

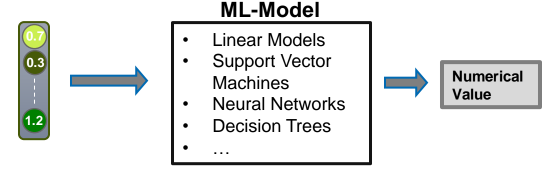


Traditional numerical representations for proteins

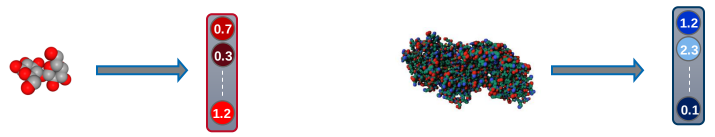
How can we make predictions for molecules?



Machine Learning models require numerical representations as input:



We need numerical representations of molecules

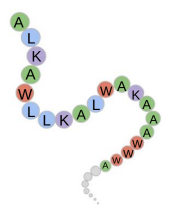


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Amino Acid Composition



- Amino Acid Composition (AAC):
 - Percentages of each amino acid in the sequence of that protein
 - Results in a vector $(f_1, f_2, ..., f_{20})$
 - Example:





Number of residues = 537

Molecular weight = 60673.44

$$\vec{f} = (4.5, 0, 1.5, ..., 3.2, 0)$$

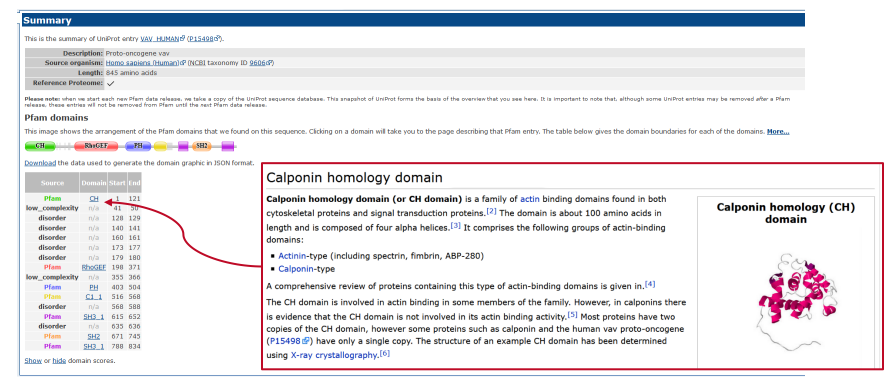
k-mer Features



- k-mer (=n-gram) features:
 - Creating a list of all possible amino acid sequences of length k
 - Calculate the frequencies of these strings in the protein amino acid sequence
 - Use the resulting features in a vector $(f_1, f_2, ..., f_n)$, $n = 20^k$
 - If n becomes too large, only use the most common substrings
 - For k = 1, we get the amino acid composition
 - Example for k = 3:
 - Calculating the frequency of each of the subsequences AAA, AAR, AAN, AAD, AAE, AAG, AAC, AAQ, AAH, (8000 possible 3-mer) within the protein amino acid sequence: $(f_1, f_2, ..., f_{8000})$
 - These subsequences can be overlapping:
 - "AAAM" contains both AAA and AAM

Pfam entries / Protein domains





http://pfam.xfam.org/ http://pram.xfam.org/

Protein Domains



- Protein Domains:
 - Subsequences of proteins with specific structural or functional characteristics
 - Pfam database: 20k different protein domains



The amino acid sequence of every protein can be mapped via the webservice of Pfam to functional domains

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