

# Predicting properties of biological sequences using n-gram analysis

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*In silico* research allows scientists to more efficiently design and conduct experimental studies.

Examples:

- prediction of protein properties (presence of signal peptides, amyloidogenicity),
- predicting culture conditions of bacteria.

Machine learning models can help in the understanding of biological phenomena provided that they are not black boxes.

Create efficient methods for analysis of amyloids that have human-readable decision rules.

n-grams

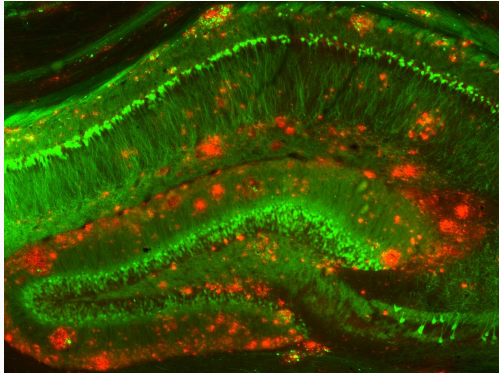
Simplified alphabets

Prediction of amyloidogenicity

Perspectives and summary

# Amyloid proteins

Amyloid are aggregate-forming proteins associated with various diseases (e.g., Alzheimer's, Creutzfeldt-Jakob's and Huntington's diseases).



Amyloid aggregates (red) around neurons (green). Strittmatter Laboratory, Yale University.

# Amyloid proteins

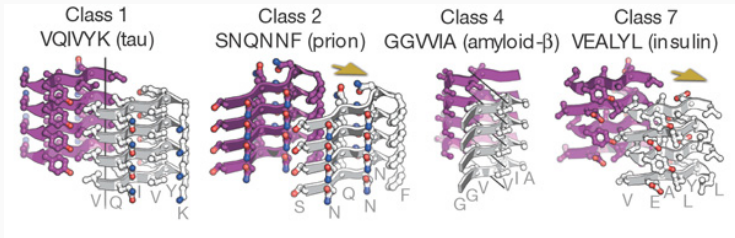
Functional amyloids:

- Pmel17,
- RIP1 and RIP3,
- acrosomal matrix proteins,
- HET-s,
- proteinaceous scaffolds of biofilms.

# Amyloid proteins

Hot-spots:

- short (6-15 amino acids),
- very high variability of amino acid composition,
- initiate amyloid aggregation,
- create specific "zipper-like"  $\beta$ -structures.



Sawaya et al. (2007)



## n-grams

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Computational analysis of biological sequences requires converting them to features understandable by machines.

The optimal conversion of information:

- loss-less,
- concise.

n-grams (k-tuples, k-mers):

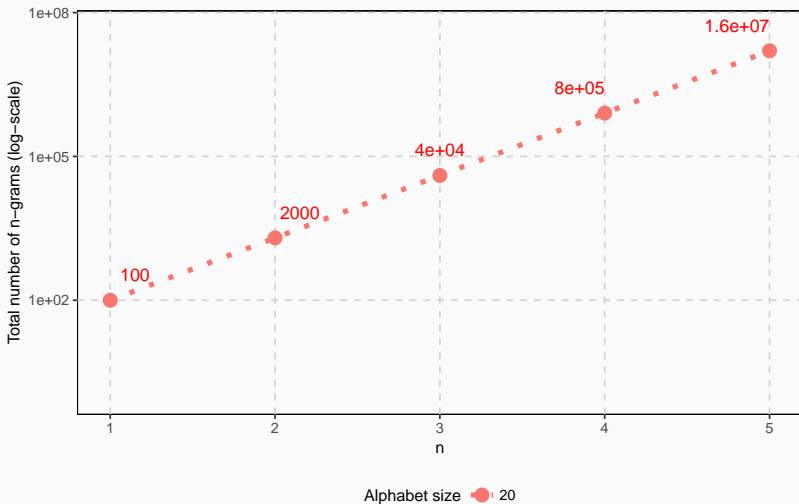
- subsequences (continuous or gapped) of  $n$  residues,
- considers the context of a specific residue.

	P1	P2	P3	P4	P5
S1	M	R	K	L	Y

2-grams: MR, RK, KL, LY

2-grams (gap 1): M – K, R – L, K – Y

3-grams: MRK, RKL, KLY



Longer n-grams are more informative, but create larger feature spaces, which are hard to process and analyze.

