# UNIVERSITY OF BELGRADE FACULTY OF MATHEMATICS 

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# Data mining on protein sequences: n-gram analysis of ordered and disordered protein regions 

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# UNIVERZITET U BEOGRADU MATEMATIČKI FAKULTET 

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# Istraživanje podataka na proteinskim niskama: n-gramska analiza uređenih i neuređenih regiona proteina 

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#### Abstract

Proteins with intrinsically disordered regions are involved in large number of key cell processes including signaling, transcription, and chromatin remodeling functions. On the other side, such proteins have been observed in people suffering from neurological and cardiovascular diseases, as well as various malignancies. Process of experimentally determining disordered regions in proteins is a very expensive and longterm process. As a consequence, a various computer programs for predicting position of disordered regions in proteins have been developed and constantly improved.


In this thesis a new method for determining Amino acid sequences that characterize ordered/disordered regions is presented. Material used in research includes 4076 viruses with more than 190000 proteins. Proposed method is based on defining correspondence between n-grams (including both repeats and palindromic sequences) characteristics and their belonging to ordered/disordered protein regions. Positions of ordered/disordered regions are predicted using three different predictors.

The features of the repetitive strings used in the research include mole fractions, fractional differences, and z -values. Also, data mining techniques association rules and classification were applied on both repeats and palindromes. The results obtained by all techniques show a high level of agreement for a short length of less than 6 , while the level of agreement grows up to the maximum with increasing the length of the sequences. The high reliability of the results obtained by the data mining techniques shows that there are n-grams, both repeating sequences and palindromes, which uniquely characterize the disordered/ordered regions of the proteins. The obtained results were verified by comparing with the results based on n-grams from the DisProt database which contains the positions of experimentally verified disordered regions of the protein. Results can be used both for the fast localization of disordered/ordered regions in proteins as well as for further improving existing programs for their prediction.

## Keywords

n-gram, data mining, ordered/disordered regions, association rules, proteins

## Scientific field

Computer Science

## Scientific subfield

Data Mining

## Podaci o doktorskoj disertaciji

Naslov doktorske disertacije: Istraživanje podataka na proteinskim niskama: ngramska analiza uređenih i neuređenih regiona proteina

Rezime: Proteini koji imaju neuređene regione učestvuju u velikom broju ćelijskih procesa kao što su prenos signala, transkripcija i remodelovanje funkcija hromatina. Sa druge strane, pojava takvih proteina je uočena kod osoba koje boluju od neuroloških i kardiovaskularnih bolesti, kao i različitih oblika maligniteta. Eksperimentalno određivanje neuređenih regiona protiena je vrlo skup i spor proces. Zbog toga su razvijeni i stalno se usavršavaju različiti računarski programi za predviđanje pozicija neuređenih regiona u proteinu.

U radu je prikazana nova metoda za određivanje niski amino kiselina koje karakterišu neuređene i uređene regione proteina. Materijal nad kojim je vršeno istraživanje obuhvata 4076 virusa sa preko 190000 proteina. Metoda je zasnovana na ispitivanju osobina n-grama (koji obuhvataju ponavljajuće i palindromske niske) i njihove pripadnosti uređenim i neuređenim regionima proteina. Pozicije neuređenih /uređenih regiona u proteinima su određene korišćenjem tri programa za predviđanje. Osobine ponavljajućih niski koje su korišćene u istraživanju uključuju molske frakcije, frakcijske razlike i z-vrednost. Takođe, na ponavljajuće niske kao i na palindromske niske primenjene su određivanje pravila pridruživanja i klasifikacija, kao tehnike istraživanja podataka. Rezultati dobijeni svim tehnikama pokazuju visok nivo saglasnosti, za niske dužine manje od 6, dok nivo saglasnosti rezultata raste sve do maksimalnog sa porastom dužine niski. Visoka pouzdanost rezultata dobijenih tehnikama istraživanja podataka, pokazuje da postoje n-grami, kako ponavljajuće sekvence tako i palindromi, koji jednoznačno karakterišu neuređene/uređene regione proteina. Dobijeni rezultati su provereni upoređivanjem sa rezultatima zasnovanim ngramima iz DisProt baze koja sadrži pozicije eksperimentalno verifikovanih neuređenih regiona proteina, i mogu da budu korišćeni kako za brzu lokalizaciju neuređenih/uređenih regiona u proteinima tako i za dalje poboljšanje postojećih programa za njihovo predviđanje.

Ključne reči
n-gram, istrživanje podataka, uređeni/neuređeni regioni, pravila pridruživanja, proteini
Naučna oblast
Računarstvo

Naučna podoblast<br>Istraživanje podataka

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## 1 Introduction

### 1.1 Bioinformatics

From its very beginning until the fourth quarter of the 20th century biology has been observational and experimental science. Recent development and using computers not altered completely this orientation, but introduce new methods and algorithms in processing of biological material. Nature of data has changed - data have become discret and more precise. The quantity of available data grow rapidly bringing to the scene a new discipline capable to provide efficient processing of data in the new conditions - Bioinformatics. Bioinformatics has a lot of subdisciplines and research directions [1]. Most pressing task of bioinformatics has moved to analyze and interpret various types of data, including nucleotide and amino acid sequences, protein structures and interactions, and so on. To meet the new requirements arising from the new tasks, researchers in the field of bioinformatics are working on the development of new algorithms (mathematical formulas, statistical methods, etc) and software tools which are designed for assessing relationships among large data sets stored, such as methods to locate a gene within a sequence, predict protein structure and/or function, understand diseases at gene expression level and etc.

A particular active area of research in bioinformatics is the application and development of data mining techniques to solve biological problems. Analyzing large biological data sets requires making sense of the data by inferring structure or generalizations from the data. Examples of this type of analysis include protein structure prediction, gene classification, cancer classification based on microarray data, clustering of gene expression data, statistical modeling of protein-protein interaction, etc.

### 1.2 Proteins

Proteins are biological macromolecules, of polymeric nature, that are built by forming of so called "peptide bond" (polypeptides) between their basic constituents amino acids (Figure 1). Amino acids (AA) are organic molecules that posses at least one amino $\left(-\mathrm{NH}_{2}\right)$ and carboxyl ( -COOH ) group. There are $20(+2)$ amino acids that constitute all, so far, known proteins. Protein structure and function are mainly determined by so called "protein primary structure", which represents amino acid content of protein molecule, its number and sequence (Figure 1). In bioinformatics amino acids are represented by one or three letter code as shown in Appendix table A1.



Figure 1. Forming of peptide bond between two amino acids (a), schematic representation of protein primary structure (b). Primary structure is "reed" from N- to C-terminal of polypeptide (protein) chain.

Protein "secondary structure" may be defined by so called "torsion angles", ( $\varphi$ and $\psi$ ), from Ramachandran diagram [2], between successive amino acids, that forms backbone of polypeptide chain (Figure 2.A and 2.B). If three or more pairs of torsion angles are the same, than there is a regular secondary structure. Secondary structure results from forming secondary, noncovalent H -bonds between $\mathrm{C}=\mathrm{O}$ and $\mathrm{H}-\mathrm{N}$ groups; the exact pattern of them is different in different forms of secondary structure. Two of most represented secondary structures in proteins are alpha ( $\boldsymbol{\alpha}$ ) helix structure and the beta ( $\boldsymbol{\beta}$ ) pleated sheet (Figure 3.).



Figure 2. A: Part of polypeptide chain with $\varphi$ and $\psi$ torsion angles between $\mathrm{C}_{\alpha}$ and N atom (from amino group) and $\mathrm{C}_{\alpha} \mathrm{C}$ atom from carboxyl group, $\omega$ torsion angle that corresponds to peptide link is small and usually neglected. B: Ramachandran diagram with marked areas that correspond to certain secondary structures.
Source: A - Jane S. Richardson, The Anatomy and Taxonomy of Protein Structure. In Advances in protein chemistry, Vol. 34 (1981)
B - https://www.studyblue.com/notes/note/n/protein-structure/deck/7778686


Figure 3. The $\alpha$-helix (A, B) and the $\beta$-pleated sheet (C, D) are the two principal secondary structures found in protein. $\mathrm{C}, \mathrm{N}, \mathrm{O}$ and H atoms involved in polypeptide chain forming.
Source: Bruce Alberts, Alexander Johnson, Julian Lewis, David Morgan, Martin Raff, Keith Roberts, Peter Walter, Molecular biology of the cell. 6 ed, (2015),
Garland Science, Taylor \& Francis Group, 711 Third Avenue, New York, NY 10017, US 3 Park Square, Milton Park, Abingdon, OX14 4RN, UK, ISBN 978-0-8153-4432-2

Protein tertiary structure refers to the spatial arrangement of a polypeptide chain through folding and coiling to produce a compact globular shape. It may be defined by knowing positions of all atoms that protein consists of [3, 4].

### 1.2.1 Intrinsically disordered proteins/protein regions (IDP/IDPR)

In last 15 years, it became more and more evident that a significant number of proteins, under physiological conditions, do not possess a well defined 3 dimensional ordered structure (Figure 4). They exhibit a variety of conformational isomers in which the atom positions and the polypeptide backbone torsion angles of the Ramachandran plot vary over time, with no specific equilibrium values, typically involving noncooperative conformational changes [5]. They may be completely or partially disordered and may undergo a disorder-to-order, or vice versa, transition upon interaction with other molecules. Thanks to their high structural mobility they readily interact with other molecules/proteins and carry out mostly regulatory functions related to molecular recognition, signal transduction, protein-protein, and protein-nucleic acid interaction.


Figure 4. SUMO-1 protein (PDB:1a5r), with central part that shows relatively ordered structure. The N - and C-terminal regions (left and right, respectively) are intrinsically disordered (grey disordered regions). Secondary structure elements: $\alpha$-helices (red), $\beta$ strands (blue arrows).
Source: http://www.rcsb.org/pdb/explore/explore.do?structureId=1a5r
In accordance to arising function, they are classified into, at least, 16 structural/functional categories, as listed in the DisProt database, that currently contain 803 experimentally determined IDP/IDPRs [6]. Taxonomically, IDPs are represented in
the proteomes of all of the three superkingdoms (Archaea, Bacteria and Eukarya), as well as in viruses. Primary structure of IDP/IDPRs are characterized by low sequence complexity (i.e. often consist of repetitive short fragments) and are biased toward polar and charged, but against bulky hydrophobic and aromatic AA residues (Figure 5), i.e., they are enriched in Ala, Arg, Gly, Gln, Ser, Glu, Lys and Pro and depleted in orderpromoting Trp, Tyr, Phe, Ile, Leu, Val, Cys, Asn AAs [7]. Experimentally, IDP/IDPRs may be detected by more than 20 various biophysical and biochemical techniques such as: x-ray diffraction crystallography, heteronuclear multidimensional NMR, circular dichroism, etc. Since IDP/IDPRs experimental study is costly and difficult (because of the lack of unique structure in the isolated form), a number of prediction tools have been developed [8].


Figure 5. Fractional differences in composition between disordered and ordered sets of regions, calculated on the basis of data from DisProt DB. On right part of diagram are amino acids with higher propensity to disorder.
Source: Predrag Radivojac, Lilia M. Iakoucheva, Christopher J. Oldfield, Zoran Obradovic, Vladimir N. Uversky, and A. Keith Dunker, Intrinsic Disorder and Functional Proteomics. Biophysical Journal Volume 92 March 2007 1439-1456. doi: 10.1529/biophysj.106.094045

## Disorder prediction

Disordered regions of the protein chain are important for the protein function. Today there are special programs (disorder predictors) that can predict them. IDP/IDPRs predictors can be grouped according characteristics or methods used for prediction. For example, one group include those that use physico-chemical properties of amino acids in proteins (PONDR, FoldUnfold, IUPred, GlobPlot, PreLINK, and FoldIndex), the second one those that use alignment of homologous protein sequences (Ronn, Disopred), etc. [9]. A summary of these methods can be found in Appendix Table A2.

Programs of the first group differ by the property of amino acids in proteins used for prediction of disordered regions. For example, PONDR uses local amino acid composition and hydrophobicity, FoldUnfold uses number of expected contacts, PreLINK uses propensity of a chain region to form a hydrophobic cluster, and IUPred uses estimation of the energy interaction between neighbouring amino acids. In the second group, the RONN program uses a neural network and compares the given sequence with a number of sequences whose structure can be a priori determined (ordered/disordered/mixture), while DISOPRED uses the network trained to distinguish regions that are missed in the structure obtained by x-ray analysis [10, 11, 12, 13].

### 1.3 Viruses

Viruses are small infectious agent that proliferates only inside the cells of all life forms: Archaea, Bacteria and Eukaryote. Outside of a cell viruses exist in the form of a virion, that consist of two, or three parts: (i) the genetic material made from either DNA or RNA; (ii) a protein coat, called the capsid, which surrounds and protects the genetic material; and in some cases (iii) an lipid envelope that surrounds the protein coat when they are outside a cell [14].

Genomic organization of viruses shows an enormous variety (as a group, they contain more structural genomic diversity than in all of three superkingdoms). Genome size varies greatly: in general, RNA viruses have smaller genome sizes than DNA viruses, although the smallest viral genome is that of ssDNA circoviruses (family Circoviridae), have a genome size of only two kilobases and code for only two proteins. The largest-genome size is that of the pandoraviruses of around two megabases, which code for about 2500 proteins. Virus genes are often arranged so that they overlap and rarely have introns.


Figure 6. Viral classification according to host and morphology.
(Source: Nucleic Acids Research 2011;39 (Database issue) :D576-D582)

Viral proteins exhibit distinct and structural features than the host proteins. There are several potentially unique characteristics of viral proteins, that include (a) the low contact densities, (b) the high occurrence of random coil segments and short disordered regions and (c) the lower destabilizing effects of mutations [15]. It has been shown that viruses have the largest variation range of the disordered residue fractions in their proteomes (human coronavirus NL63 has only 7.3\% disordered residues, while Avian carcinoma virus proteome has $77.3 \%$ disordered residues). Also, some viral species are highly enriched in intrinsic disorder. With the increase of proteome size, the fractions of disordered residues seem to converge to a range between 20 and $40 \%$. IDP/IDPRs help viruses to deal with their hostile habitats, in managing of their gene expression and generally, better adaptability and functioning of their proteins [16].

There are probably millions of different types of viruses, although only about 5,000 species have been described in detail. At the beginning of 2017 year, the NCBI Virus genome database has more than 7000 complete virus genomes. Viruses may be classified according to different criteria: their host and morphology (as shown on Figure 6), their morphology (symmetry and possession of envelope), genome organization (ds or ss; DNA or RNA) and in the case of Baltimore classification on mechanisms of viral genome replication (i.e., mechanism of viral mRNA production). This classification places viruses into seven groups as shown on Figure 7.

### 1.4 Topic of the dissertation

Because of importance of disorder regions for protein function, the research topic in this dissertation is to find amino-acids strings that characterize ordered/disordered protein regions. The aim is not to produce new disorder predictor, but to discover are there any AA (or series of AAs) that can be used as 'indicators' of region type, without pretension to determine exact boundaries of such regions.

The characteristics of AA can be mapped to the problem of finding characteristics of sequence of AAs (called n-gram) where the length of the sequence can be $1,2, \ldots, N$. There are different methods for characterization such n-grams in some environment (e.g. string), but no one can, in advance, determine characteristics that can be used as indicators, with high accuracy. This research will use set of viral proteins as material. Viruses from different phyla are used as material to minimize potential influence of group of specific phyla.


