# Simplified alphabets in protein analysis

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## Outline

Simplified alphabets

Signal peptides

Methodology

Results

# Simplified alphabets

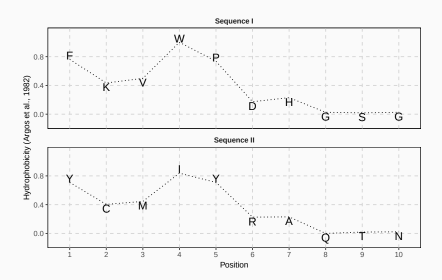
#### Simplified alphabets:

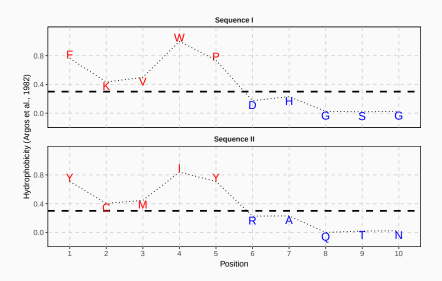
- are based on grouping amino acids with similar physicochemical properties,
- ease computational analysis of a sequence (Murphy et al., 2000),
- create more explicite models.

Two sequences that are drastically different considering their amino acids composition can have the same physicochemical properties.

Sequence I: FKVWPDHGSG

Sequence II: YCMIYRAQTN





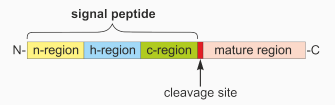
Subgroup	Amino acid
1	C, I, L, K, M, F, P, W, Y, V
2	A, D, E, G, H, N, Q, R, S, T

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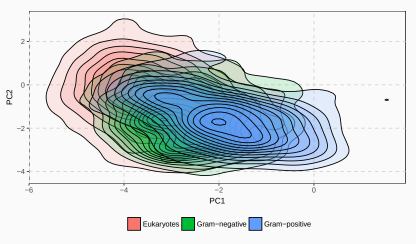
# Signal peptides



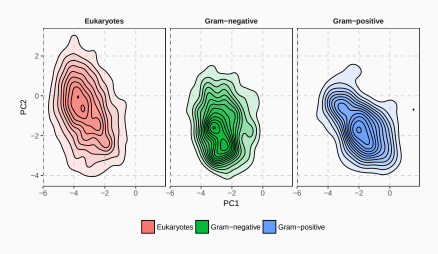
Signal peptides possess three distinct domains with variable length and characteristic amino acid composition (Hegde and Bernstein, 2006):

- n-region: mostly basic residues (Nielsen and Krogh, 1998),
- h-region: strongly hydrophobic residues (Nielsen and Krogh, 1998),
- c-region: a few polar, uncharged residues, similar to the mature protein.

Amino acid composition of signal peptides differ between eukaryotes and bacteria (gram positive and gram negative). Therefore, predictors of signal peptides are not universal and had to be taxon-specific.



PCA of amino acid frequency in signal peptides.



PCA of amino acid frequency in signal peptides.

#### Aim

Obtain a simplified alphabet to create a unified signal peptide predictor.

# Methodology

The are several algorithm for the effective probing of the alphabet space. They were created mostly with the protein folding/alignment in mind, but can be altered to work in prediction models.

#### Algorithms:

- branch and bound (Cannata, 2002),
- genetic algorithm without mutation (Palensky, 2006),
- genetic algorithm with mutation (Lenckowski and Walczak, 2007).

Genetic algorithm: cadidate solutions are individuals in a evolving population, where survival is depending on the quality of the solution.

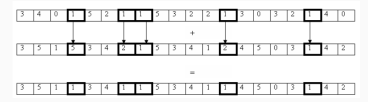
New individuals are created through cross-over and mutation of the fittest individuals.

# **Mutation operator**

Randomly change assignments of amino acids.

#### **Crossover operator**

Assignments of amino acids from a single subgroup belonging to a chosen parent alphabet overwrite assignments of amino acids belonging to the other chosen parent alphabet.



#### Caveat

Keep the number of groups constant. If any group is lost, randomly revert chosen amino acids to the missing groups.

#### **Fitness function**

Fitness function: a mean value of multiple  $\chi^2$  statistics.

We compare six data sets: 1-grams from signal peptides and mature proteins belonging to gram negative bacteria, gram positive bacteria and eukaryotes. The fitness function for a single comparison  $f_{single}$  is the value of  $chi^2$  statistic for following data (assuming the simplified alphabet of length 6):

	I	П	Ш	IV	V	VI
Taxon A	$x_{A,I}$	$x_{A,II}$	X <sub>A,III</sub>	$x_{A,IV}$	$x_{A,V}$	X <sub>A</sub> ,VI
Taxon B	x <sub>B,I</sub>	$x_{B,II}$	X <sub>B,III</sub>	X <sub>B,IV</sub>	$x_{B,V}$	x <sub>B,VI</sub>

 $x_{TAXON,N-GRAM}$  denotes the count of a specific 1-gram (a latin number) in a specific taxon (A or B).