# Permutation Tests

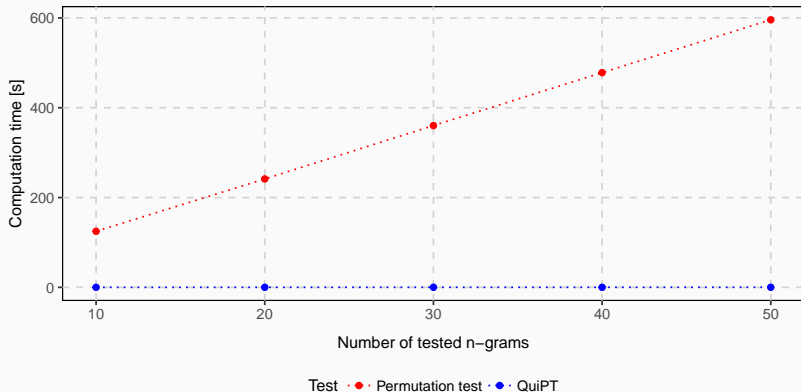
Informative n-grams are usually selected using permutation tests.

During a permutation test we shuffle randomly class labels and compute a defined statistic (e.g. information gain). Values of statistic for permuted data are compared with the value of statistic for original data.

$$\text{p-value} = \frac{N_{T_P > T_R}}{N}$$

$N_{T_P > T_R}$ : number of cases, where  $T_P$  (permuted test statistic) has more extreme values than  $T_R$  (test statistic for original data).

$N$ : number of permutations.



QuiPT (available as a part of the **biogram** R package) is faster than classical permutation tests and returns exact p-values.

# **Simplified alphabets**

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## Simplified alphabets:

- are based on grouping amino acids with similar physicochemical properties,
- ease computational analysis of a sequence (Murphy et al., 2000),
- create more explicit models.



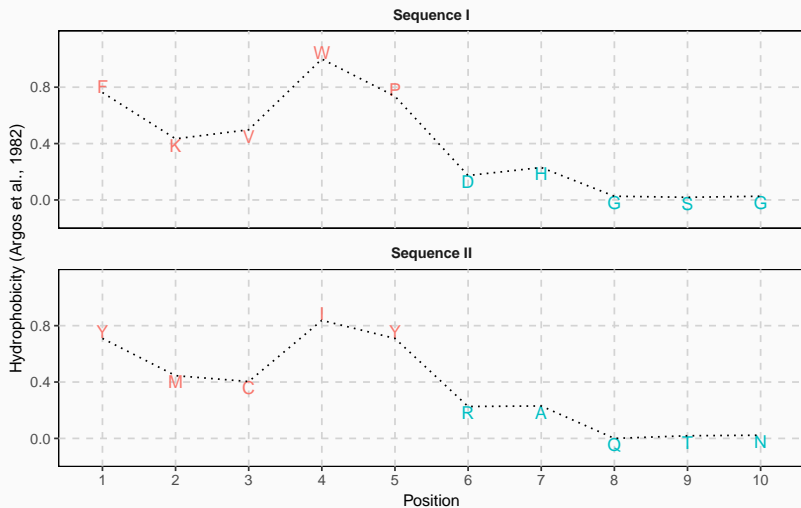
Two sequences that have drastically different amino acids composition may have very similar physicochemical properties.

Sequence I:

FKVWPDHGSG

Sequence II:

YMCIYRAQTN



Subgroup	Amino acid
1	C, I, L, K, M, F, P, W, Y, V
2	A, D, E, G, H, N, Q, R, S, T