Figure 7. Baltimore classification of viruses (Source: http://www.nlv.ch/Virologytutorials/Classification.htm)

## 2 Methods for determining characteristics strings in protein regions

In this chapter idea for construction of specific model for determining n-grams that characterize disorder/order ( $\mathrm{D} / \mathrm{O}$ ) protein regions is described. First part of the chapter includes description of methods for n-gram characterization in the specific string. In the second part, idea for discover n-gram characteristics that are related to ordered/disordered regions in proteins is described. Second part also includes discussion about quality of proposed model.

### 2.1 N-gram analysis

There are many definitions of n-gram. In this research we used the following one [17]:

Definition 1: Given a sequence of letters $S=s_{1} s_{2} \ldots s_{N+(n-1)}$ over the alphabet $A$, with $N$ and $n$ a positive integer, an $n$-gram of the sequence $S$ is an n-long subsequence of consecutive letters. The i-th n-gram of $S$ is the sequence $s_{i} s_{i+1} \ldots s_{i+n-1}$.

There are $N$ such $n$-grams in $S$. For an alphabet $A$ with $|A|$ distinct letters, there are $|A|^{n}$ possible unique $n$-grams. Gram is a Greek word; depends on value of $n$, n-grams are denoted as monograms ( $n=1$ ), bigrams ( $n=2$ ), trigrams ( $n=3$ ), tetragrams ( $n=4$ ), pentagrams ( $n=5$ ), hexagrams ( $n=6$ ), etc. Some authors prefer using names unigram, bigram, trigram, quadrigram..., etc.

Simple n-gram analysis includes counting of specific n-gram occurrences in observed (analyzed) areas, as well as calculating the difference and, if applicable, the standard deviation of its occurring in those areas compared to the whole material. In this research the $n$-gram analysis for the occurrence of amino acids in the ordered/disordered
regions of proteins has been performed. N -Grams belong to any of the three regions including: disordered region (D), ordered region ( O ) and borderline transition from ordered to disordered region or vice versa ( N ) in the proteins, whereas monograms can belong to either D or O region only. For example, the amino acids in the sequence RAVERSQVSEN in a protein may correspond to the following ordered/disordered regions: OODODDDOOOO. The set of monograms in the sequence is $\{$ R A V E R S Q V S E N\} and their corresponding disordered/ordered characteristics are \{O O D O D D D O O O O\}. The set of bigrams for the above amino acids sequence is \{RA AV VE ER RS SQ QV VS SE EN\}, while corresponding ordered/disordered regions characteristics are $\{\mathrm{O} N \mathrm{~N} N \mathrm{D}$ D N O O O $\}$. Analogously, the set of the trigram representations of the above amino acids sequence is \{RAV AVE VER ERS RSQ SQV QVS VSE SEN\}, with the corresponding ordered/disordered region characteristics \{N N N N D N N O $\mathrm{O}\}$.

N -gram analysis has also been performed at the level of nucleotide sequence. Because nucleotide sequences are widespread across whole genome sequence, there are four (compared to three in the case of proteins) possible regions: disordered regions (D) which corresponds to the positions (in the genome sequence) of the disorder regions in proteins, ordered region (O) which corresponds to the positions (in the genome sequence) of the order regions in proteins, intergenic regions (I) which corresponds to the parts of the genome sequence that did not corresponds to any of the proteins, and borderline transition ( N ) between some of the previous three kinds of regions. In this research, the objects of n-gram analysis are nucleotide sequences that correspond to amino-acid sequences in proteins, so they belong to D , O or N regions only.

### 2.2 Repeats

Repeats can be considered as a special type of n-grams. Various kinds of repeats can be defined based on underlying n-gram characteristics. The following definition of repeats is taken from [18]:

Definition 2: Let $A=\{a, b, c, d, \ldots\}$ denote an alphabet with arbitrary symbols and $\mathrm{L}=\left\{\mathrm{l}_{1}, \mathrm{l}_{2}, \ldots, \mathrm{l}_{\mathrm{n}}\right\}$ is a language over alphabet $A$ which includes strings over $A$ with an arbitrary length, including empty string, and let $|\mathrm{s}|$ denote length of string $s \in L$, which is equal to the number of symbols (letters) from alphabet $A$.

An ordered triplet ( $x, \mathrm{~s}, p_{x}$ ) denotes a substring $x \in L$ of string $s \in L$ at the position $p_{x} \geq 1$ if $\exists y, z \in L: s=y x z \wedge|s|=|x|+|y|+|z| \wedge|x| \geq 1$. where $|y|=p_{x}$

Let the following functions be defined as:
(a) $f: L \rightarrow L \quad f(x)=z, \quad$ if $|x|=1 \quad$ for some $z \in A$
$f\left(x_{1}\right) f\left(x_{2}\right), \quad$ if $x=x_{1} x_{2} \in L \wedge|x|>1$
(b) $g: L \rightarrow L$

$$
\begin{aligned}
& g(x y)=y x \\
& g(x y)=y g(x) \\
& g(x y)=g(y) x \\
& g(x y)=g(y) g(x)
\end{aligned}
$$

if $|x|=1 \wedge|y|=1$
if $|x|>1 \wedge|y|=1$
if $|x|=1 \wedge|y|>1$
otherwise
then, for all string $s \in L$ the following four types of repeats can be defined (Figure 8):

1) The substring pair ( $a, \mathrm{~s}, p_{a}$ ) and (b,s, $\mathrm{p}_{\mathrm{b}}$ ) is a direct non-complementary repeat $(D N)$ if and only if $a=b \wedge p_{a}<p_{b}$
2) The substring pair ( $a, s, p_{a}$ ) and (b,s, $\mathrm{p}_{\mathrm{b}}$ ) is a inverse non-complementary repeat (IN) if and only if $a=g(b) \wedge p_{a} \leq p_{b}$
3) The substring pair ( $a, s, p_{a}$ ) and (b,s, $\mathrm{p}_{\mathrm{b}}$ ) is a direct complementary repeat ( $D C$ ) if and only if $a=f(b) \wedge p_{a}<p_{b}$
4) The substring ( $a, \mathrm{~s}, p_{a}$ ) and (b,s, $\mathrm{p}_{\mathrm{b}}$ ) is a inverse complementary repeat (IC) if and only if $a=f(g(b))=g(f(b)) \wedge p_{a} \leq p_{b}$

a) direct non-complementary

c) direct complementary

b) inverse non-complementary

d) inverse complementary

Figure 8. Graphical presentation of repeat types. In the examples, $f(x)=l, f(y)=k, f(z)=g$ is used for complementary mapping.

Extracting repeats from protein sequences is done using StatRepeats program [18]. Two different alphabets were used:

- $A=\{A, C, G, T\}$, when extracting repeats from (protein) nucleotide sequences, and
- $A=\{A, C, D, E, F, G, H, I, K, L, M, N, P, Q, R, S, T, V, W, Y, U, O\}$, when extracting repeats from (protein) amino-acids sequences.
By using nucleotide complementary characteristics ( $\mathrm{A} \leftrightarrow \mathrm{T}, \mathrm{G} \leftrightarrow \mathrm{C}$ ) in definition of functions $f$ and $g$, it is possible to obtain all four types of repeats for nucleotide sequences. For amino-acid sequences only non-complementary repeats are correct. Although, on first sight, looks that, except for monograms, set of direct noncomplementary repeats is equal to set of n-grams, this is not correct because StatRepeats extracts maximal repeats (i.e. repeats that not belongs to longer one). On the other side, StatRepeats can extract all (maximal) repeats or just subset of statistically significant repeats which can be used for additional checking of results.


### 2.3 Mole Fractions and fractional difference

Mole fractions are one way of representing the concentrations of the various chemical elements. In chemistry, mole fraction $x$ is a way of expressing the composition of a mixture. The mole fraction $x_{i}$ of each component $i$ is defined as its amount of substance $k_{i}$ divided by the total amount of substance in the system, $k_{\text {sum }}$ :

$$
x_{i}=\frac{k_{i}}{k_{\text {sum }}} \text { where } k_{\text {sum }}=\sum_{i=1}^{N} k_{i}
$$

$k_{\text {sum }}$ is calculated over all components, including the solvent in the case of a chemical solution. Consequence of such definition is that the sum of all the mole fractions is equal to 1 .

$$
\sum_{i=1}^{N} x_{i}=\sum_{i=1}^{N} \frac{k_{i}}{k_{\text {sum }}}=\frac{\sum_{i=1}^{N} k_{i}}{k_{\text {sum }}}=1
$$

In our research the mole fractions of amino-acid and nucleotide $n$-grams in regions was used as additional method for discovering $n$-grams that characterize specific type of regions. Mole fraction of specific $n$-gram in some region is calculated as quotient of number of $n$-gram occurrences in region and region length.

As a measure for difference of occurrences the same n-gram in different regions, the fractional difference (FD) is used. Fractional difference of occurrences of n-gram ngr in region $\mathrm{reg}_{1}$ related to the region $\mathrm{reg}_{2}$ can be defined as

$$
\mathrm{FD}\left(\text { ngr, } \text { reg }_{1}, \text { reg }_{2}\right)=\left(x_{\text {ngr-reg } 1}-x_{\text {ngr-reg } 2}\right) / x_{\text {ngr-reg } 2}
$$

where $x_{\text {ngr-regi }}$ denotes mole fraction of the n-gram ngr in the region regi. Thus a negative value for FD indicates a poorer concentration of n-gram ngr in the region $\mathrm{reg}_{1}$, while a positive value of FD indicates a richer concentration of n-gram ngr in the region $r e g_{1}$ then in the region reg $_{2}$.

### 2.4 Z-score

A z-score (also known as z-value, standard score, or normal score) is a measure of the divergence of an individual experimental result from the most probable result, the mean. Z-Score is a statistical measurement of a score's relationship to the mean in a group of scores, and is expressed in terms of the number of standard deviations from the mean value. A Z-score of 0 denotes that the score is equal to the mean. A Z-score can also be positive or negative, indicating how many standard deviations it is above or below the mean [18]. Prerequisites for applying z -score test are normal (or approximately normal) distribution of data and existence of standard deviation. In general, z -values are calculated according to the following formula:

$$
z=\frac{X-\mu}{\sigma}
$$

where X is experimentally observed mean in N items, $\mu$ is the mean value, and $\sigma$ is the standard deviation.

Most statistical tests begin by identifying a null hypothesis. The Z score is a test of statistical significance that helps to decide whether or not to reject the null hypothesis. P-value, or probability value, is a statistical measure that also helps to decide if hypotheses are correct. It is directly related to the significance level, which is an important component in determining whether the data obtained from scientific research is statistically significant. In the other words, the p-value is the probability of incorrectly rejecting the null hypothesis. Z-score and p-value are connected. The judgment of rejecting the null hypothesis is often connected to some confidence levels. Typical confidence levels are $90 \%$, $95 \%$, or $99 \%$. A confidence level of $99 \%$ indicates that null hypothesis will not be rejected unless the probability that the pattern was created by random chance is less than a $1 \%$ probability. The Table 1 shows the critical p-values and z-scores for different confidence levels.

Table 1. Values of $\mathbf{z}$-score and p-value for some confidence levels

| z-score (standard deviations) | p-value (probability) | Confidence level |
| :---: | ---: | :---: |
| $<-1.65$ or $>+1.65$ | $<0.10$ | $90 \%$ |
| $<-1.96$ or $>+1.96$ | $<0.05$ | $95 \%$ |
| $<-2.58$ or $>+2.58$ | $<0.01$ | $99 \%$ |

In analysis the null hypothesis is that all $n$-grams have the (almost) similar number of occurrence in all region types. Because we work with n-grams (e.g. sequences), the meaning of p-value can be stated as the probability that at least one sequence will produce the same score by chance, while z -value for some n-gram measures how much standard deviations above the mean of the score distribution is number of its occurrences. In this research for evaluation of the results obtained from ngram extracted, the statistic z-score with p-value 0.01 has been used. Presumptions of normal distributed data and existence of standard deviations for $n$-grams hold. Z-value for n-gram $X=L_{1} L_{2} \ldots . L_{n}$ where $L_{i}$ denotes amino acid or nucleotide is calculated as following [29]:

$$
X_{Z}\left(L_{1} L_{2} \ldots L_{n}\right)=\frac{N\left(L_{1} L_{2} \ldots L_{n}\right)-\mu}{\sigma}
$$

where $N\left(L_{1} L_{2} \ldots L_{n}\right)$ denotes the number of occurrences of n -gram X . The mean value $\mu$ is equal to

$$
\mu=\frac{N\left(L_{1} L_{2} \ldots L_{n-1}\right) \times N\left(L_{2} \ldots L_{n}\right)}{N\left(L_{2} \ldots L_{n-1}\right)}
$$

and the standard deviation $\sigma$ is equal to

$$
\sigma=\frac{\sqrt{\mu} \times \sqrt{\left[N\left(L_{2} \ldots L_{n-1}\right)-N\left(L_{1} \ldots L_{n-1}\right)\right] \times\left[N\left(L_{2} \ldots L_{n-1}\right)-N\left(L_{2} \ldots L_{n}\right)\right]}}{N\left(L_{2} \ldots L_{n-1}\right)}
$$

### 2.5 Data mining techniques

Data mining is the process of extracting interesting information or patterns from large information store such as: relational database, data warehouses, XML repository, etc. Also data mining is known as one of the core processes of Knowledge Discovery in Database (KDD). There are various types of data mining techniques such as association rules, classifications and clustering, etc. In this research two methods were used: association rules and classification.

Classification is a data mining technique which uses input data to build classification model. Classification uses a learning algorithm to identify model that best fits the relationship between attribute set and class label of the input data [20, 21]. Direct application of classification (for example, tree based algorithm) on complete material used in this research do not bring satisfactory results. Quality of such model is between $50 \%$ and $60 \%$, which can not guarantee correct results of prediction. Instead of that classification was applied on parts of material (more precisely on groups of organisms that belong to the same family). Corrections and accuracy of model obtained can be measured with different measures, depends on applied classification algorithm. Detailed information about different classification algorithms and appropriate measures can be found in [20, 21, 22].

Association rules are relationships between seemingly unrelated data in a relational database or other information repository, with aim to extract interesting correlations [20, 21]. An association rule is an implication expression of the form of $\mathrm{X} \rightarrow \mathrm{Y}$, where X and Y are disjoint sets of items called itemsets. X is called the body (or the antecedent) of the rule, and $Y$ the head (or the consequent) of the rule.

There are two important basic measures for association rules quality, support (denoted as s) and confidence (denoted as c). Support is defined as the percentage/fraction of records that contain $\mathrm{X} \rightarrow \mathrm{Y}$ to the total number of records in the database. Support reflects frequency of a set of items. Confidence is defined as the percentage/fraction of the number of transactions that contain $\mathrm{X} \rightarrow \mathrm{Y}$ to the total number of records that contain X . Confidence is a measure of strength of the association rules, The higher the confidence and support, the rule is more significant [20, 21].

The formal definition of support and confidence are:

Support $\quad \mathrm{s}(\mathrm{X} \rightarrow \mathrm{Y})=\frac{\sigma(X \bigcup Y)}{N}$

Confidence $\quad c(X \rightarrow Y))=\frac{\sigma(X \cup Y)}{\sigma(X)}$
where $\sigma(\mathrm{X} \rightarrow \mathrm{Y})$ denotes number of occurrences of an item $\mathrm{X} \rightarrow \mathrm{Y}, \mathrm{N}$ is the total number of items and $\sigma(\mathrm{X})$ denotes number of occurrences of an item X .

