

N-gram analysis of amyloid data

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Aim

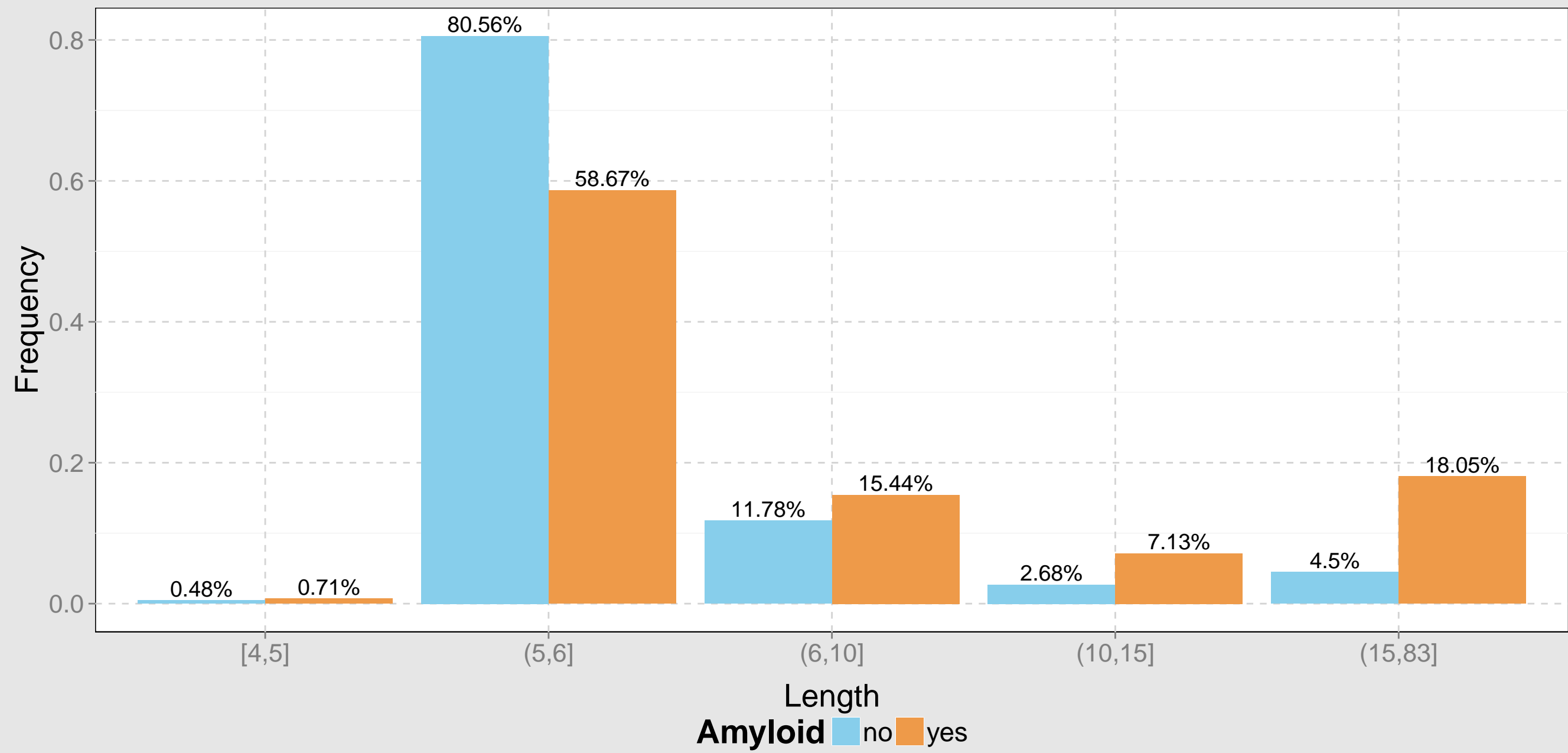
Investigate and predict sequences responsible for amyloidogenicity, the cause of various clinical disorders (e.g. Alzheimer’s or Creutzfeldt-Jakob’s diseases).

Introduction

All amyloidogenic sequences (amyloids), despite their variability in size and amino acid composition, form mostly cytotoxic aggregates (Breydo and Uversky, 2015). The hallmark trait of amyloids is the presence of hot-spots, short sequences of amino acids that play key role in the aggregation process (Fändrich, 2012).

AmyLoad database

The sequences used in the study (1044 non-amyloids and 421 amyloids) were extracted from AmyLoad database (Wozniak and Kotulska, 2015).



Clustering of amino acids

1. Nine scales representing properties important in the amylogenicity: hydrophobicity, size polarity and solvent accessibility from AAIndex database (Kawashima et al., 2008) were chosen. Additionally, two frequencies of forming contact sites (Wozniak and Kotulska, 2014) were added. All scales were normalized.
2. All combinations of characteristics (each time selecting only one scale per the property) were clustered using Euclidean distance and Ward’s method.
3. Each clustering was divided into 3 to 6 groups creating 144 encodings of amino acids.
4. Redundant 51 encodings (identical to other encodings) were removed.

Evaluation

1. Sequences shorter than 6 amino acids were discarded.
2. From each sequence overlapping 5-grams were extracted. All n-grams were labelled as their sequence of the origin (e.g. all 5-grams extracted from amyloid sequence were labelled as positive).
3. For each encoding features were filtered by the QuiPT and used to train the Random Forests (Liaw and Wiener, 2002). This procedure was performed independently on three training sets: a) 6 amino acids, b) 10 amino acids or shorter, c) 15 amino acids or shorter creating three classifiers.
4. All classifiers were evaluated in the 5-fold cross-validation. The sequence was labelled as positive (amylogenic), if at least one 5-gram was assessed as amylo-genic.
5. The cross-validation was repeated 8 times.

Encoding distance

The encoding distance between **A** and **B** is defined as the minimum number of amino acids that have to be shifted between subgroups of encoding **A** to make it identical to **B** (order of subgroups in the encoding and amino acids in a group is unimportant).

Group	Elements
1	a, b, c
2	d, e

Encoding **A**.

Group	Elements
1	a, b
2	d, e
3	c

Encoding **B**.

The encoding distance between **A** and **B** is 1 (element *c* must be moved from Group 3 to Group 1).

Specificity versus sensitivity

Training length	Number of groups	Encoding ID	AUC	Specificity	Sensitivity
6	3	6	0.7955	0.8221	0.6181
6	4	45	0.8183	0.9014	0.5038
<11	4	2	0.6615	0.4304	0.8307
<11	3	15	0.8088	0.8329	0.6060
<16	3	16	0.8162	0.7477	0.7374
<16	6	87	0.8320	0.5186	0.9195

Encodings with the best sensitivity and specificity for each training set type.

The best specificity encoding (training length 6, 4 groups, encoding ID 45) and the best sensitivity (training length <16, 6 groups, encoding ID 87) seem to have the different areas of the competence.

The committee of the best specificity and best sensitivity classifiers has overall **0.8911** AUC, **0.7473** sensitivity and **0.08684** specificity.

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