediction of protein subcellular localization using deep learning. Bioinformatics (Oxford, England) 33(21), 3387–3395 (2017). doi:10.1093/bioinformatics/btx431
14. Lopez-Del Rio, A., Nonell-Canals, A., Vidal, D., Perera-Lluna, A.: Evaluation of Cross-Validation Strategies in Sequence-Based Binding Prediction Using Deep Learning. Journal of Chemical Information and Modeling 59(4), 1645–1657 (2019). doi:10.1021/acs.jcim.8b00663
15. Hochreiter, S., Heusel, M., Obermayer, K.: Fast model-based protein homology detection without alignment. Bioinformatics 23(14), 1729–1736 (2007). doi:10.1093/bioinformatics/btm247
16. Di Lena, P., Nagata, K., Baldi, P.: Deep architectures for protein contact map prediction. Bioinformatics 28(19), 2449–2457 (2012). doi:10.1093/bioinformatics/bts475

17. Li, Y., Wang, S., Umarov, R., Xie, B., Fan, M., Li, L., Gao, X.: DEEPPre: Sequence-based enzyme EC number prediction by deep learning. Bioinformatics 34(5), 760–769 (2018). doi:10.1093/bioinformatics/btx680

18. Domingos, P.: A few useful things to know about machine learning. Communications of the ACM 55(10), 75–87 (2012). doi:10.1145/2347736.2347755

19. Jing, X., Dong, Q., HONG, D., Lu, R.: Amino acid encoding methods for protein sequences: a comprehensive review and assessment. IEEE/ACM Transactions on Computational Biology and Bioinformatics PP(1), 1–1 (2019). doi:10.1109/tcbb.2019.2911677

20. Ahmad, S., Sarai, A.: PSSM-based prediction of DNA binding sites in proteins. BMC Bioinformatics 6, 1–6 (2005). doi:10.1186/1471-2105-6-33

21. Lenselink, E.B., Ten Dijke, N., Bongers, B., Papadatos, G., van Vlijmen, H.W.T., Kowalczyk, W., IJzerman, A.P., van Westen, G.J.P.: Beyond the hype: deep neural networks outperform established methods using a ChEMBL bioactivity benchmark set. Journal of cheminformatics 9(1), 45 (2017). doi:10.1186/s13321-017-0232-0

22. Strömbergsson, H., Lapins, M., Kleywegt, G.J., Wågberg, J.E.S.: Towards Proteome-Wide Interaction Models Using the Proteochemometrics Approach. Molecular Informatics 29(6-7), 499–508 (2010). doi:10.1002/minf.201000052

23. Yang, K.K., Wu, Z., Bedbrook, C.N., Arnold, F.H.: Learned protein embeddings for machine learning. Bioinformatics 34(15), 2642–2648 (2018). doi:10.1093/bioinformatics/bty178

24. Asgari, E., Moftad, M.R.K., Bluhm, W., Christie, C., Dimitropoulos, D., Dutta, S.: Continuous Distributed Representation of Biological Sequences for Deep Proteomics and Genomics. PLOSONE 10(11), 0141287 (2015). doi:10.1371/journal.pone.0141287

25. Mazzaferro, C.: Predicting Protein Binding Affinity With Word Embeddings and Recurrent Neural Networks (2017). doi:10.1101/128223

26. Kimothi, D., Soni, A., Biyani, P., Hogan, J.M.: Distributed Representations for Biological Sequence Analysis. arXiv.org, 1608–059492 (2016). doi:10.1145/1235. arXiv:1608.05949v2

27. Juritz, V.I., Johansen, A.R., Nielsen, M., Almagro Armenteros, J.J., Nielsen, H., Sønderby, C.K., Winther, O., Sønderby, S.K.: An introduction to deep learning on biological sequence data: examples and solutions. Bioinformatics 33(22), 3685–3690 (2017). doi:10.1093/bioinformatics/btx531

28. Mirabella, C., Wallner, B.: RAWMSA: End-to-end Deep Learning using raw Multiple Sequence Alignments. PLoS ONE 14(8) (2019). doi:10.1371/journal.pone.0220182

29. Dwarampudi, M., Reddy, N.V.S.: Effects of padding on LSTMs and CNNs. arXiv 1903.07288 (2019). 1903.07288

30. Vinyals, O., Bengio, S., Kudlur, M.: Order matters: Sequence to sequence for sets. 4th International Conference on Learning Representations, ICLR 2016 - Conference Track Proceedings, 1–11 (2016). 1511.06391

31. Shroten, C., Koshgoftar, T.M.: A survey on Image Data Augmentation for Deep Learning. Journal of Big Data 6(1) (2019). doi:10.1186/s40537-019-0197-0

32. Shen, H.B., Chou, K.C.: EzyPred: A top-down approach for predicting enzyme functional classes and subclasses. Biochemical and Biophysical Research Communications 364(1), 53–59 (2007). doi:10.1016/j.bbrc.2007.09.098

33. Furnham, N., Sillitoe, I., Holliday, G.L., Cuff, A.L., Laskowski, R.A., Orenco, C.A., Thornton, J.M.: Exploring the Evolution of Novel Enzyme Functions within Structurally Defined Protein Superfamilies. PLoS Computational Biology 8(3), 1002403 (2012). doi:10.1371/journal.pcbi.1002403

34. Holliday, G.L., Andreini, C., Fischer, J.D., Rahman, S.A., Almonacid, D.E., Williams, S.T., Pearson, W.R.: MACIE: Exploring the diversity of biochemical reactions. Nucleic Acids Research 40(D1), 783–789 (2012). doi:10.1093/nar/gkr799

35. Shrikumar, A., Greenside, P., Kundaje, A.: Learning Important Features Through Propagating Activation Scores. arXiv.org, 1710–05623 (2017). doi:10.1101/171005623

36. Pan, X., Shen, H.-B.: Predicting RNA–protein binding sites and motifs through combining local and global deep convolutional neural networks. Bioinformatics 34(20), 3427–3436 (2018). doi:10.1093/bioinformatics/bty364. http://oup.prod.sis.lan/bioinformatics/article-pdf/34/20/3427/26024370/bty364.pdf

37. Zheng, X., Xu, S., Zhang, Y., Huang, X.: Nucleotide-level convolutional neural networks for pre-mirna classification. Scientific reports 9(1), 628 (2019).