Using support and confidence as a measure for quality of association rules in some cases can give wrong result [20]. The reason is the fact that the confidence ignores the support of the itemset appearing in the rule consequent. One way to overcome this pitfall is to use lift as a metric. Lift is evaluated as the ratio between rule's confidence and the support of the itemset in the rule consequent: Lift $=\mathrm{c}(\mathrm{X} \rightarrow \mathrm{Y}) / \mathrm{s}(\mathrm{Y})$

The lift is a value between 0 and infinity:
(a) A lift value greater than 1 indicates that the rule body and the rule head appear more often together than expected, which makes such rule interesting.
(b) A lift smaller than 1 indicates that the rule body and the rule head appear less often together than expected. This means that the occurrence of the rule body has a negative effect on the occurrence of the rule head. Such rule can be interesting as indicate absence of rule body constituents in the case of rule head occurring.
(c) A lift value near 1 indicates that the rule body and the rule head appear almost as often together as expected, so such rule will not be considered in this research.

In this research rule head can contain only two possible forms (including "order" and "disorder"). From this point of view association rules can be considered as auxiliary method for classification.

### 2.6 Disorder prediction

In this research the IUPred-long [10, 23], VSL2b [13, 24] and IsUnstruct [12, 25] predictors have been used for predicting ordered and disordered level for each protein in all dataset. Three predictors with different prediction algorithms have been used in order to minimize influence of prediction algorithm to results of prediction.

Disorder predictors are very complex programs. For example, architecture of VSL2b consists of three component predictors in two-level (VSL2B-M1 and VSL2B-M2) architectures (Figure 9). At the first level, there are two specialized predictors: a short disorder predictor, VSL2b-S, for disordered regions of $\leq 30$ residues, and a long disorder predictor, VSL2b-L, for disordered regions of $>30$ residues. At the second level, there is a metapredictor that combines outputs of the two specialized predictors into the final prediction. All component predictors are built as binary classifiers that approximate the posterior class probability $\mathrm{p}(\mathrm{c}=1 \mid \mathrm{x})$, where x is the feature (input) vector and c is the class label [24].
A. VSL2-MI

B. VSL2-M2


Figure 9. VSL2b predictor architectures (taken from [24])

### 2.6.1 IUPred predictor

IUPred assumes that globular proteins have larger numbers of effective interresidue interactions (negative free energy) than disordered proteins due to the different types of amino acids involved in possible residue contacts. The core of IUPred is a method that enables the direct estimation of the interaction energies using the protein sequence alone. The estimated energy for each residue depends on the amino acid type but also on the amino acid composition in the neighbourhood. Generally, residues with less favourable predicted energies are more likely to be disordered [10].

The IUPred server takes a single amino acid sequence as an input and calculates the pairwise energy profile along the sequence. The energy values are then transformed into a probabilistic score ranging from 0 (complete order) to 1 (complete disorder). Residues with a score above 0.5 can be regarded as disordered. Optional is the prediction of long disorder, short disorder, and structured domains, each using slightly different parameters. The main profile is to predict context-independent global disorder that encompasses at least 30 consecutive residues of predicted disorder [23].

### 2.6.2 VSL2b predictor

VSL2b predictor is a combination of neural network predictors for both short and long disordered regions. It marks residues of length at least 30 as long disordered regions; otherwise regions are marked as short. Each individual predictor is trained by the dataset containing sequences of that specific length. The final prediction is a weighted average determined by a second layer predictor. VSL2b applies not only the sequence profile, but also the result of sequence alignments from PSI-blast and secondary structure prediction from PHD and PSI-pred.

### 2.6.3 IsUnstruct predictor

IsUnstruct is program based on the Ising model for prediction of disordered residues from protein sequence. IsUnstruct searches not only for disordered regions but also for individual disordered residues in a protein chain. It takes an amino acid sequence in the FASTA format as an input and calculates probabilities for each residue. A residue is considered as disordered if the probability is larger than 0.5. In IsUnstruct, the interaction term between neighbours has been replaced by a penalty for a state change (the energy of border). This allows applying dynamic programming to the Ising problem.

The energy of each residue in one state or the other depends on the type of residue in our model. To estimate the energy of any state we introduce the energy of the border between ordered and disordered residues and the energies of initiation of disordered state at the ends [12, 25]. The energy of the $j$-th state of a protein chain is calculated according the following formula:
$E_{j}=\sum_{i=1}^{L} \omega\left(a_{i}, s_{i j}\right)+k_{j} . \omega_{g}+\delta_{N, j} \cdot \omega_{N}+\delta_{C, j} . \omega_{C}$
where $a_{i}$ is the type of amino acid residue, $s_{i j}$ describes the state of the $i$ residue in the $j$ conformation ( 1 in the case of disordered residue and 0 in the case of ordered state), $\omega_{g}$ is the energy of border, $k_{j}$ is the number of borders between ordered and disordered residues in the $j$ conformation, $\omega_{N}, \omega_{c}$ are the energies of initiation of disordered state at the ends, and $\delta_{N j}, \delta_{C, j}$ are equal to 0 if the corresponding terminal residue is in the ordered state and to 1 in the opposite case, and $L$ is the length of protein chain.

### 2.7 Model for determining region-characteristic n-grams in proteins

The basic idea for model construction is a very simple but effective: to combine the results of previously described methods. Using the n-gram analysis, repeat analysis and $z$-score technique we determine sets $S_{n}, S_{r}$, and $S_{z}$ of $n$-grams which have, in some region, peak values (for example, the number of occurrences) either below or above
mean value of other n-grams. Additionally, set of n-grams $\mathrm{S}_{\mathrm{FD}}$ is defined based on fractional difference. Finally, applying association rule mining methods on the sets $\mathrm{S}_{\mathrm{n}}$ and $\mathrm{S}_{\mathrm{rz}}$ (intersection of the sets $\mathrm{S}_{\mathrm{r}}$, and $\mathrm{S}_{\mathrm{z}}$ ), the additional set $\mathrm{S}_{\mathrm{AR}}$ will be obtained. Appropriate quality depends on the following factors:

1) Confidence. Only rules that have confidence of at least $50 \%$ can provide support for determining n -grams that characterize regions in general. If intention is to find some n-grams that are close to "absolute" (>50\%) confidence in the set of three possible values ('O', 'D', 'N) association rules with confidence lower than $50 \%$ can be searched in material ${ }^{1}$.
2) Support. Only rules with sufficient support will be taken. Sufficient support depends on body-n-gram length and n-gram constituents and is equal to the probability of the n-gram occurrence. The initial probability ('weight') for monograms (individual AA) is equal to the probability of occurrence of AA in the analyzed material. Probability for single AA occurrence in some region(s) is ${ }^{2}$

$$
w_{A A}=x_{A A}=\frac{n_{A A}}{\text { reg_len }} \text { where } \sum_{i=1}^{20} w_{i(A A)}=1
$$

where $\mathbf{w}_{\mathrm{AA}}$ denotes probability ('weight') of AA. In calculation probability for $n$-grams ( $n \geq 2$ ) the model assumed that $n$-gram constituents are independent. Thus, probability for the $n$-gram of length $n$ is equal to the product of probabilities of its monograms. For the $n$-gram $i$ in some region the probability of its occurrence is

$$
w_{i}=\prod_{j=1}^{n} w_{A A j}
$$

where $w_{A A j}$ denotes probability of the AA in the $j$-th position in the $n$-gram $i$. If for specific n-gram calculated probability is lower than support obtained from association rule mining where this n-gram occurs in the body of the rule, then such rule is preserved, otherwise rejected. Association rules selected in this

[^0]process give information that some dependency between region type and n-gram in specific region exists.
3) Lift. Only rules with lift $\geq 1.05$ or $\operatorname{lift} \leq 0.95$ are considered [20].
4) Only rules with 'unique' both left and right sides are considered. 'Unique' means that do not exist two or more rules with the same body that cover all types of regions. For example, none of the rules $A B C \rightarrow D$ and $A B C \rightarrow O$ is considered if both are suggested. Using threshold of $50 \%$ for confidence automatically reject all such rules.
5) Rules with body that is extension of the body of some other rule are rejected. For example, rule $A B C \rightarrow R$ is rejected if exist rule $B \rightarrow R$ with similar support, confidence and lift.

Sets $\mathrm{S}_{\mathrm{FD}}, \mathrm{S}_{\mathrm{z}}$ and $\mathrm{S}_{\mathrm{AR}}$ are determined for each type of region. Their intersection will give the set $S$ which include $n$-grams that characterize regions type. N-grams are characteristic n-grams for such region type if they:

- are rare or frequent in this type of region (from FD)
- have very high confidence (from Z-score), and
- their statistically significant occurrence (from association rules mining or from statistically significant repeats) is connected only to specific type of region.

Although n-grams have been already used in research for finding some genome characteristics [26, 27, 28, 29], the presented approach is new and original and, according to available literature in the time of doing research described in this thesis, not previously used for determining characteristic regions in protein.

## 3 Material

Viral genomes material used in this research was downloaded from NCBI site: ftp://ftp.ncbi.nlm.nih.gov/genomes/Viruses/. Material includes amino acids and nucleotide sequences of viral proteins and, among others, taxonomic information. During research, different versions of data have been used. New versions commonly represent extension of the old ones with addition on several new genomes. Corrections of obtained results have been checked on sets of such new genomes. Results presented in this thesis are produced on data downloaded at January 2017.

After downloading, data were passed through the process of checking and cleansing. Incomplete and duplicate genomes and their proteins have been removed as well as individual proteins with some failure or incompatibility (for example, proteins with non-continual code, proteins with different length of amino acid and nucleotide codes, etc.). In order to eliminate influence of possible noise and outliers, classes with small number $(<10)$ of genomes were eliminated from further processing. Finally the set of 190626 proteins is used as research material. Proteins are sourced from 4076 viruses which belong to 8 phyla and 31 different classes. Proteins in selected sets were coded with two translation tables ${ }^{3}$ : 11 (190493 proteins) and 4 (133 proteins). As these translation tables differ only in TGA nucleotide triplet (coded as stop codon in translation table 11 and amino acid W (Tryptophan) in the translation table 4), and because only 583 W amino acid (coded with TGA or TGG) exists in the proteins that have translation table 4 (which is $0.09 \%$ of total occurrence of amino acid W in the dataset), all proteins in the dataset are considered as they have translation table 11.

For additional verification of obtained results the proteins from DisProt database (http://www.disprot.org/) have been used. Total of 803 proteins with 2167 disorder regions was used from DisProt (Version 7.03, September 2016). DisProt database includes proteins with experimentally verified disordered regions. Because there is no guarantee that the rest of the protein (not belongs to verified disordered region) is completely order, data from DisProt database can be used primarily for verifying

[^1]characteristics related to disordered regions ${ }^{4}$, while verification related to ordered regions can be taken with some caution ${ }^{5}$.

### 3.1 Determining threshold for n-grams

Observing individual n-gram mole fractions can not give satisfactory (strong) prediction of n-grams which characterize (dis)order regions. The reason is meaning of mole fraction (concentration of object, i.e. n-gram) which can not adequately cover object probability of occurrences and its uniqueness or majority. For example, hypothetically, if some n-gram $\mathbf{N}$ occurs once in ordered region(s) and ten times in disordered region(s), and if length of these disordered regions is 9.99 times larger than length of the ordered region, then mole fraction of $\mathbf{N}$ is lower in disordered than in ordered region. On the other side, this single occurrence of $\mathbf{N}$ in ordered region can have smaller probability than (single) occurrence o $\mathbf{N}$ in disordered region. The similar situation is with fractional difference (in previous example $\mathbf{F D}(\mathbf{N})_{\text {D_o }}$ will be negative but close to zero, so it can be concluded that $\mathbf{N}$ can not characterize neither ordered nor disordered region). Additionally, large number of n-grams with small numbers of occurrences produce a noise that, although not affect the results, can make data mining process significantly slower. Parameters related to material used in the research are presented in Table 2 (number of AA n-grams) and Table 3 (number of regions, in material and in DisProt).

Table 2. Number of AA n-grams in material

| N-gram length | Total number of AA n-grams <br> in material | Number of unique AA n-grams in material |  |
| ---: | ---: | ---: | ---: |
|  | Present |  |  |
| 1 | $46,413,638$ | 20 | Missing |
| 2 | $46,223,012$ | 400 | 0 |
| 3 | $46,032,386$ | 8,000 | 0 |
| 4 | $45,841,760$ | 159,988 | 12 |
| 5 | $45,651,134$ | $2,886,848$ | 313,152 |
| 6 | $45,460,508$ | $17,821,832$ | $46,178,168$ |
| 7 | $45,269,882$ | $27,652,049$ | $1,252,347,951$ |

[^2]| 8 | $45,079,256$ | $29,360,800$ | $25,570,639,200$ |
| ---: | ---: | ---: | ---: |
| 9 | $44,888,630$ | $29,907,712$ | $511,970,092,288$ |
| 10 | $44,698,004$ | $30,278,252$ | $10,239,969,721,748$ |

Table 3. Number of regions in material
For length $>20$ average protein length (AA) is shown in brackets

| Region |  | Number of regions |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Type | Length | DisProt | IUPred-L | IsUnstruct | VSL2b |
|  | 1 | 1 | 230.615 | 20.424 | 60.776 |
|  | 2 |  | 118.577 | 39.258 | 49.362 |
|  | 3 |  | 67.201 | 47.393 | 60.373 |
|  | 4 |  | 42.984 | 44.031 | 82.299 |
|  | 5 | 31 | 31.454 | 40.605 | 104.052 |
|  | 6 | 38 | 24.789 | 35.228 | 90.843 |
|  | 7 | 40 | 18.406 | 31.011 | 70.737 |
|  | 8 | 29 | 14.730 | 27.388 | 55.485 |
|  | 9 | 26 | 12.197 | 24.160 | 42.595 |
|  | 10 | 32 | 10.191 | 21.030 | 32.208 |
|  | 11 | 38 | 8.595 | 19.158 | 27.730 |
|  | 12 | 36 | 7.780 | 17.315 | 23.215 |
|  | 13 | 19 | 6.936 | 15.847 | 19.227 |
|  | 14 | 28 | 6.227 | 14.440 | 16.171 |
|  | 15 | 27 | 5.472 | 13.197 | 13.640 |
|  | 16 | 29 | 5.224 | 12.216 | 12.027 |
|  | 17 | 23 | 4.367 | 10.901 | 10.527 |
|  | 18 | 15 | 4.122 | 10.248 | 9.319 |
|  | 19 | 15 | 3.908 | 9.241 | 7.847 |
|  | 20 | 20 | 3.534 | 8.430 | 7.259 |
|  | >20 | 761 [114,59] | 60.842 [47,77] | 140.415 [44,86] | 141.647 [53,83] |
|  | 1 | 27 | 140.217 | 12.015 | 30.914 |
|  | 2 | 5 | 76.420 | 7.012 | 15.230 |
|  | 3 | 8 | 45.108 | 5.947 | 10.497 |
|  | 4 | 8 | 29.669 | 5.331 | 10.082 |
|  | 5 | 5 | 23.450 | 5.435 | 11.158 |
|  | 6 | 3 | 19.612 | 5.113 | 10.969 |
|  | 7 | 10 | 15.517 | 4.876 | 11.318 |
|  | 8 | 11 | 13.105 | 4.591 | 12.140 |
|  | 9 | 7 | 11.163 | 4.557 | 12.232 |
|  | 10 | 8 | 9.321 | 4.095 | 11.905 |
|  | 11 | 5 | 8.538 | 4.090 | 12.497 |
|  | 12 | 12 | 7.863 | 3.818 | 12.529 |
|  | 13 | 8 | 7.489 | 3.882 | 11.763 |
|  | 14 | 3 | 6.845 | 3.640 | 11.885 |
|  | 15 | 7 | 6.203 | 3.719 | 12.226 |
|  | 16 | 15 | 5.498 | 3.615 | 11.884 |
|  | 17 | 9 | 4.898 | 3.631 | 11.253 |
|  | 18 | 12 | 4.757 | 3.387 | 10.921 |
|  | 19 | 14 | 4.408 | 3.408 | 10.327 |
|  | 20 | 15 | 3.869 | 3.341 | 9.616 |
|  | >20 | 1.148 [272,06] | 368.846 [105,72] | 315.851 [112,86] | 508.965 [60,69] |

That problem can be alleviated by observing only those n-grams that appear (in disordered or ordered regions) more times than a predefined threshold. Threshold must be defined to eliminate n-grams with very small probability (i.e. can occur by chance). Also, threshold must not be too strong to eliminate n-grams that include possibly important information. Based on n-grams distribution show in Table 4, the following rule is used to define threshold: All n-grams that appear once in the complete material will not be taken into account in the research.