Sequence I: FKVWPDHGSG                      1111122222

Sequence II: YMCIIYRAQTN                      1111122222

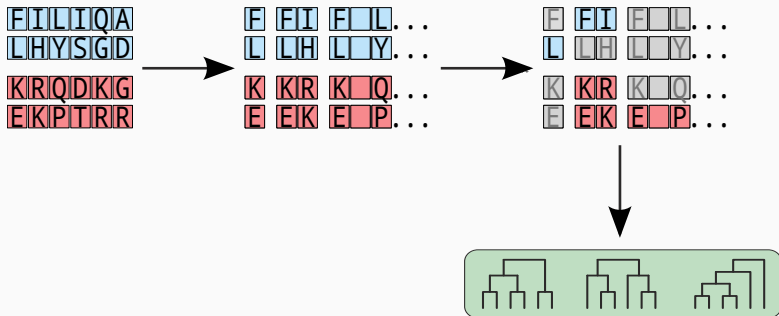
# Prediction of amyloidogenicity

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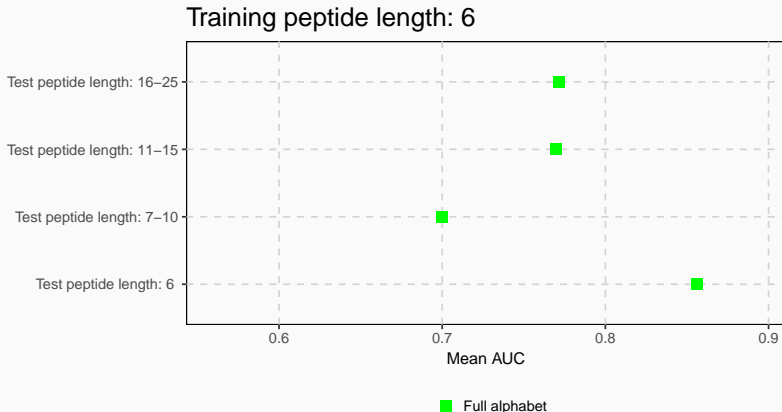
AmyLoad: a database of amyloid fragments (Wozniak and Kotulska, 2015).

- 1465 fragments,
- 11915 residues,
- 421 aggregation-prone fragments,
- 4312 residues (36.19%) in aggregation-prone fragments.

Can we predict amyloid fragments using n-gram data?



# Cross-validation



AUC (Area Under the Curve) measures the performance of a classifier (1 - classifier always properly recognizes amyloid proteins, 0 - classifier never properly recognizes amyloid proteins).

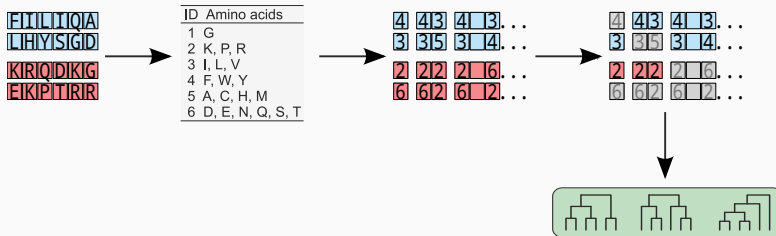


Does amyloidogenicity depend on the exact sequence of amino acids?

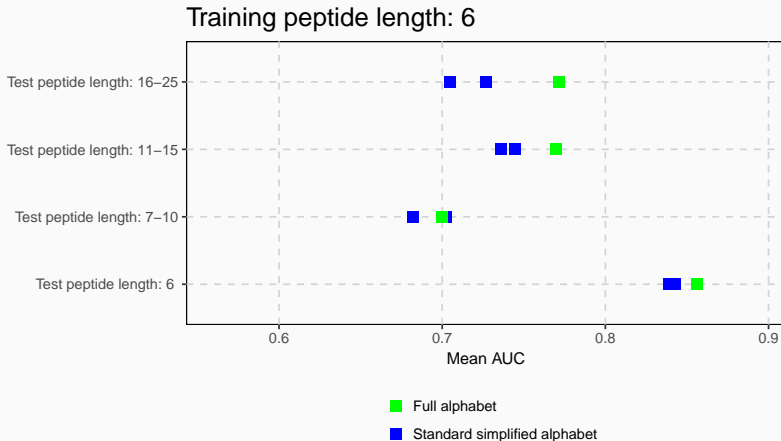
## Standard simplified amino acid alphabets

To date, several simplified amino acid alphabets have been proposed, which have been applied to (among others) protein folding and protein structure prediction (Kosiol et al., 2004; Melo and Marti-Renom, 2006).