38. Consortium, U.: UniProt: the universal protein knowledgebase. Nucleic Acids Research 45(D1), 158–169 (2017). doi:10.1093/nar/gkw1099

39. Srivastava, N., Hinton, G., Krizhevsky, A., Sutskever, I., Salakhutdinov, R.: Dropout: A Simple Way to Prevent Neural Networks from Overfitting. J. Mach. Learn. Res. 15, 1929–1958 (2014)

40. Kingma, D.P., Ba, J.L.: Adam: A Method for Stochastic Optimization. arXiv.org, 1412–6980 (2014)

41. Chollet, F.: Keras. GitHub. https://keras.io (accessed September 10, 2018) (2015)

42. Abadi, M., Agarwal, A., Barham, P., Brevdo, E., Chen, Z., Citro, C., Corrado, G.S., Davis, A., Dean, J., Devin, M., Ghemawat, S., Goodfellow, I., Harp, A., Irving, G., Isard, M., Jia, Y., Jozefowicz, R., Kaiser, L., Kurach, M., Levenberg, J., Mané, D., Monga, R., Moore, S., Murray, D., Olah, C., Schuster, M., Shlens, J., Steiner, B., Sutskever, I., Talwar, K., Tucker, P., Vanhoucke, V., Vasudevan, V., Viégas, F., Vinyals, O., Warden, P., Wattenberg, M., Wicke, M., Yu, Y., Zheng, X.: TensorFlow: Large-Scale Machine Learning on Heterogeneous Systems. (accessed September 13, 2018) (2015). http://tensorflow.org/

43. Wilcoxon, F.: Individual Comparisons by Ranking Methods. Technical Report 6 (1945). http://www.jstor.org/about/terms.html.

44. Picart-Armada, S., Barrett, S.J., Wille, D.R., Perera-Lluna, A., Gutteridge, A., Essaizy, B.H.: Benchmarking network propagation methods for disease gene identification. PLOS Computational Biology 15(9), 1–24 (2019). doi:10.1371/journal.pcbi.1007276

45. R Core Team: R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna, Austria (2015). R Foundation for Statistical Computing. https://www.R-project.org/
46. Benjamini, Y., Hochberg, Y.: Controlling the false discovery rate: A practical and powerful approach to multiple testing. J. R. Stat. Soc. Series B Stat. Methodol., 289–300 (1995)

Figures

Figure 1 Schematic explanation of one-hot encoding, zero-padding and truncation of amino acid sequences A. Amino acid sequences of different lengths are shaped to the common dimension of 7 by truncating or padding zeros at the end. B. Amino acid sequence at common length $L$ is transformed to a binary matrix $(n + 1) \times L$, being $n$ the number of different amino acids and placeholders. Each column of this matrix is full of zeros, being one only in the position of the corresponding amino acid.

Figure 2 Macro F1-score on test for each type of padding in each tested architecture. Each boxplot comprises 10 data points (which is the number of folds).

Figure 3 Density representation of the PCA of the activations of the convolutional layer. Figure shows PC1 vs PC2 of the activations of the 1D Convolutional layer for the 1_conv model in task 2, for the 10 folds. This representation comes from applying a PCA to the convolutional filter activations after the prediction of 14 enzymatic sequences of different EC number using each padding type. Then, the graphical representation was stratified by padding and enzyme type. We can see that according to the structure of the distributions, there seems to be two different groups of enzymes: 2, 3, 4 and 1, 5, 6, 7. Regarding types of padding, the activations for dense paddings are similar between them (two clusters separated along PC1) and different from sparse paddings.

Figure 4 Linear models on F1-score for both tasks 1 and 2 focusing on five specific questions. The “No interaction” facet represent estimates of additive terms of the model, while the other facet represent the interaction between two factors. Models regarding questions A, B and C have no interaction terms and thus they only have the “No interaction” facet. Only coloured tiles correspond to model coefficients; the white ones are outside the model specification. The colour of each category of a term represents the value of the estimate: red tiles correspond to positive estimates, specified with a “+”; blue tiles correspond to negative estimates, specified by “-” and grey tiles are close to zero. Framed categories are those that have a significant effect on each question (adjusted p-value < 0.05). An example on how to interpret this figure: in Model B task 2, switching from post_padding (reference) to pre_padding in stack_conv (reference) implies a decrease in performance.

Figure 5 Different types of padding analysed in this study. Each color bin represents an amino acid. Black lines represents zero padding.

Figure 6 Hierarchical enzyme classification with two levels. Task 1 classifies enzyme/non-enzyme and if enzyme, task 2 classifies by the first digit of the EC number.

Additional Files
Additional file 1 — Supporting Information
Detailed information and supplementary figures on data distribution, deep learning model parameters and architectures and supplementary model performance and linear models estimates tables and figures.

Additional file 2 — Data
CSV file with the Uniprot release 2019,05 reviewed entries for taxonomy Archaea.