Although in literature [26, 27, 28, 29] was found that AA n-grams with length less than four can not be used to give precise characterization, because the threshold is weak, all monograms, bigrams, trigrams and almost all tetragrams will be used in the research, while the number of eliminated n-grams increase (up to $55 \%$ for n-grams with length 10) as increase their length. This is especially important for data mining, because it decrease the number of different objects (here n-grams) used in the mining process. Regardless those n-grams which appear exactly twice can also be considered as object with low probability that holds approximately $6 \%$ of the material for longer ones, they are not eliminated from the research. One of the reason was that such (pair of) n-grams represents direct non-complementary repeats. The same principle is also applied on nucleotide n-grams. Nucleotide n-grams are calculated from length 1 up to the length of 30 which corresponds to the AA n-grams of length 10 . Also, nucleotide n-grams that appear only once are eliminated from research (percents are similar to the percents in the case of AA n-grams. For example, from initial 133712762 n-grams of length 30, after eliminating 91903558 n-grams that appear only once (about 69\%) in research remain 41809204 n-grams).

Table 4. Threshold for AA n-grams and percentage of eliminated n-grams

| N -gram length | Number of n-grams |  |  | Percentage related to total number of n-grams |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  | Total | Appear once | Appear twice | N -grams that appear once | N -grams appear less than three times |
| 1 | 46,413,638 | 0 | 0 | 0\% | 0\% |
| 2 | 46,223,012 | 0 | 0 | 0\% | 0\% |
| 3 | 46,032,386 | 0 | 0 | 0\% | 0\% |
| 4 | 45,841,760 | 33 | 61 | 0\% | 0\% |
| 5 | 45,651,134 | 280,445 | 237,269 | 0.61\% | 1.13\% |
| 6 | 45,460,508 | 9,256,115 | 3,531,199 | 20.36\% | 28.12\% |
| 7 | 45,269,882 | 20,955,158 | 3,519,377 | 46.28\% | 54.06\% |


| 8 | $45,079,256$ | $23,357,355$ | $3,155,572$ | $51.81 \%$ | $58.81 \%$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 9 | $44,888,630$ | $24,082,091$ | $3,093,361$ | $53.64 \%$ | $60.53 \%$ |
| 10 | $44,698,004$ | $24,575,343$ | $3,060,423$ | $54.98 \%$ | $61.82 \%$ |

### 3.2 Repeats and data mining

Because set of direct non-complementary repeats is equal to set of n-grams (that appear at least twice), by nature, and complementary repeats are not applicable to AAs, only inverse non-complementary repeats are determined for protein AA codes. For protein nucleotide codes inverse non-complementary, direct complementary and inverse non-complementary repeats are determined. Repeats are calculated in two versions - all repeats and statistically significant repeats.

For data mining application (classification) both amino acids and nucleotide ngrams and repeats were divided into two parts: model and test. Proteins from each phylum was divided related to their number and length in proportion belongs to [68, 72] interval for model and [28,32] for test. In the cases where proteins could not be divided according to both criteria, a division with a weaker proportion ([65, 75] for model and [25, 35] for test) was used. Distribution of proteins over groups and their phyla are shown in Table A3 in Appendix.

Number of determined amino-acids and nucleotide repeats from the used material and amino-acids repeats from DisProt are shown on Table 5. Determination of nucleotide repeats started with length 6 which corresponds to 2 AAs. It is interesting that numbers of all repeats and statistically significant repeats are the same for the lengths greater than 7 for AA repeats and 15 for nucleotide repeats ( 16 for in nucleotide repeats). Direct complementary repeats were not determined because they are included in the set of already determined n-grams.

## Table 5. Determined amino-acid and nucleotide repeats

Legend: in - inverse non-complementary repeats
ic - inverse complementary repeats
dc - direct complementary repeats
all - all repeats
ssr - statistically significant repeats

| Repeat length |  | Amino acids repeats |  | Nucleotide repeats |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | in |  | dc |  | ic |  | in |  |
|  |  | all | ssr | all | ssr | all | ssr | all | ssr |
| 2 | disprot <br> model <br> test | $\begin{gathered} \hline 3,734,384 \\ 37,582,276 \\ 16,706,373 \end{gathered}$ | $\begin{gathered} \hline 3,536,416 \\ 24,616,388 \\ 10,903,875 \end{gathered}$ |  |  |  |  |  |  |
| 3 | disprot <br> model <br> test | $\begin{gathered} 327,782 \\ 4,523,944 \\ 2,005,362 \end{gathered}$ | $\begin{gathered} 272,105 \\ 2,620,175 \\ 1,158,337 \end{gathered}$ |  |  |  |  |  |  |
| 4 | disprot <br> model <br> test | $\begin{gathered} 27,926 \\ 377,681 \\ 164,839 \end{gathered}$ | $\begin{gathered} 25,326 \\ 322,862 \\ 141,687 \end{gathered}$ |  |  |  |  |  |  |
| 5 | disprot <br> model <br> test | $\begin{gathered} 6,387 \\ 195,007 \\ 86,328 \end{gathered}$ | $\begin{gathered} 6,008 \\ 154,603 \\ 68,538 \end{gathered}$ |  |  |  |  |  |  |
| 6 | model <br> test <br> disprot | $\begin{gathered} \hline 24,500 \\ 10,231 \\ 985 \end{gathered}$ | $\begin{gathered} \hline 24,412 \\ 10,185 \\ 982 \end{gathered}$ | $\begin{aligned} & \hline 7,536,950 \\ & 7,536,950 \end{aligned}$ | $\begin{aligned} & \hline 3,963,612 \\ & 3,963,612 \end{aligned}$ | $\begin{aligned} & 8,639,587 \\ & 8,639,587 \end{aligned}$ | $\begin{aligned} & 4,737,143 \\ & 4,737,143 \end{aligned}$ | $\begin{aligned} & 9,220,206 \\ & 9,220,206 \end{aligned}$ | $\begin{aligned} & \hline 5,302,808 \\ & 5,302,808 \end{aligned}$ |
| 7 | model test disprot | $\begin{gathered} 22,884 \\ 8,966 \\ 604 \end{gathered}$ | $\begin{gathered} 22,817 \\ 8,937 \\ 604 \end{gathered}$ | $\begin{aligned} & 2,108,140 \\ & 2,108,140 \end{aligned}$ | $\begin{aligned} & 1,200,145 \\ & 1,200,145 \end{aligned}$ | $\begin{aligned} & 2,324,517 \\ & 2,324,517 \end{aligned}$ | $\begin{aligned} & 1,480,539 \\ & 1,480,539 \end{aligned}$ | $\begin{aligned} & 3,098,382 \\ & 3,098,382 \end{aligned}$ | $\begin{aligned} & 1,820,441 \\ & 1,820,441 \end{aligned}$ |
| 8 | model test disprot | $\begin{gathered} 5,254 \\ 2,021 \\ 196 \end{gathered}$ | $\begin{gathered} 5,254 \\ 2,021 \\ 196 \end{gathered}$ | $\begin{aligned} & 594,551 \\ & 594,551 \end{aligned}$ | $\begin{aligned} & 397,164 \\ & 397,164 \end{aligned}$ | $\begin{aligned} & 807,787 \\ & 807,787 \end{aligned}$ | $\begin{aligned} & 556,359 \\ & 556,359 \end{aligned}$ | $\begin{aligned} & 885,763 \\ & 885,763 \end{aligned}$ | $\begin{aligned} & 638,963 \\ & 638,963 \end{aligned}$ |
| 9 | model <br> test <br> disprot | $\begin{gathered} \hline 4,464 \\ 1,761 \\ 218 \end{gathered}$ | $\begin{gathered} \hline 4,464 \\ 1,761 \\ 218 \end{gathered}$ | $\begin{aligned} & 338,002 \\ & 165,073 \end{aligned}$ | $\begin{aligned} & 232,807 \\ & 118,026 \end{aligned}$ | $\begin{aligned} & 394,544 \\ & 191,207 \end{aligned}$ | $\begin{aligned} & 312,527 \\ & 156,137 \end{aligned}$ | $\begin{aligned} & 789,931 \\ & 380,354 \end{aligned}$ | $\begin{aligned} & 510,954 \\ & 261,928 \end{aligned}$ |
| 10 | model <br> test <br> disprot | $\begin{gathered} \hline 1,285 \\ 525 \\ 74 \end{gathered}$ | $\begin{gathered} \hline 1,285 \\ 525 \\ 74 \end{gathered}$ | $\begin{aligned} & \hline 96,345 \\ & 48,652 \end{aligned}$ | $\begin{aligned} & 73,336 \\ & 38,456 \end{aligned}$ | $\begin{gathered} 194,610 \\ 94,035 \end{gathered}$ | $\begin{gathered} 132,360 \\ 66,837 \end{gathered}$ | $\begin{aligned} & 242,117 \\ & 117,226 \end{aligned}$ | $\begin{gathered} 180,646 \\ 90,903 \end{gathered}$ |
| 11 | model <br> test <br> disprot | $\begin{gathered} 8,314 \\ 3,033 \\ 612 \end{gathered}$ | $\begin{gathered} 8,314 \\ 3,033 \\ 612 \end{gathered}$ | $\begin{aligned} & 28,645 \\ & 14,566 \end{aligned}$ | $\begin{aligned} & 24,284 \\ & 12,525 \end{aligned}$ | $\begin{aligned} & \hline 34,828 \\ & 17,153 \end{aligned}$ | $\begin{aligned} & 30,926 \\ & 15,488 \end{aligned}$ | $\begin{gathered} 143,070 \\ 67,466 \end{gathered}$ | $\begin{aligned} & 98,384 \\ & 48,235 \end{aligned}$ |
| 12 | model test |  |  | $\begin{aligned} & 8,527 \\ & 4,291 \end{aligned}$ | $\begin{aligned} & 7,853 \\ & 4,014 \end{aligned}$ | $\begin{aligned} & 32,305 \\ & 15,863 \end{aligned}$ | $\begin{aligned} & 27,267 \\ & 13,577 \end{aligned}$ | $\begin{aligned} & 44,917 \\ & 21,928 \end{aligned}$ | $\begin{aligned} & \hline 39,507 \\ & 19,607 \end{aligned}$ |
| 13 | model test |  |  | $\begin{aligned} & 2,533 \\ & 1,314 \end{aligned}$ | $\begin{aligned} & 2,470 \\ & 1,278 \end{aligned}$ | $\begin{aligned} & 3,410 \\ & 1,691 \end{aligned}$ | $\begin{aligned} & 3,362 \\ & 1,662 \end{aligned}$ | $\begin{aligned} & \hline 36,610 \\ & 17,538 \end{aligned}$ | $\begin{aligned} & \hline 32,984 \\ & 15,886 \end{aligned}$ |
| 14 | model test |  |  | $\begin{aligned} & \hline 791 \\ & 430 \end{aligned}$ | $\begin{aligned} & 788 \\ & 428 \end{aligned}$ | $\begin{aligned} & 3,093 \\ & 3,680 \end{aligned}$ | $\begin{aligned} & 7,962 \\ & 3,607 \end{aligned}$ | $\begin{gathered} \hline 11,483 \\ 5,088 \end{gathered}$ | $\begin{gathered} \hline 11,329 \\ 5,020 \end{gathered}$ |
| 15 | model test |  |  | $\begin{aligned} & 262 \\ & 127 \end{aligned}$ | $\begin{aligned} & 262 \\ & 127 \end{aligned}$ | $\begin{aligned} & 415 \\ & 226 \end{aligned}$ | $\begin{aligned} & 415 \\ & 226 \end{aligned}$ | $\begin{gathered} 10,761 \\ 4,722 \end{gathered}$ | $\begin{gathered} 10,655 \\ 4,672 \end{gathered}$ |
| 16 | model test |  |  | $\begin{aligned} & 78 \\ & 40 \end{aligned}$ | $\begin{aligned} & 78 \\ & 40 \end{aligned}$ | $\begin{aligned} & 2,179 \\ & 1,108 \end{aligned}$ | $\begin{aligned} & 2,179 \\ & 1,108 \end{aligned}$ | $\begin{aligned} & 3,739 \\ & 1,629 \end{aligned}$ | $\begin{aligned} & 3,739 \\ & 1,629 \end{aligned}$ |
| 17 | model <br> test |  |  | $\begin{aligned} & 34 \\ & 13 \end{aligned}$ | $\begin{aligned} & 34 \\ & 13 \end{aligned}$ | $\begin{gathered} 105 \\ 33 \end{gathered}$ | $\begin{gathered} 105 \\ 33 \end{gathered}$ | $\begin{aligned} & 3,459 \\ & 1,536 \end{aligned}$ | $\begin{aligned} & 3,459 \\ & 1,536 \end{aligned}$ |


| 18 | model | 12 | 12 | 743 | 743 | 1,159 | 1,159 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | test | 9 | 9 | 367 | 367 | 773 | 773 |
| 19 | model | 12 | 12 | 17 | 17 | 1,187 | 1,187 |
|  | test | 7 | 7 | 8 | 8 | 588 | 588 |
| 20 | model | 1 | 1 | 329 | 329 | 589 | 589 |
|  | test | 2 | 2 | 146 | 146 | 202 | 202 |
| 21 | model | 1 | 1 | 4 | 4 | 510 | 510 |
|  | test | 6 | 6 | 9 | 9 | 246 | 246 |
| 22 | model |  |  | 123 | 123 | 191 | 191 |
|  | test |  |  | 55 | 55 | 91 | 91 |
| 23 | model | 2 | 2 | 12 | 12 | 320 | 320 |
|  | test | 1 | 1 | 5 | 5 | 133 | 133 |
| >23 | model | 3 | 3 | 170 | 170 | 1,258 | 1,258 |
|  | test |  |  | 91 | 91 | 485 | 485 |

## 4 Results

Results will be presented for each of the previously described methods and their combinations, and compared with corresponding DisProt data.

### 4.1 Mole fractions

### 4.1.1 Mole fractions of AA n-grams

It is not expected that mole fractions (especially for longer n-grams) can be used for finding n -grams that characterize either order or disorder regions. But, mole fractions can be good markers for compatibility of material used in research with material in DisProt version 7.03. Comparison of mole fractions for monograms in complete material and DisProt is presented on Figure 10.