# Standard simplified amino acid alphabets



# Cross-validation

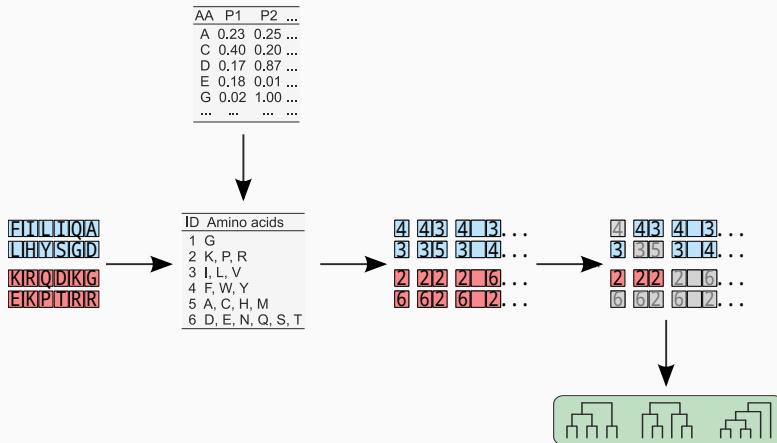


Standard simplified amino acid alphabets do not enhance discrimination between amyloidogenic and non-amyloidogenic proteins.

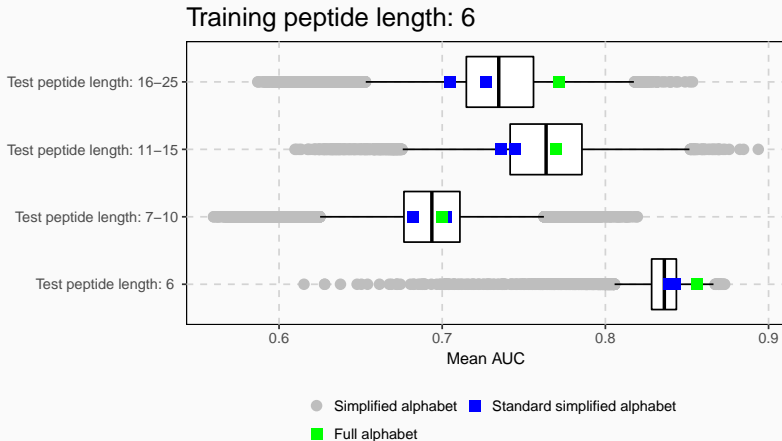
# Novel simplified amino acid alphabets

- 17 measures handpicked from AAIndex database:
  - size of residues,
  - hydrophobicity,
  - solvent surface area,
  - frequency in  $\beta$ -sheets,
  - contactivity.
- 524 284 amino acid simplified alphabets with different level of amino acid alphabet reduction (three to six amino acid groups).