#### **Fitness function**

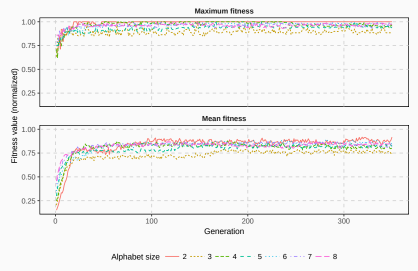
 $f_{\it final}$  is an **arithmetic mean** of multiple  $\chi^2$  statistics for all possible taxon comparisons:

Signal peptide origin	Mature peptide origin		
gram negative bacteria	gram negative bacteria		
gram positive bacteria	gram negative bacteria		
eukaryotes	gram negative bacteria		
gram negative bacteria	gram positive bacteria		
gram positive bacteria	gram positive bacteria		
eukaryotes	gram positive bacteria		
gram negative bacteria	eukaryotes		
gram positive bacteria	eukaryotes		
eukaryotes	eukaryotes		

# **Results**

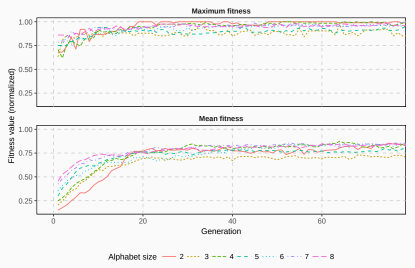
#### **Fitness function**

#### Fitness plot



# Fitness function (close-up)

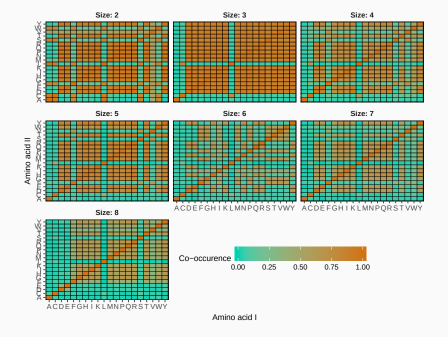
#### Fitness plot



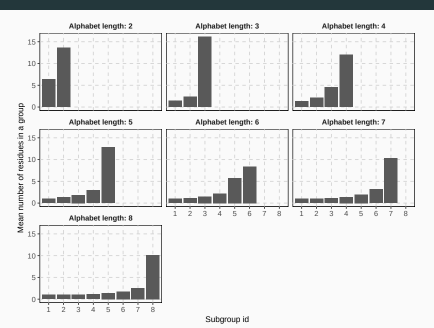
# Best simplified alphabets

To be absolutely sure that we have only alphabets in the fitness plateau, I considered only alphabets from generations 200-350.

To find the most common groupings of amino acids, I computed the co-occurence of amino acids in groups. The co-occurence is defined as the fraction of best-fitness alphabets, where the amino acid I is in the same group as the amino acid II.



# Preference for small groups



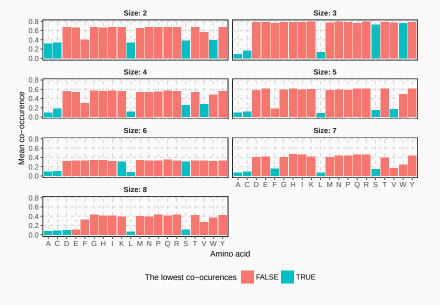
# Preference for small groups

The prevalence of groups of a length 1 may stem from the algorithm itself. Since the Walczak's algorithm keeps the alphabet length constant, when due to the crossover or mutation, a single group is missing, **randomly chosen amino acid** is altered to be in this group.

# Frequency standardization

Normalization by the global amino acid frequency removes the taxon-specific amino acid bias.

The global amino acid frequency is very similar to the amino acid frequency in mature proteins, because they are the majority in our data set.



Only fours residues had constantly the lowest co-occurence: A, C, L and S.

#### Mature protein С Е D G Н K Low co-occurence amino acids М N Q R **FALSE** TRUE Υ s Т ٧ W

euk gm gp

euk gm gp

euk gm gp

source

Normalized amino acid frequency

euk gm gp

euk gm gp

#### Signal peptide С D Е F 2 -G Н Κ Low co-occurence amino acids R **FALSE** Ν Р Q TRUE

W

euk gm gp

Υ

euk gm gp

٧

euk gm gp

source

Т

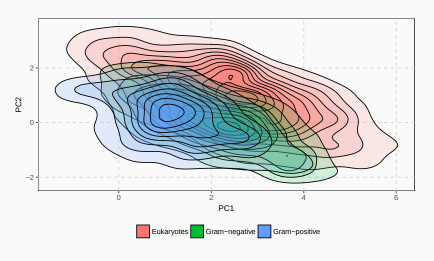
euk gm gp

Normalized amino acid frequency

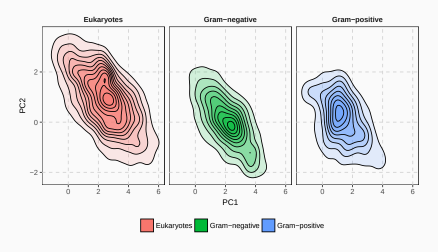
2 -

s

euk gm gp



PCA of amino acid frequency in signal peptides (reduced alphabet).



PCA of amino acid frequency in signal peptides (reduced alphabet).

## Summary

- Mean  $\chi^2$  is not the appropriate fitness measure.
- Repeats of spefic amino acids (A, C, L and S) are typical for signal peptides (Labaj et al., 2010).

## **Acknowledgments**

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#### References I

#### References

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