Figure 10. Comparison of mole fraction in used material and DisProt database V7.03

In general, most amino acids have similar mole fractions in both series. Larger differences exists for amino Glutamic acid (E), Proline (P) and Serine (S) (higher level in DisProt), and Phenylalanine (F), Isoleucine (I), Asparagine (N) and Tyrosine (Y) (higher level in our material). Number of significant differences became even smaller if mole fractions are compared separately in predicted disordered with experimentally found disordered regions from DisProt (Figure 11), and in predicted ordered regions with non-disordered regions from DisProt (Figure 12).


Figure 11. Comparison of mole fraction in disordered regions of used material (predicted by disorder predictors) and disordered regions from DisProt database

Different disorder predictors predict different regions and consequently have different mole fractions for individual AAs. But, from the Figures 11 and 12 it is evident that content of AAs in the predicted regions (later used in the research) have very similar behaviour (related to dis/ordered regions) to material in DisProt.


Figure 12. Comparison of mole fraction in ordered regions of used material (predicted by disorder predictors) and non-disordered regions from DisProt database

### 4.1.2 Mole fractions of nucleotide n-grams

Because nucleotide sequences are three times longer then corresponding aminoacid sequences and nucleotide n-gram can start at arbitrary positions, set of nucleotide
n-grams not completely correspond to the set of amino-acids n-grams. More precisely, only a third of nucleotide n-grams have their equivalent amino acid n-grams. Mole fractions of individual n-grams also depend on codon usage. For this reason, in some analysis, set of nucleotide n-grams is divided in three parts, according to their starting positions (i.e. relative offset to a closest codon starting position, takes a value from 0 to 2). Because of these dependences which can not guarantee correctness in the case of generalization to other material, the obtained mole fractions were used just as a method for sorting n-grams according to their abundance in the material. Nevertheless, some interesting information related to nucleotide n-grams mole fractions were found in the material. Mole fractions related to monograms for all material and grouped according their starting positions ("ORF", open reading frame) are shown in Table 6.

Table 6. Mole fractions of nucleotide monograms in all material and grouped according to their starting positions

| ORF | n-gram | Mole fractions all | Mole fractions order | Mole fractions disorder |
| :---: | :---: | :---: | :---: | :---: |
| all | A | 0.286286328169 | 0.279469194991 | 0.311333252030 |
|  | C | 0.224269613742 | 0.217597454352 | 0.248783887397 |
|  | G | 0.244571161031 | 0.239782427068 | 0.262165515114 |
|  | T | 0.244871597151 | 0.263149516567 | 0.177716439107 |
| 1 | A | 0.296621113820 | 0.292812695421 | 0.310613676849 |
|  | C | 0.190309343990 | 0.182180318583 | 0.220176309346 |
|  | G | 0.332583474710 | 0.328734266434 | 0.346725904394 |
|  | T | 0.180485787388 | 0.196272390646 | 0.122484008703 |
| 2 | A | 0.330734492305 | 0.321099981484 | 0.366132782567 |
|  | C | 0.224520517008 | 0.209108422513 | 0.281146307835 |
|  | G | 0.167926784795 | 0.165927007687 | 0.175274193692 |
|  | T | 0.276818205890 | 0.303864588314 | 0.177446715904 |
| 3 | A | 0.231503378382 | 0.224494908069 | 0.257253296673 |
|  | C | 0.257978980229 | 0.261503621959 | 0.245029045009 |
|  | G | 0.233203223586 | 0.224686007081 | 0.264496447256 |
|  | T | 0.277310798175 | 0.289311570740 | 0.233218592713 |

We can observe some characteristics of the n-grams:

- There are only 180 non-ACGT nucleotides in complete material (mole fractions 0.000001292720 ), so such nucleotide codes were not considered as separate group
- Percentage of GC nucleotides is almost half (51.09\%) in complete material
- N -grams belong to $\mathrm{ORF}=3$ set (i.e. are on the third position in the AA codon) have similar GC percent (50.95\%); n-grams that belong to ORF=1 (first nucleotide in AA codon) are richer (56.69\%) while nucleotides in the middle of AA codons are poorer (45.64\%) in GC nucleotides
- As expected, n-grams that correspond to amino-acids homorepeats occur also in the nucleotide level. But, because individual amino acids have different corresponding codons at nucleotide level, the corresponding nucleotide repeats are not necessary homorepeats of appropriate trigrams. An interesting observation is that nucleotide sequences that are homorepeats or include homorepeats occur more often than sequences that are random sequences of (codon) trigrams. For example, on amino-acid level hexagram 'PPPPPP' occurs 1145 times in disorder regions. Corresponding nucleotide n-grams (for ORF=1) occurs in 285 variations; among them the most numerous groups are 'pure' homorepeats 'CCACCACCACCACCACCA' and 'CCGCCGCCGCCGCCGCCG' (each occurs 29 times), followed by tandem repeats ('ССТССАССТССАССТССА' - 18 times and 'ССАССТССАССТССАССТ'- -16 times, 'CCACCACCTCCACCACCT' - 8 times, 'CCGCCGCCACCGCCGCCG' - 8 times, 'CCACCACCTCCACCACCT' - 8 times, etc. This diversity of AAs translation into codons gives additional opportunity to more precisely describe characteristic n-grams.


### 4.2 Fractional difference

### 4.2.1 Fractional differences of AA n-grams

Fractional difference of some n -grams indicates their richer or poorer concentration in disordered or ordered regions. Fractional difference disorder/order of the n-gram $N$ with length $n$ (in the rest of the text $\mathrm{FD}_{\mathrm{n}}\left(\mathrm{d} \_0, \mathrm{~N}\right.$ ) is positive if disorder region is richer of this n-gram, and negative if disorder region is poorer in this n-gram (i.e. order region is richer). If fractional difference disorder/order of some n-gram is positive, than this n-gram characterize disordered region; as opposite it characterize ordered region. Fractional difference for monograms is shown on Figure 13 together
with fractional differences of the monograms in DisProt database (version V7.03, September 2016).


Figure 13. Comparison of disorder/order fractional difference of monograms from material used in research and material from DisProt database

There are some differences in predicted FD compared to the FD in DisProt material (VSL2b and IUPred-L have 4 and IsUnstruct 3 differences). FD1(d_o,T) from DisProt and FD1(d_o,T) predicted by IsUnstruct are very close to zero but with opposite signs, and such variations are expected. A little bit larger incompatibilities are for AAs Metionine ( M ) and Arginine ( R ) that are also close to the transition from positive to negative. It must be noticed that FD values for current version of DisProt differs from version 3.4 (shown of Figure 5) both in order of AAs related to FD and in orientation. For example, AAs M and R are disorder oriented in version 3.4, but order oriented in DisProt version 7.03. The reason can be the different percentage of AAs in ordered/disordered regions in later added proteins to DisProt database. Also the results from predictors better fits to FD in DisProt 3.6 than in 7.03. The probable reason for
that can be that the set of proteins used for predictors training which is more similar to set in version 3.6 than in 7.03.

From previous figures it can be seen that all three predictors produce similar results. For this reason, even though the calculation was done for all three predictors, in the further text the most of the results will be illustrated only for IsUnstruct predictor, while the results or other two will be presented only if there are large differences in results.

Number of n-grams that have positive FD(d_o) is shown on Table 7. Observing that only fractional differences can not give precise characterization of disordered or ordered regions because numbers are too high. Even if consider only those n-grams that occur only in disordered regions their number remains too high. Appendix table A4 contains list of some n-grams that occur only in disordered regions predicted by IsUnstruct predictor.

Table 7. Number of n-grams with positive $\mathrm{FD}(\mathrm{d}$ _o $)$ regions and their percentage in the sample

| N-gram length | Number of grams with positive FD(d_o) |  |  | Total number of n-grams | Percent of positive n-grams |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | VSL2b | IsUnstruct | IUPred-L |  | VSL2b | IsUnstruct | IUPred-L |
| 1 | 12 | 11 | 10 | 20 | 60.00\% | 55.00\% | 50.00\% |
| 2 | 160 | 156 | 154 | 400 | 40.00\% | 39.00\% | 38.50\% |
| 3 | 2,702 | 2,643 | 2,482 | 8,000 | 33.77\% | 33.03\% | 31.02\% |
| 4 | 51,234 | 48,278 | 45,588 | 159,955 | 32.03\% | 30.18\% | 28.50\% |
| 5 | 900,484 | 795,388 | 637,986 | 2,606,403 | 34.54\% | 30.51\% | 24.47\% |
| 6 | 3,064,465 | 2,427,476 | 1,436,112 | 8,565,717 | 35.77\% | 28.33\% | 16.76\% |
| 7 | 2,286,561 | 1,623,033 | 1,023,642 | 6,696,891 | 34.14\% | 24.23\% | 15.28\% |
| 8 | 2,076,342 | 1,416,707 | 948,566 | 6,003,445 | 34.58\% | 23.59\% | 15.80\% |
| 9 | 2,082,655 | 1,393,438 | 960,963 | 5,825,621 | 35.74\% | 23.91\% | 16.49\% |
| 10 | 2,111,453 | 1,391,255 | 982,448 | 5,702,909 | 37.02\% | 24.39\% | 17.22\% |

One criterion for selecting "better" n-grams (the ones that not only appear in disordered regions, but also have positive fractional difference in disordered regions), can be the position of such n-grams in the list of n-grams ordered according to their mole fractions in descending order. For each n-gram length, first 100 n -grams with the highest mole fractions are shown in Appendix Table A5. It is interesting that among the n-grams that occur only in predicted disordered regions the most of the n-grams include some kind of homorepeats [30], either partial or full (for example EEEEG, PPPSPPPS,

SSSSSSSS, etc), or a repeat structure (for example PAPAPA). These repeat structures also can be found in many of the $n$-grams that prefer disordered regions (for example EEE, DDD, PPPSP, etc., see Table A5) where such n-grams are included in longer ngrams that appear only in disordered regions. Similar tables are presented for characteristic n-grams in ordered regions (Appendix tables A6 and A7) and for borders between ordered and disordered regions (Appendix tables A8 an A9). Characteristic ngram of smaller length are combined with order promoted AAs (for example WIC, CYW, LCYL, VLYV, etc.) or rudimentary (homo)repeats (like YYVV or ILILL) but for longer n-grams no clear pattern can be observed except that trigram LLL appears as a part of various longer n-grams. Although there are a lot of n-grams in Tables A8 and A9, no clear pattern for border n-grams can be observed. Because of the huge number of n-grams in previously described sets (for example, set of n-grams of any length that appear only in disordered regions have cardinality of 3.5 M , while set of n-grams that have positive disorder fractional difference and appear not only in disordered regions have cardinality of 2.7 M ), additional restriction can be provided by increasing threshold for eliminating $n$-grams accepting the rule that $n$-grams with very small mole fractions will be removed from sets. This procedure will be used for sets produced as combination of different approaches.

As a verification of the method of using mole fraction and fractional differences for determining characteristic n-grams, a comparison with fractional differences of identical n-grams available from DisProt proteins was performed. The comparison results are presented in Table 8. Percentage of identical n-grams that belong to the same type of region grows up to $99.38 \%$ as n-gram length increase. Additional information about widespread of n-grams over proteins shows that the most of the n-grams appear in different proteins and in proteomes of different phyla and classes of viruses. different classes of viruses. For example, n-gram GGGGGGG belongs to 450 different proteins in material (from 3 phyla and 12 classes), and to 5 proteins from DisProt, n-gram GGGSGGG to 70 proteins (3 phyla, 9 classes) in material and 4 proteins from DisProt, etc.

### 4.2.2 Fractional differences of nucleotide n-grams

Nucleotide n-grams can also be ordered by the concentration in disordered or ordered regions. Fractional differences of nucleotide monograms divided into three sets
according to their starting positions ("ORF" on figure) are shown on Figure 14. Concentration is almost uniform regardless of ORF-s: nucleotide T has larger concentration in ordered regions while other nucleotides have large concentration in disordered regions with exception of C in ORF3.

Table 8. Number of matched regions according to fractional difference of AA n-grams that appear in predicted regions and regions from DisProt database.
Number of equal - number of n-grams available both in materials used in research and in DisProt
Number of matched - number of n-grams that belongs to the same type of region (comparing FD related to predicted regions and FD related to regions from DisProt)
Number of non-matched - number of n-grams that belongs to the opposite type of region (comparing FD related to predicted regions and FD related to regions from DisProt)
Matched/number of equal - percent of n-grams with matched FD related to total number of n-grams

| N-gram <br> length | Number of equal | Number of matched | Number of non-matched | matched/number of equal |
| :---: | ---: | ---: | ---: | ---: |
| 1 | 20 | 17 | 3 | $85.00 \%$ |
| 2 | 400 | 344 | 56 | $86.00 \%$ |
| 3 | 7,213 | 5,276 | 1,937 | $73.14 \%$ |
| 4 | 45,224 | 24,118 | 21,106 | $53.33 \%$ |
| 5 | 61,346 | 42,455 | 18,891 | $69.20 \%$ |
| 6 | 20,705 | 19,273 | 1,432 | $93.08 \%$ |
| 7 | 2,950 | 2,910 | 40 | $98.64 \%$ |
| 8 | 1,140 | 1,131 | 9 | $99.21 \%$ |
| 9 | 799 | 794 | 5 | $99.37 \%$ |
| 10 | 648 | 644 | 4 | $99.38 \%$ |



Figure 14. Fractional differences of nucleotide monograms grouped according to their starting positions. Ordered/disordered regions are predicted using IsUnstruct predictor.

Fractional differences of nucleotide trigrams can be used for determining if there are any difference related to ordered/disordered regions and AA codon usage. Figure 15 presents fractional differences of nucleotide trigrams ordered according AA codon usage (related to translation table 11). The most interesting are values for ORF1. Such differences exist for some AAs but these results are predictor depending and can not be generalized without further verification. For example, depending on predictor used order/disorder codon dependencies are

- VSL2b: amino acid A:
o GCT-order, all other codons - disorder
- IsUnstruct (shown on Figure 15):
o amino acid G: GGG-order, all other codons - disorder
o amino acid N : AAC-disorder AAT-order
o amino acid T: ACT-order, all other codons - disorder
- IUPred-L:
o amino acid H : CAC-disorder CAT-order
o amino acid K: AAA-order, AAG-disorder
o amino acid N : AAC-disorder AAT-order


### 4.3 Z-score

Z-score value is used as an additional confirmation if specific n-gram characterize ordered or disordered region. Z-score is calculated only for n-grams in disordered or ordered regions. It is not possible to calculate it for n-grams in N (border) regions because there is not guarantee for n-gram in border region that all its sub-ngrams also belong to the border region (which is necessary for z -score calculation). Also, some n-grams and their sub-n-grams can occur many times in proteins in the same type of region, but only once in any of (individual) proteins. Such n-grams have z-score equal to zero and do not satisfy any confidence level, despite they are potential markers for some type of region.