# Novel simplified amino acid alphabets

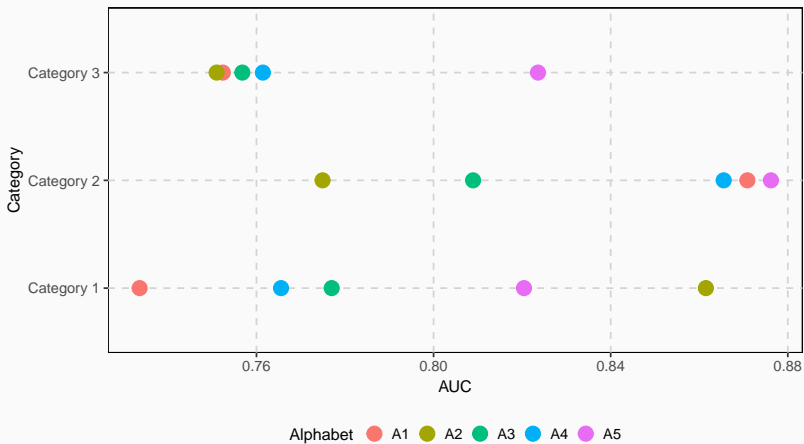


# Cross-validation



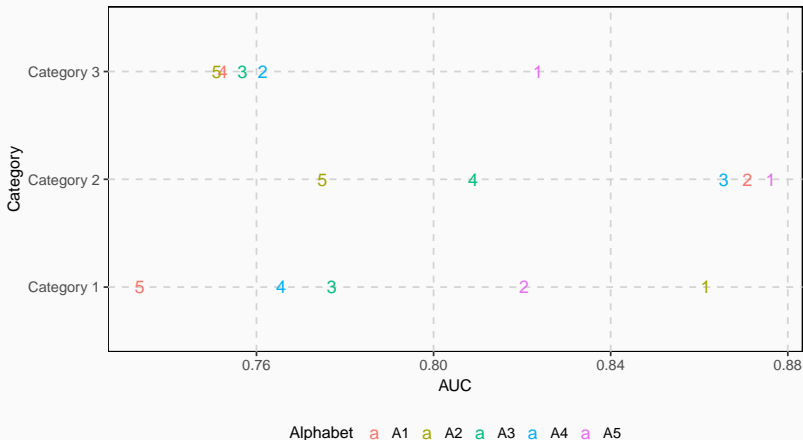
Hinges of boxes correspond to the 0.25 and 0.75 quartiles. The bar inside the box represents the median. The gray circles correspond to the simplified alphabets with the AUC outside the 0.95 confidence interval.

# Ranking alphabets



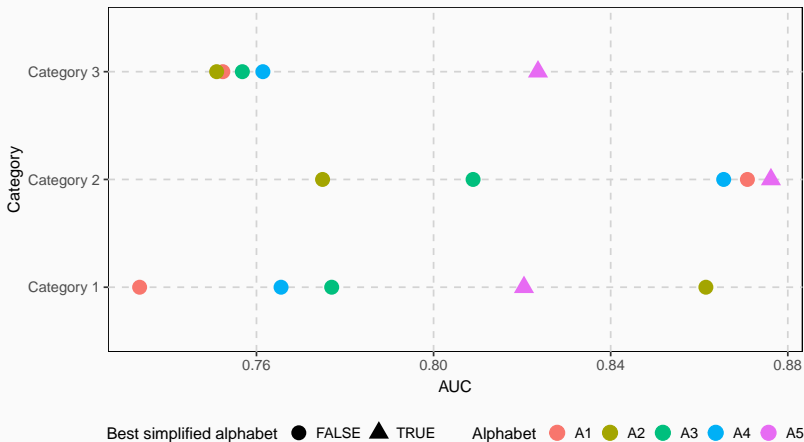


# Ranking alphabets



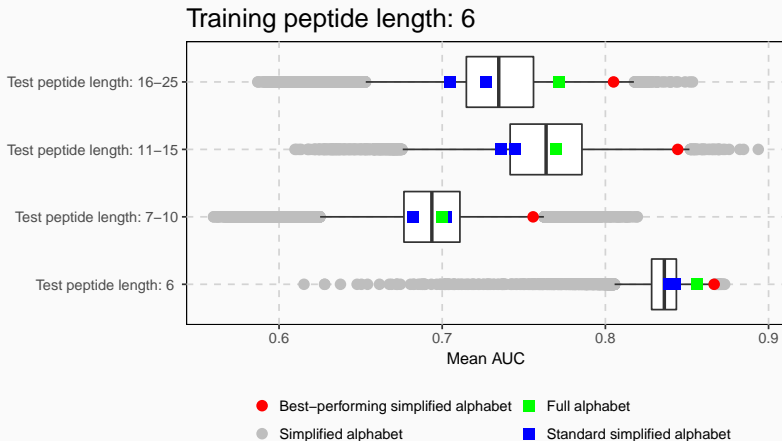
We rank alphabets separately in all length categories assuming the rank 1 for the best AUC, rank 2 for the second best AUC and so on.

# Ranking alphabets



The best-performing alphabet has the lowest sum of ranks.

# The best-performing simplified alphabet



# The best-performing simplified alphabet

Subgroup ID	Amino acids
1	G
2	K, P, R
3	I, L, V
4	F, W, Y
5	A, C, H, M
6	D, E, N, Q, S, T

## The best-performing simplified alphabet

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Group 3 and 4 - hydrophobic amino acids.