The most restrictive criterion for z -values is chosen by selecting n-grams with confidence level 99\% (see Table 1). The selection algorithm for n-gram $N$ and region with type $R$ (ordered, disordered) can be illustrated with the following pseudocode:



Figure 15. Fractional differences of nucleotide trigrams. Part one: amino acids A, C, D, E, F, G, H, I, K, L, M. Pat two: amino acids N, P, Q, R, S, T, V, W, Y. Ordered/disordered regions are predicted with IsUnstruct predictor.

```
for each n-gram N an region type R
    if exist z-score for N in type R regions
        then if abs(z-score)>2.58
            then if exist z-score for N in opposite-type regions
                        then if abs(z-score opposite-type)<1.65
                        then N characterize R type regions
                        else without-characterization
                        else without-characterization
            else N characterize R type regions (exclusive)
        else without-characterization
```

Applying the previous algorithm, n-grams that characterize only one type of region (requirement abs(z-score)>2.58) but not the opposite (requirement abs(zscore)<1.65) were selected. N -grams that characterize specific regions are shown in

Appendix tables A10 (ordered regions) and A11 (disordered regions). In both tables ngram patterns with similar structures (homorepeats and repeats) can be seen for disordered and ordered regions. As in the previous methods, for both ordered and disordered regions, numbers of selected n-grams have peak for length 6 and 7, and decrease as n-gram length increase or decrease (Table 9).

Table 9. Number of selected characteristic n-grams based on z-score values

| N-gram |  |  |  |
| ---: | ---: | ---: | :---: |
| length | number /disordered regions | number /ordered regions |  |
| 3 | 568 | 1532 |  |
| 4 | 12789 | 31699 |  |
| 5 | 126827 | 330136 |  |
| 6 | 641019 | 2215554 |  |
| 7 | 406744 | 5121903 |  |
| 8 | 61952 | 56721 |  |
| 9 | 8638 | 54022 |  |
| 10 | 2374 | 13331 |  |

### 4.4 Combination of fractional difference, z-score and mole fractions

### 4.4.1 Combination of Fractional difference and Mole fractions for AA n-grams

Numbers of significant n -grams decrease in a very small percent (about 2.65\%) if z -score method combined with method based on fractional difference. More significant reduction is obtained if combination includes fractional difference, z-score and n-grams with mole fractions larger than specific value (which is increasing threshold level, see section 3.1). Percentages of decreasing n-gram numbers depending of mole fractions are shown in Table 10.

Number of n-grams that characterize ordered regions is reduced much faster than number of n -grams that characterize disordered ones. This is caused by average number and standard deviation of n-gram occurrences which is both between 2 and 3 for n-gram length $>5$, but with lower average number and higher standard deviation of n-gram occurrences in disordered compared to ordered regions, which is especially emphasized
for n-gram lengths 6 and 7. Related to number of n-grams in Table 10 middle (but satisfactory) level of reducing is obtained by taking condition "mole fraction>1E-6". Ngrams that satisfy this condition are shown in Appendix Tables A12 (ordered regions) and A13 (disordered regions). Among the n-grams that characterize ordered regions, the same pattern as in the previous tables is observed (for example 'LLL'), but for disordered regions patterns are more uniform than in previous cases. On the top of the list for all n-gram lengths are homorepeats ('QQQ', 'SSSS', 'GGGG', 'PPPPP', 'EEEEEEE', etc.), tandem repeats ('APAP', 'SRSRSR', 'PEPEPE', 'AATTTAATTT', etc.) or palindromes ('APAPA', 'SDSDSDS', 'PKPAPKP', 'DEDDEDDED', etc.) or their shorter versions of disorder promoting AAs (see Figure 13) combined with some other AAs.

Table 10. Number of n-grams and percentage of initial n-grams for different mole fractions used
Initial n -gram number - n -grams that satisfy fractional difference and z -score conditions

| Mole fractions > |  |  | 5E-6 |  | 1E-6 |  | 5E-7 |  | 1E-7 |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Type | N-gram length | Initial ngram number | n-gram number | Percent of initial | n-gram number | Percent of initial | n-gram <br> number | Percent of initial | n-gram <br> number | Percent of initial |
| $\begin{aligned} & \stackrel{\rightharpoonup}{0} \\ & \text { on } \\ & \text { on } \end{aligned}$ | 3 | 242 | 242 | 100.00 | 242 | 100.00 | 242 | 100.00 | 242 | 100.00 |
|  | 4 | 7176 | 5678 | 79.1248 | 7102 | 98.9687 | 7163 | 99.8188 | 7176 | 100.00 |
|  | 5 | 108560 | 4471 | 4.1184 | 55374 | 51.0077 | 79551 | 73.2783 | 108560 | 100.00 |
|  | 6 | 635966 | 813 | 0.1278 | 73993 | 11.6347 | 216916 | 34.1081 | 635966 | 100.00 |
|  | 7 | 406634 | 423 | 0.1040 | 63583 | 15.6364 | 149604 | 36.7908 | 406634 | 100.00 |
|  | 8 | 61939 | 166 | 0.2680 | 20641 | 33.3247 | 36584 | 59.0645 | 61939 | 100.00 |
|  | 9 | 8633 | 89 | 1.0309 | 3590 | 41.5846 | 5822 | 67.4388 | 8633 | 100.00 |
|  | 10 | 2371 | 46 | 1.9401 | 1147 | 48.3762 | 1741 | 73.4289 | 2371 | 100.00 |
| 苞 | 3 | 1218 | 1216 | 99.8357 | 1218 | 100.00 | 1218 | 100.00 | 1218 | 100.00 |
|  | 4 | 24662 | 10579 | 42.8959 | 22100 | 89.6115 | 23789 | 96.4601 | 24620 | 99.8296 |
|  | 5 | 281140 | 333 | 0.1184 | 59787 | 21.2659 | 140114 | 49.8378 | 265399 | 94.4010 |
|  | 6 | 2144995 | 126 | 0.0058 | 7860 | 0.3664 | 78970 | 3.6815 | 1205359 | 56.1940 |
|  | 7 | 3081651 | 104 | 0.0033 | 4096 | 0.1329 | 45270 | 1.4690 | 1114921 | 36.1793 |
|  | 8 | 563499 | 33 | 0.0058 | 1484 | 0.2633 | 23994 | 4.2580 | 333600 | 59.2015 |
|  | 9 | 53794 | 12 | 0.0223 | 289 | 0.5372 | 3600 | 6.6921 | 40082 | 74.5101 |
|  | 10 | 13283 | 2 | 0.0150 | 103 | 0.7754 | 992 | 7.4681 | 9507 | 71.5726 |

N -grams determined under these conditions can be compared with n-grams generated from DisProt database proteins. The percents of agreement of predicted characteristic ngrams with corresponding $n$-grams in disordered and ordered regions are shown on Figure 16. Shorter n-grams (length<5) more precisely characterize ordered regions than disordered. For disordered regions, longer n-grams agreed with n-grams from DisProt
database in high (9-grams) or very high (other n-grams, $n>4, n \neq 9$ ) percent; for ordered regions pentagrams agreed in high percent while longer n-grams agreed in very high percent.

Based on these results, it is expected that also the data mining analysis will confirm that regions are more precisely characterized with longer n-grams. This expectation is in compliance with the results presented in the next chapter. Also, because using z -score values excludes set of n-grams that characterize border regions, for border regions the final results will be produced by intersecting sets obtained with fractional difference and mole fractions methods with set of n-grams produced with data mining.


Figure 16. Agreement in characterization regions with identical n-grams from used material and DisProt database. D - disordered regions; O - ordered regions. Ordered/disordered regions are predicted with IsUnstruct predictor.

### 4.4.2 Combination of Fractional difference and Mole fractions for nucleotide n-grams

Z-score values were not calculated for nucleotide n-grams. In addition to the previously mentioned disadvantage, it is not possible to calculate z -scores for nucleotide n-grams divided into ORF groups because sub-n-grams (necessary for z-score calculation) belong to different ORF group. If methods that include combination of fractional difference and mole fractions are applied on nucleotide n-grams, some interesting facts can be observed:

- Percentages of retain (initial) n-grams are comparable for all ORF-s; also number of n-grams have the same order of magnitude in all ORF-s for the same mole fractions restriction level. Table 11 presents how the number of n-grams is related to increasing mole fractions for $\mathrm{ORF}=1^{6}$.
- Number of order related n-grams decrease more rapidly compared to disorder ones, as mole fraction increase, regardless their number significantly exceed number of disordered related n-grams. This leads to conclusion that longer order related n-grams have a smaller cardinality of occurrences than longer disorder related n-grams, i.e. for all n-grams lengths exists some disordered related ngrams with sufficient number of occurrences that with high probability can be considered as markers for disordered regions.
- For each n-gram lengths some significant nucleotide n-grams exist in different ORF-s. Some of these n-grams (with length divided by 3) corresponds to AAs ngrams that are also among significant ones (for example, 'GGTCAGCACATTTCCATCCGA' with corresponding AA n-gram 'GQHISIR', 'AATCCAGCTCCGACGTCAAGTCCT' which correspond 'NPAPTSSP', etc).

[^3]Table 11. Number of nucleotide n-grams and percent of retain initial n-grams for different mole fractions used. N -grams belong to ORF=1 i.e. start on position correspond to AAs n-grams

| Mole fractions > |  |  | 5E-6 |  | 1E-6 |  | 5E-7 |  | 1E-7 |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Type |  | Initial ngram number | n-gram number | Percent of initial | n-gram number | Percent of initial | n-gram number | Percent of initial | n-gram number | Percent of initial |
| $\begin{aligned} & \ddot{ \pm} \\ & 000 \\ & 0 . \end{aligned}$ | 1 | 3 | 3 | 100.00 | 3 | 100.00 | 3 | 100.00 | 3 | 100.00 |
|  | 2 | 10 | 10 | 100.00 | 10 | 100.00 | 10 | 100.0 | 10 | 100.00 |
|  | 3 | 37 | 36 | 97,2972 | 36 | 97,2972 | 37 | 100.00 | 37 | 100.0 |
|  | 7 | 6.556 | 6.533 | 99,6491 | 6.533 | 99, 6491 | 6.533 | 99,6491 | 6.553 | 99,9542 |
|  | 8 | 24.842 | 23.968 | 96,4817 | 24.805 | 99,851 | 24.806 | 99,855 | 24.836 | 99,9758 |
|  | 9 | 99.374 | 49.350 | 49,6608 | 96.335 | 96,9418 | 98.963 | 99,5864 | 99.361 | 99,9869 |
|  | 13 | 3.913.917 | 662 | 0,0169 | 88.176 | 2,2528 | 340.739 | 8,7058 | 3.455.361 | 88,2839 |
|  | 14 | 3.196.310 | 402 | 0,0125 | 70.169 | 2,1953 | 270.353 | 8,4582 | 2.721.386 | 85,1414 |
|  | 15 | 2.320.596 | 254 | 0,0109 | 60.434 | 2,6042 | 229.094 | 9,8722 | 1.913.555 | 82,4596 |
|  | 16 | 1.831 .777 | 192 | 0,0104 | 53.916 | 2,9433 | 200.902 | 10,9676 | 1.377 .924 | 75,2233 |
|  | 17 | 1.645.754 | 169 | 0,0102 | 52.923 | 3,2157 | 195.910 | 11,9039 | 1.208.295 | 73,4189 |
|  | 18 | 1.560.665 | 127 | 0,0081 | 63.902 | 4,0945 | 186.066 | 11, 9222 | 1.130 .913 | 72,4635 |
|  | 19 | 1.545.559 | 106 | 0,0068 | 59.336 | 3,8391 | 172.069 | 11,1331 | 1.043.707 | 67,5294 |
|  | 20 | 1.535.807 | 105 | 0,0068 | 58.830 | 3,8305 | 170.658 | 11,1119 | 1.034.296 | 67,3454 |
|  | 21 | 1.513 .175 | 97 | 0,0064 | 56.494 | 3,7334 | 164.093 | 10, 8442 | 1.015.957 | 67,1407 |
|  | 24 | 1.500.663 | 79 | 0,0052 | 50.556 | 3,3689 | 146.869 | 9,7869 | 941.656 | 62,7493 |
|  | 27 | 1.492 .550 | 182 | 0,0121 | 60.052 | 4,0234 | 240.903 | 16,1403 | 877.990 | 58,8248 |
|  | 30 | 1.485.934 | 171 | 0,0115 | 54.535 | 3,67 | 219.899 | 14,7987 | 821.506 | 55,2854 |
| 范 | 1 | 1 | 1 | 100.00 | 1 | 100.00 | 1 | 100.00 | 1 | 100.00 |
|  | 2 | 6 | 6 | 100.00 | 6 | 100.00 | 6 | 100.00 | 6 | 100.00 |
|  | 3 | 26 | 26 | 100.00 | 26 | 100.00 | 26 | 100.00 | 26 | 100.00 |
|  | 7 | 8.597 | 8.351 | 97,1385 | 8.351 | 97,1385 | 8.352 | 97,1501 | 8.420 | 97,9411 |
|  | 8 | 35.159 | 31.692 | 90,139 | 34.728 | 98,7741 | 34.730 | 98,7798 | 34.771 | 98, 8964 |
|  | 9 | 128.503 | 44.411 | 34,5602 | 120.629 | 93,8725 | 126.937 | 98,7813 | 127.696 | 99,3719 |
|  | 13 | 10.246.752 | 4 | 0 | 4.171 | 0,0407 | 60.288 | 0,5883 | 1.880.747 | 18,3545 |
|  | 14 | 8.724.060 | 4 | 0 | 1.097 | 0,0125 | 28.198 | 0,3232 | 1.330 .687 | 15,253 |
|  | 15 | 6.289.576 | 1 | 0 | 539 | 0,0085 | 18.155 | 0,2886 | 1.020.736 | 16,229 |
|  | 16 | 4.930.514 | 1 | 0 | 401 | 0,0081 | 14.973 | 0,3036 | 900.736 | 18,2686 |
|  | 17 | 4.444.240 | 1 | 0 | 362 | 0,0081 | 14.253 | 0,3207 | 868.765 | 19,5481 |
|  | 18 | 4.169.242 | -- | -- | 238 | 0,0057 | 12.408 | 0,2976 | 808.236 | 19,3856 |
|  | 19 | 4.042.036 | -- | -- | 200 | 0,0049 | 11.778 | 0,2913 | 785.001 | 19,4209 |
|  | 20 | 4.012 .001 | -- | -- | 182 | 0,0045 | 11.640 | 0,2901 | 779.692 | 19,4339 |
|  | 21 | 3.906 .581 | -- | -- | 147 | 0,0037 | 18.179 | 0,4653 | 739.870 | 18,939 |
|  | 24 | 3.742.557 | -- | -- | 87 | 0,0023 | 16.069 | 0,4293 | 688.272 | 18,3904 |
|  | 27 | 3.600.261 | -- | -- | 49 | 0,0013 | 14.361 | 0,3988 | 644.961 | 17,9142 |
|  | 30 | 3.472.107 | -- | -- | 39 | 0,0011 | 12.943 | 0,3727 | 607.449 | 17,4951 |

### 4.5 Data mining

Previously described sets of n-grams and repeats were used as the input to Data Mining process. Two different data mining techniques were applied: association rules and classification. In process of determining association rules, the complete set of ngrams (repeats) is used as input. In classification process, data were divided into two subsets: model and test (see section 3.2). Classification models were built using model subset as input and verified on test subset. For both techniques results were obtained using IBM Intelligent miner [31].

### 4.5.1 Association rules

Association rules were obtained using SIDE (Simultaneous Depth-first Expansion) algorithm [32] with the following parameters: confidence>=51\%, support $>=0.0001$ and lift $>=1.05$ or lift $<=0.95$. Association rules were obtained for each n-gram or repeat length from 2 to 10 . Typical result produced by Intelligent miner is shown on Figure 17 and includes association rules, rule support, confidence, lift, absolute support (number of n-grams that satisfy rule), rule body, rule head, number of items in rule body and rule head, group (rules having head 'ORDER_LEVEL_IU='D' belong to group 2, while rules indicating order level ' O ' belong to group 1), and weight mean (here empty). More information about meaning of each field can be found in [33].


Figure 17. Association rules for n-grams with length=10 produced by IBM Intelligent miner. Information about each rule includes rule and related support, confidence, lift, absolute support, number of items in rule body and rule head, group, rule body, rule head and weight mean.

Association rules can also be represented graphically. If number of rules is large, presenting all rules on a single picture would make the picture cumbersome and ambiguous. For this reason on Figure 18, for example, only the rules related to disordered regions are shown. Rule head is in the middle of the figure while n-grams
that belong to rule bodies are on the circle. Two of three measure parameters (support, confidence, lift) can be (arbitrary) selected for presentation on the figure:

- by line colour; confidence level is presented on the figure by colour spectrum from highest (ocher in tone) to lowest (blue).
- by line width; support is presented on the figure by line width - n-grams with higher support are connected to rule head with wider line.
- by numbers; numerical values of the parameters presented by colour and width are shown on corresponding line.


Figure 18. Graphical presentation of association rules

### 4.5.1.1 Association rules of AA n-grams

Total numbers of discovered rules per n-gram lengths are shown in Table 12. Each rule includes only one n-gram. Although rules for ordered regions are more numerous and have larger support, they have significantly lower average value of lift, and uniform but
small standard deviation of lift. As higher lift means, by default, more interesting rule, the conclusion that can be derived from Table 12 is that rules for disordered regions are, in general, more significant, and that n-grams much better characterize disordered than ordered regions. Appendix tables A14, A15 and A16 contains first 100 rules for each ngram length that characterize all three types of regions.

Parameters used in association rules results in significantly lower number of rules (i.e. n-grams) compared with corresponding number of n-grams for z -score values (Table 9) or fractional difference (Table 7). The reasons are:

- different meaning of rules compared to classification (for example confidence>=51\% implies that majority of n-grams appears in specific region)
- more restrictive support level than in mole fractions or fractional difference method (support>=0.0001 can be considered as mole fractions threshold equal to 1E-6 on global level, not on the level of individual order level as mole fractions are) ${ }^{7}$
- additional filtering with lift interval which discards rules with low level of interestingness (i.e. rule that are expected to occur).

Combining the results obtained from the association rules, mole fractions, fractional difference and z -score methods produce smaller set of $n$-grams that characterize regions from different points of view and very high confidence. Numbers of n-grams in the intersection set are shown in Table 13. Numbers of n-grams in this table are relatively small because of the different characteristics of methods. For example, because of confidence $\geq 51 \%$ for association rules, first condition that some n-gram can be marked as characteristic one, for some region type, is that more than half occurrences of that n gram are found in the regions of such type. On the other side, fractional difference or z score values can be higher for n-gram in such region if majority of occurrences of this n-gram belongs to region with different type. Also, some n-grams have standard deviation equal to zero and hence their z-score can not be calculated (this is especially expressed for n-grams with length 8, 9 and 10). Percentages of order levels agreement

[^4]between methods are shown on Figure 19 （A：for n－grams in disordered；B：for n－grams in ordered regions）．

Table 12．Association rules characteristics for disordered and ordered regions． Parameters used for discovering rules are：confidence＞＝51\％，support＞＝0．0001 and lift＞＝1．05 or lift $<=0.95$

| Rules for disordered regions |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Lift |  |  |  | Support |  | Confidence |  |
|  | 』 Z 0 0 0 む 乙 |  |  | $\bar{\Sigma}$ | ${\underset{\Sigma}{x}}_{\text {肴 }}$ | $\begin{aligned} & \text { ~0 } \\ & \text { ơ } \\ & 0 \\ & \stackrel{2}{4} \end{aligned}$ |  | $\begin{aligned} & \text { 品 } \\ & \text { ©id } \\ & \text { 交 } \end{aligned}$ |  |
| 2 | 1 | 2，88 | 0 | 2，88 | 2，88 | 0，1479133 | 0 | 58，17 | 0 |
| 3 | 150 | 2，95 | 0，33 | 2，63 | 4，45 | 0，0098004 | 0， 0059749 | 56，20 | 6，30 |
| 4 | 5.119 | 3，29 | 0，44 | 2，79 | 5，58 | 0，0005698 | 0， 0005616 | 59，05 | 7，89 |
| 5 | 6.778 | 4，28 | 0，77 | 2，96 | 5，92 | 0，0001690 | 0， 0001894 | 72，30 | 13，09 |
| 6 | 781 | 5，72 | 0，74 | 3，13 | 6，26 | 0，0002048 | 0，0002725 | 91，48 | 11，86 |
| 7 | 339 | 6，23 | 0，72 | 3，33 | 6，61 | 0，0002100 | 0， 0002423 | 94，26 | 11，00 |
| 8 | 187 | 6，51 | 0，83 | 3，51 | 6，96 | 0，0002178 | 0， 0002045 | 93，50 | 12，04 |
| 9 | 135 | 6，81 | 0，90 | 3，70 | 7，32 | 0，0002103 | 0， 0001729 | 93， 03 | 12，36 |
| 10 | 97 | 7，00 | 1，03 | 3，93 | 7，68 | 0，0002160 | 0，0001553 | 91，13 | 13，43 |


| Rules for ordered regions |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Lift |  |  |  | Support |  | Confidence |  |
|  |  | $\begin{aligned} & \text { ~ } \\ & 0 \\ & 0 \\ & 0 \\ & 0 \\ & <~ \end{aligned}$ |  | $\stackrel{\Xi}{\Sigma}$ | $\underset{\Sigma}{\text { x }}$ | $$ |  | $\begin{aligned} & \text { ~0 } \\ & 00 \\ & 000 \\ & \gtrless \\ & \hline \end{aligned}$ | иоџ̣џ!̣иәр рхериеıS |
| 2 | 329 | 1，03 | 0，16 | 0，64 | 1，26 | 0，1874637 | 0，1163087 | 80，87 | 12，96 |
| 3 | 6.533 | 1，07 | 0，17 | 0，64 | 1，29 | 0，0093590 | 0， 0076481 | 82，93 | 13，25 |
| 4 | 109.586 | 1，08 | 0，17 | 0，64 | 1，29 | 0，0005483 | 0， 0004824 | 83， 81 | 13，70 |
| 5 | 99.911 | 1，14 | 0， 14 | 0，65 | 1，30 | 0，0001429 | 0， 0000515 | 87，60 | 11，33 |
| 6 | 3.167 | 1，27 | 0， 09 | 0，65 | 1，31 | 0，0001556 | 0， 0000874 | 97，13 | 6，93 |
| 7 | 1.811 | 1，30 | 0， 07 | 0，66 | 1，32 | 0， 0001674 | 0， 0000963 | 98，41 | 5，42 |
| 8 | 1.363 | 1，31 | 0， 07 | 0，66 | 1，33 | 0，0001718 | 0， 0000977 | 98，38 | 5，56 |
| 9 | 1.146 | 1，32 | 0， 07 | 0，69 | 1，34 | 0，0001712 | 0， 0000977 | 98， 36 | 5，76 |
| 10 | 948 | 1，33 | 0，07 | 0， 70 | 1，35 | 0，0001734 | 0，0000980 | 98，44 | 5，46 |


| Rules for border regions |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Lift |  |  |  | Support |  | Confidence |  |
|  | $\begin{aligned} & \frac{y}{0} \\ & \text { Z } \\ & 0 \\ & 0 \\ & 0 \\ & 0 \\ & Z \end{aligned}$ |  |  | $\sum$ | $\sum_{\Sigma}^{\text {® }}$ |  |  |  |  |
| 2 3 4 |  |  |  |  |  |  |  |  |  |
| 5 | 55 | 10，13 | 1，58 | 7，64 | 13，89 | 0，0001489 | 0，0000630 | 67，66 | 10，60 |
| 6 | 57 | 9，71 | 2，00 | 6，16 | 12，32 | 0，0001545 | 0，0000557 | 78，82 | 16，23 |
| 7 | 58 | 8，56 | 1，63 | 5，28 | 10，56 | 0， 0001545 | 0， 0000534 | 81， 04 | 15，46 |
| 8 | 57 | 7，53 | 1，45 | 4，65 | 9，31 | 0，0001553 | 0，0000512 | 80，86 | 15，63 |
| 9 | 56 | 6，83 | 1，19 | 4，30 | 8，37 | 0，0001539 | 0，0000503 | 81，58 | 14，23 |
| 10 | 53 | 6，23 | 1，14 | 3，90 | 7，64 | 0，0001521 | 0，0000521 | 81，55 | 14，95 |

Table 13. Numbers of n-grams in intersection set of fractional difference, z -score and association rules methods depending on fractional difference.

Order level: FD/z-score - order level of n-gram in combination of fractional difference and z-score methods; association rules - order level of n-gram according found association rule. Blue cells: numbers of n-grams with identical order level in all methods; yellow cells: numbers of n-grams with different order level in FD/z-score and association rules methods.

| N -gram length | Order level |  | Minimal value of n-gram mole fraction |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | FD/z-score | association rules | 5.0E-6 | 1.0E-6 | 5.0E-7 | 1.0E-7 |
| 3 | D | D | 15 | 15 | 15 | 15 |
|  |  | O | 166 | 166 | 166 | 166 |
|  | O | O | 1.087 | 1.089 | 1.089 | 1.089 |
| 4 | D | D | 1.046 | 1.046 | 1.046 | 1.046 |
|  |  | O | 3.446 | 4.180 | 4.180 | 4.180 |
|  | O | O | 9.143 | 18.881 | 18.881 | 18.881 |
| 5 | D | D | 2.458 | 2.458 | 2.458 | 2.458 |
|  |  | O | 375 | 2.062 | 2.063 | 2.063 |
|  | O | O | 322 | 33.231 | 33.231 | 33.231 |
|  |  | D | 0 | 2 | 5 | 8 |
| 6 | D | D | 436 | 436 | 436 | 436 |
|  |  | O | 0 | 6 | 7 | 8 |
|  | O | O | 126 | 2.746 | 2.746 | 2.746 |
|  |  | D | 0 | 5 | 5 | 6 |
| 7 | D | D | 190 | 190 | 190 | 190 |
|  |  | O | 0 | 4 | 4 | 4 |
|  | O | O | 104 | 1.553 | 1.553 | 1.553 |
|  |  | D | 0 | 5 | 6 | 6 |
| 8 | D | D | 60 | 60 | 60 | 60 |
|  | O | O | 33 | 493 | 493 | 493 |
|  |  | D | 0 | 2 | 3 | 5 |
| 9 | D | D | 30 | 30 | 30 | 30 |
|  | O | O | 12 | 125 | 125 | 125 |
| 10 | D | D | 10 | 10 | 10 | 10 |
|  | O | O | 2 | 41 | 41 | 41 |

It is interesting that number of n-grams with identical order levels in all methods does not dramatically change for various mole fractions smaller than 5E-6. Also, as length of n-grams increase, numbers of order levels differences decrease, and for lengths 9 and 10 there are no differences in order levels for the identical n-grams. These trends remain the same if percentages are considered instead of n-grams numbers.

N -grams that belong to resulting set, without restriction related to mole fractions, are listed in Appendix tables A17 (disordered regions) and A18 (ordered regions). The minimal n-gram length is 3 because no $z$-score exists for shorter n-grams. Tables includes up to 100 n -grams (if there are so many characteristic n-grams for appropriate length) ordered according lift, confidence, and support, all in descending order.


Figure 19. Percentage of order levels agreement between FD/z-score and association rules methods. A: n-grams in disordered regions; B: n-grams in ordered regions

Final set of n-grams that characterize disordered regions includes:

1) homorepeat n-grams of various lengths (like HHHH, KKKKK, GGGGGG, NNNNNN, PPPPPPP, TTTTTTT, EEEEEEEE, DDDDDDDDD, QQQQQQQQQ, SSSSSSSSSSS, etc) of AAs that are disorder promoting (G, K, E, D, Q, S, P, E) or border promoting (N, T, H). Homorepeats of A amino acids are found in association rules but are not member of final set because they do not satisfy z-score condition - either have large z-score in both ordered and disordered regions or have smaller absolute value of $z$-score than necessary for confidence level of 99\% ( $\pm 2.58$ ).
2) their combinations with some AA (like PPPA, REEEE, TGGGGG, GAGGGGGS, RYGGGGGGG, etc)
3) tandem repeats like n-grams of disorder promoting AAs (for example, KPAPKPAP, PSPPPPPSPPP, PEPEPEPE, GGEGGEGG, etc)
4) palindromes of disorder promoting AAs (for example, QPQPQ, DEEEED, PAPAPAPAP, etc.)

Final set of n-grams that characterize ordered regions includes:

1) n-grams that include bigrams or trigrams of order promoting AAs (bigrams: VV, FF, WW, YY; trigram LLL)
2) almost all n-grams that include bigram CC or II. More than $99.5 \%$ of n-grams that include bigram II are classified as order or border characteristics, with exception n-grams where II is surrounded by disorder promoting AAs. N-grams RPADIII, IISTPA, ADIIIST, PADIII, ADIIIS, IIISTPAS, PADII are marked as disorder promoting while n-gram MKKII is marked as border promoting. Also, about $90.5 \%$ n-grams that include bigram CC characterize order region, while others characterize border regions.

Only a few of the patterns can be observed in the final set of n-grams that characterize border regions:

1) n-grams that contain HCP, PLLN, YFYDS characterize border regions only
2) n-grams that contain QID, TRS, FQI, TEG, and YFY prefer border regions but also characterize order regions. Some of them (like PLL or YFY) are sub-ngrams of n-grams that characterize border regions only

### 4.5.1.2 Comparison with data from DisProt database

Previous results can be compared with corresponding data from DisProt database. The same methods (mole fractions, fractional difference, z-score and association rules) were applied on data available from DisProt database. Due to the initial smaller number of n-grams, the final set of DisProt n-grams also have low cardinality and the intersection between this set and set of obtained results is too small. That's why results of comparison will be shown in three figures: Figure 20 includes results of comparison order levels of identical n-grams from final (intersected) sets; Figure 21 includes results of comparison of order levels generated by association rules, and Figure 22 includes results of comparison of order levels generated by fractional difference and z -score. In all three figures numbers and percentages of identical n-grams with equal/not equal order levels in DisProt and used material are presented.