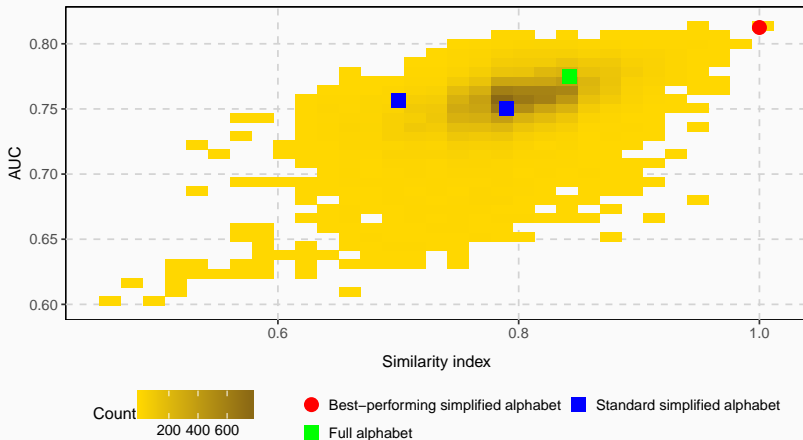
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Group 2 - charged breakers of  $\beta$ -structures.

Is the best-performing simplified amino alphabet associated with amyloidogenicity?

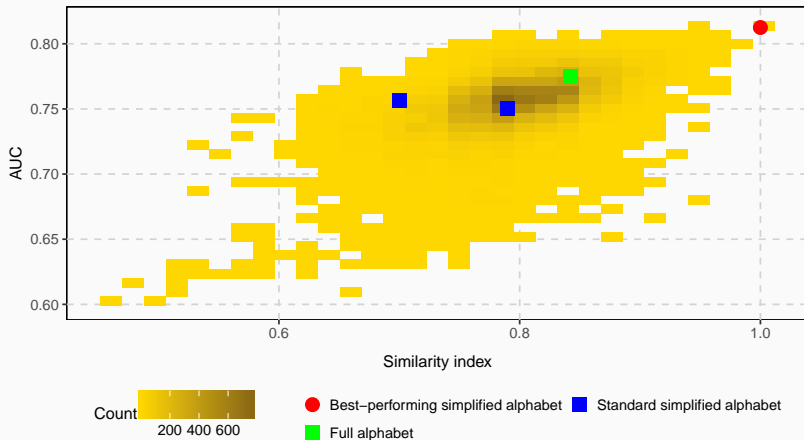
# Similarity index



Similarity index (Stephenson and Freeland, 2013) measures the similarity between two simplified alphabets (1 - identical, 0 - totally dissimilar).

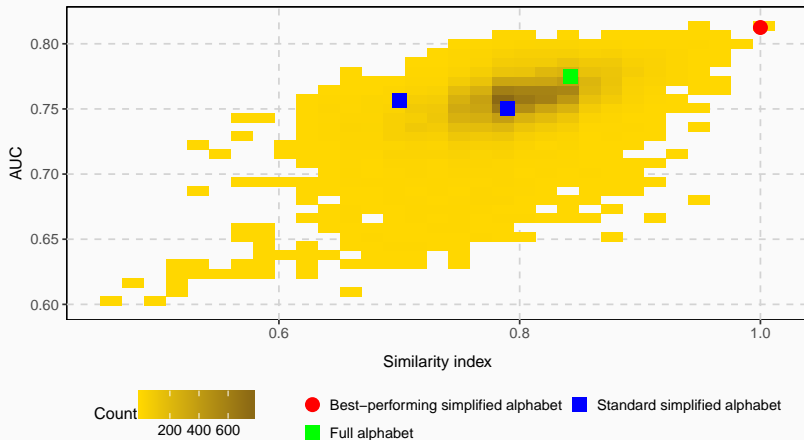


# Similarity index



The color of a square is proportional to the number of simplified alphabets in its area.

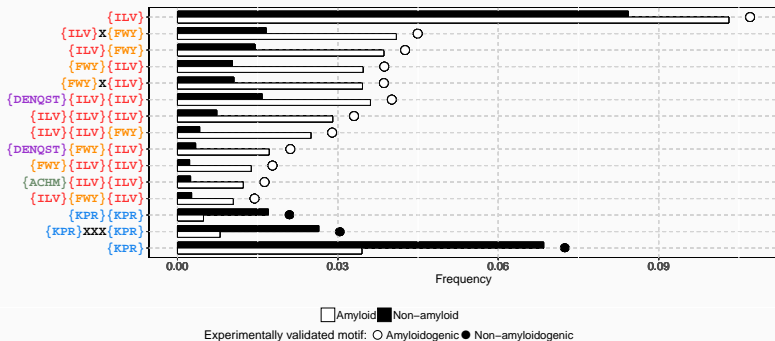
# Similarity index



The correlation between mean AUC and similarity index is significant ( $p\text{-value} \leq 2.2^{-16}$ ;  $\rho = 0.51$ ).

Are informative n-grams found by QuiPT associated with amyloidogenicity?

# Informative n-grams



Out of 65 the most informative n-grams, 15 (23%) were also found in the motifs validated experimentally (Paz and Serrano, 2004).

## Benchmark results

Classifier	AUC	MCC
AmyloGram	<b>0.8972</b>	<b>0.6307</b>
PASTA 2.0 (Walsh et al., 2014)	0.8550	0.4291
FoldAmyloid (Garbuzynskiy et al., 2010)	0.7351	0.4526
APPNN (Família et al., 2015)	0.8343	0.5823

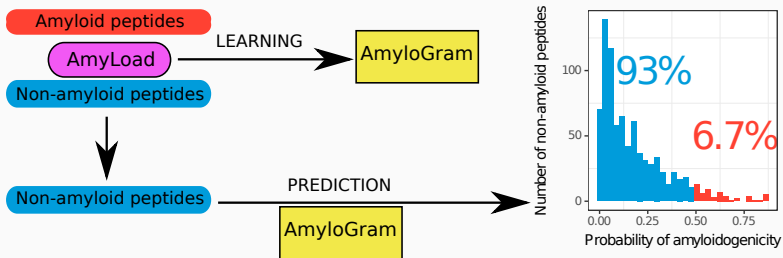
The predictor based on the best-performing alphabet, called AmyloGram, was benchmarked against the most popular tools for the detection of amyloid peptides using an external data set *pep424*.

## Benchmark results

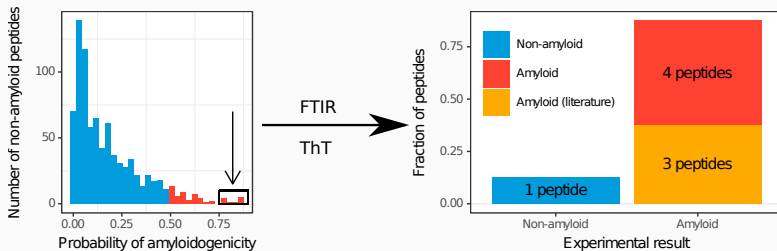
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MCC (Matthew's Correlation Coefficient) measures the performance of a classifier (1 - classifier always properly recognizes amyloid proteins, -1 - classifier never properly recognizes amyloid proteins).

# Experimental validation



# Experimental validation





## Perspectives and summary

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Improved prediction of amyloid proteins:

- hot-spots in the context of the whole protein,
- association of amino acid motifs and amyloidogenicity.

Goal: proteome-wide *in silico* detection of amyloid proteins.

Limitations: very few proteins with known aggregation-prone regions.

**AmyPro**: a database of amyloid proteins.

- 143 proteins,
- 40719 residues,
- 174 aggregation-prone regions,
- 5645 residues (13.86%) in aggregation-prone regions.

Seeding and cross-seeding: families of hot spots and relaxed seeding specificity.

- CsgA, CsgB,  $\alpha$ -synuclein,
- FapC and FapB.

Goal: identification of potential cross-seeding proteins in human microbiome.

Hot-spot specific inhibitors of amyloidogenicity (CsgC, TTR).

Goal: co-evolution of amyloid and its inhibitor.

Software packages:

- **biogram:**

<https://cran.r-project.org/package=biogram>.

- **AmyloGram:**

<https://cran.r-project.org/package=AmyloGram>.

Web servers:

- **AmyloGram:**

<http://www.smorfland.uni.wroc.pl/shiny/AmyloGram/>.

1. Created a new, accurate predictor of amyloids (Burdukiewicz et al., 2017).
2. Found a set of amino acid motifs associated with the amyloidogenic properties or the lack of them.
3. Found a simplified alphabet suitable for prediction of amyloid proteins.

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- KNOW Wrocław Center for Biotechnology.

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## **AmyloGram:**

<http://www.smorfland.uni.wroc.pl/shiny/AmyloGram/>.

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