Also, as explained in the beginning of the Chapter 3, results of comparison where n-gram is predicted to be disorder related, but in DisProt it is order related, should be taken with reserve. These numbers in the corresponding tables on figures 2022 are marked yellow, while results of comparison where n-gram is in disordered region in DisProt and ordered region in material are marked red.

|  |  |  |  |
| :---: | :---: | :---: | :---: |
|  |  | D | O |
| $\begin{aligned} & \text { 플 } \\ & \text { 促 } \end{aligned}$ | 3 |  | 45 |
|  | 4 | 34 | 398 |
|  | 5 | 20 | 148 |
|  | 6 | 1 | 2 |
|  | 8 | 1 |  |
|  | 3 | 3 |  |
|  | 4 |  | 27 |
|  | 5 |  | 44 |
|  | 6 |  | 1 |
|  | 7 |  | 3 |



Figure 20. Number and percentage of equal/not equal order levels related to identical ngrams obtained from intersection of result sets of n-grams from DisProt and used material. There are no identical n-grams with length 7 in DisProt and used material in intersection of sets.

| $\begin{aligned} & \frac{\pi}{0} \\ & \frac{0}{4} \\ & 000 \end{aligned}$ |  |  |  |
| :---: | :---: | :---: | :---: |
|  |  | D | O |
|  | 2 |  | 235 |
|  | 3 | 7 | 4.782 |
|  | 4 | 595 | 23.944 |
|  | 5 | 282 | 1.567 |
|  | 6 | 40 | 8 |
|  | 7 | 15 | 1 |
|  | 8 | 5 |  |
|  | 9 | 4 |  |
|  | 10 | 4 |  |
|  | 2 |  | 1 |
|  | 3 |  | 122 |
|  | 4 | 13 | 1.910 |
|  | 5 |  | 433 |
|  | 6 |  | 35 |
|  | 7 |  | 9 |
|  | 8 |  | 6 |
|  | 9 |  | 6 |
|  | 10 |  | 4 |



Figure 21. Number and percentage of equal/not equal order levels related to identical ngrams from association rules generated on n-grams from DisProt and used material


Figure 22. Number and percentage of equal/not equal order levels related to identical ngrams from fractional difference/z-score results based on n-grams from DisProt and used material

Although the numbers of n-grams with equal/not order level are not too high (which in some cases seems that they are not enough representative) the common trend can be observed in all three figures:

1) For the n-grams length 3 and 4 accuracy of characterization is significantly lower than $50 \%$ for disordered regions and significantly higher for ordered regions
2) N-gram length 5 is a crossing point - accuracy moved to disorder side. Further increasing n-grams length increase the accuracy of characterization for disordered regions and decrease accuracy of characterization for ordered regions.
3) It is possible that accuracy of prediction for longer n-grams is higher than presented. As previously noticed current version of DisProt database include precise information only about protein regions that are experimentally proved as disorder. Consequence is that set of disorder related n-grams selected from DisProt is not complete. It is possible that longer n-grams currently not recognized as disorder related in DisProt are actually disorder-characterize ones.

For example, differences between order levels in association rules based on DisProt and used material are related to the following n-grams which consist of disorder promoting AAs only.

```
Length N-grams
DSDSDSD, GGGGSGG, GGGSGGG, GPPGPPG,PEPEPEP,QQQQQQQ, SDSDSDS, SGGGGGG, SSSSSSS
    DSDSDSDS, EEEEEEEE,QQQQQQQQ,SDSDSDSD,SGGGGGGG,SSSSSSSS
    DSDSDSDSD,EEEEEEEEE,QQQQQQQQQ,SDSDSDSDS,SGGGGGGGG,SSSSSSSSS
    DSDSDSDSDS,QQQQQQQQQQ,SDSDSDSDSD,SSSSSSSSSS
```


### 4.5.1.2.1 Finding patterns in characteristic n-grams

Additional research was done to discover patterns related to characteristic ngrams. All substrings of n-grams with lengths great than or equal 3 was considered as potential pattern. Such substring is marked as characteristics for order (disorder) region if it is not part of any n-gram that characterize disorder (order) region. Surprisingly, number of such patterns is not too low; number of patterns for material used in this research (download from NCBI), material from DisProt database and intersection of these two sets are shown in of Table 14.

Table 14. Number of sequences (sub-n-grams) that belong to n-grams and characterize some region type.

| Order level | Pattern length | NCBI material | DisProt material | Intersection |
| :---: | :---: | :---: | :---: | :---: |
| D | 3 | 100 | 184 | 4 |
|  | 4 | 2463 | 1268 | 131 |
|  | 5 | 2641 | 811 | 27 |
|  | 6 | 470 | 188 | 3 |
|  | 7 | 206 | 34 | 2 |
|  | 8 | 77 | 17 | 1 |
|  | 9 | 36 | 7 | -- |
|  | 10 | 10 | 2 | -- |
| N | 3 | 1 | -- | -- |
|  | 4 | 37 | 9 | -- |
|  | 5 | 83 | 9 | -- |
|  | 6 | 70 | 9 | -- |
|  | 7 | 73 | 9 | -- |
|  | 8 | 70 | 9 | -- |
|  | 9 | 63 | 10 | -- |
|  | 10 | 53 | 10 | -- |
| O | 3 | 5559 | 2819 | 2060 |
|  | 4 | 38010 | 6038 | 2201 |
|  | 5 | 33946 | 4201 | 171 |
|  | 6 | 2873 | 616 | 3 |
|  | 7 | 1610 | 42 | -- |
|  | 8 | 581 | 6 | -- |
|  | 9 | 157 | 2 | -- |
|  | 10 | 41 | 1 | -- |

For example, patterns of length 6 that belong to both sets and characterize ordered regions are GGLEGL, GSGKST, TGSGKS, while patterns of the same length that characterize disordered regions are APAPAP, GGGGGG, SGSSSS. It is interesting that no intersection between sets exists for sequences that characterize borderline region. Also, it is interesting that, if hydrophobicity (according Kyte-Doolittle scale, further KD scale) of amino acids in patterns are considered then patterns that characterize disordered regions are much hydrophilic than patterns related to ordered regions. Hydrophobicity of patterns is calculated on two ways: as majority of hydrophobic/hydrophilic AA (in this case 'neutral' means that numbers of hydrophilic and hydrophobic AAs are equal), and as a sum of hydrophobic/hydrophilic values according to KD scale (see Table 15). If sum is negative than the pattern is marked as hydrophilic; if sum is positive than the pattern is marked as hydrophobic, and otherwise it is marked as neutral. It can be concluded that pattern in intersection set that characterize disordered regions and can be considered as 'proved disordered' are almost completely hydrophilic. Due to the previously mentioned reasons patterns in DisProt material (and consequently in the intersection) can not be considered as 'proved order' and not commented here.

Table 15. Hydrophobicity of n-gram patterns that characterize regions.
Majority of hydrophobic/hydrophilic AA - majority of AAs in pattern are hydrophobic or hydrophilic
Neutral - pattern consists of equal number of hydrophilic and hydrophobic AAs
Hydrophobic/hydrophilic value - sum of hydro-values of AAs from pattern denotes hydrophilic/hydrophobic object
Neutral value - sum of hydro-values of AAs from pattern is equal to 0
All values are according Kyte-Doolittle scale of AAs hydrophobicity

| $\begin{aligned} & \ddot{U} \\ & \text { O} \\ & \text { in } \end{aligned}$ |  | Disordered regions |  |  |  |  |  | Borderline regions |  |  |  |  |  | Ordered regions |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Percentage of pattern with |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
|  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
|  | 3 | 86,41 | 77,71 | 13,58 | 22,28 | 0,00 | 0,00 |  |  |  |  |  |  | 68,96 | 57,21 | 31,03 | 42,42 | 0,00 | 0,35 |
|  | 4 | 74,05 | 79,33 | 4,25 | 20,34 | 21,68 | 0,31 | 22,22 | 33,33 | 33,33 | 66,66 | 44,44 | 0,00 | 54,38 | 60,69 | 13,36 | 38,65 | 32, 24 | 0,64 |
|  | 5 | 88,03 | 77,80 | 11,96 | 21,82 | 0, 00 | 0,36 | 33,33 | 33,33 | 66,66 | 66,66 | 0, 00 | 0,00 | 74,45 | 58,98 | 25,54 | 40, 89 | 0, 00 | 0,11 |
|  | 6 | 88, 82 | 85,10 | 2,65 | 13,82 | 8,51 | 1,06 | 33,33 | 33,33 | 22,22 | 66,66 | 44,44 | 0,00 | 67,85 | 64,12 | 10,55 | 35,55 | 21,59 | 0,32 |
|  | 7 | 97,05 | 88,23 | 2,94 | 11,76 | 0, 00 | 0,00 | 66,66 | 55,55 | 33,33 | 44,44 | 0, 00 | 0,00 | 92,85 | 90,47 | 7,14 | 9,52 | 0,00 | 0,00 |
|  | 8 | 94,11 | 94,11 | 0,00 | 5,88 | 5,88 | 0,00 | 66,66 | 55,55 | 11,11 | 44,44 | 22, 22 | 0,00 | 100,00 | 100,00 | 0,00 | 0, 00 | 0, 00 | 0,00 |
|  | 9 | 100,00 | 85,71 | 0, 00 | 14,28 | 0,00 | 0,00 | 90,00 | 50,00 | 10,00 | 50,00 | 0, 00 | 0,00 | 100,00 | 100,00 | 0,00 | 0, 00 | 0,00 | 0,00 |
|  | 10 | 100,00 | 100,00 | 0,00 | 0,00 | 0,00 | 0,00 | 60,00 | 50,00 | 0,00 | 50,00 | 40,00 | 0,00 | 100,00 | 100,00 | 0,00 | 0, 00 | 0,00 | 0,00 |
| $\begin{aligned} & \text { E. } \\ & \text { UU } \\ & \text { W. } \\ & \text { H. } \end{aligned}$ | 3 | 100,00 | 75,00 | 0,00 | 25,00 | 0,00 | 0,00 |  |  |  |  |  |  | 63,39 | 51,01 | 36,60 | 48,68 | 0, 00 | 0,29 |
|  | 4 | 81,67 | 92,36 | 0,76 | 7,63 | 17,55 | 0,00 |  |  |  |  |  |  | 36,48 | 41,52 | 21,26 | 57, 74 | 42, 25 | 0,72 |
|  | 5 | 92,59 | 85,18 | 7,40 | 14,81 | 0,00 | 0,00 |  |  |  |  |  |  | 50,87 | 27,48 | 49,12 | 72,51 | 0,00 | 0,00 |
|  | 6 | 66,66 | 66,66 | 0,00 | 33,33 | 33,33 | 0,00 |  |  |  |  |  |  | 100,00 | 66,66 | 0, 00 | 33,33 | 0,00 | 0,00 |
|  | 7 | 50, 00 | 50,00 | 50,00 | 50,00 | 0,00 | 0,00 |  |  |  |  |  |  |  |  |  |  |  |  |
|  | 8 | 0,00 | 0,00 | 0,00 | 100,00 | 100,00 | 0,00 |  |  |  |  |  |  |  |  |  |  |  |  |
|  | 3 | 97,00 | 95,00 | 3,00 | 5,00 | 0,00 | 0,00 | 100,00 | 100,00 | 0,00 | 0,00 | 0,00 | 0,00 | 65,49 | 54,65 | 34,50 | 45,04 | 0,00 | 0,30 |
|  | 4 | 88,63 | 93,78 | 0,85 | 6,21 | 10, 51 | 0,00 | 75,67 | 83,78 | 0,00 | 16,21 | 24,32 | 0,00 | 41,88 | 46,50 | 18,74 | 52,85 | 39,36 | 0,64 |
|  | 5 | 95,11 | 89,85 | 4,88 | 10,03 | 0,00 | 0,11 | 86,74 | 73,49 | 13,25 | 26,50 | 0,00 | 0,00 | 61,68 | 38,89 | 38,31 | 60,69 | 0, 00 | 0,41 |
|  | 6 | 87,65 | 84,04 | 4,68 | 15,95 | 7,65 | 0,00 | 78,57 | 71,42 | 2,85 | 28,57 | 18,57 | 0,00 | 55,72 | 54,12 | 15,94 | 45,59 | 28,33 | 0,27 |
|  | 7 | 91, 26 | 86,40 | 8,73 | 13,59 | 0,00 | 0,00 | 91,78 | 83,56 | 8,21 | 15,06 | 0, 00 | 1,36 | 76,83 | 58,38 | 23,16 | 41,55 | 0, 00 | 0,06 |
|  | 8 | 87,01 | 84,41 | 6,49 | 15,58 | 6,49 | 0,00 | 88,57 | 84,28 | 0,00 | 15,71 | 11,42 | 0,00 | 65,92 | 58,86 | 12,04 | 40,96 | 22,03 | 0,17 |
|  | 9 | 91, 66 | 86,11 | 8,33 | 13,88 | 0,00 | 0,00 | 98,41 | 90,47 | 1,58 | 9,52 | 0,00 | 0,00 | 77,07 | 56,68 | 22,92 | 42,67 | 0, 00 | 0,63 |
|  | 10 | 100,00 | 90, 00 | 0,00 | 10,00 | 0,00 | 0,00 | 98,11 | 84,90 | 0,00 | 15,09 | 1,88 | 0,00 | 73,17 | 63,41 | 14,63 | 36,58 | 12,19 | 0,00 |

### 4.5.1.3 Association rules of nucleotide n-grams

Discovering association rules for complete set of n-grams exceeds computational capability of computer system used for this research. Due to a huge number of n-grams (ranging from 42M to 140M) association rules were discovered on smaller subsets of direct non-complementary nucleotide repeats (n-grams) only, but not on other sort of nucleotide repeats (direct complementary, inverse complementary and inverse non-complementary).

For each n-gram length, set of n-grams is divided in three parts, according to their corresponding ORF-s. Number of discovered rules rapidly decrease as n-gram length increase, and is smaller than number of rules of AAs of corresponding length because of different codon usage tables used for translating AAs. Number of association rules for nucleotide n-gram lengths $15,18,21,24,27$ and 30 is shown in Table 16, and results of the comparison of their translation (using translation table 11) to corresponding AA ngrams is shown in Table 17. It can be seen that longer n-grams are mostly related to disordered regions regardless of ORF.

Table 16. Number of discovered association rules for nucleotide n-grams

| N -gram length | ORF |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 1 |  | 2 |  | 3 |  |
|  | D | O | D | O | D | O |
| 15 | 16 | 15 | 13 | 15 | 14 | 14 |
| 18 | 11 | 12 | 11 | 9 | 9 | 8 |
| 21 | 6 | 7 | 6 | 3 | 4 | 3 |
| 24 | 4 | 2 | 4 | 1 | 4 | 1 |
| 27 | 2 | -- | 3 | -- | 2 | -- |
| 30 | 1 | -- | 2 | -- | 1 | -- |

In all three ORFs nucleotide n-grams behave regularly as well as the corresponding AA n-grams. Longer nucleotide n-grams more precisely characterize both types of regions. Additionally, n-grams with length 27 and 30 characterize only disordered regions, as in the case of similar (with lengths 9 and 10) AA n-grams. Also, some of the n-grams that have different order level than corresponding AA n-grams (equivalent to their translation), are homorepeats of disorder promoting AAs (as S or Q) which are possible disorder related, as previously mentioned.

Table 17. Number and percentage of equal/not equal order levels related to translations of nucleotide n-grams and identical AA n-grams

|  |  | ORF1 |  |  |  | ORF2 |  |  |  | ORF3 |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | D |  | O |  | D |  | O |  | D |  | O |  |
|  |  | num | perc | num | perc | num | perc | num | perc | num | perc | num | perc |
| $\stackrel{\text { 즐 }}{\underset{y}{2}}$ | 15 | 23 | 92.0 | 15 | 100.0 | 12 | 80.00 |  |  | 10 | 71.42 |  |  |
|  | 18 | 14 | 93.33 | 12 | 100.0 | 8 | 100.0 |  |  | 5 | 83.33 |  |  |
|  | 21 | 8 | 100.0 | 7 | 100.0 | 4 | 100.0 |  |  |  |  |  |  |
|  | 24 | 5 | 100.0 | 2 | 100.0 | 3 | 100.0 |  |  |  |  |  |  |
|  | 27 | 2 | 100.0 |  |  | 2 | 100.0 |  |  |  |  |  |  |
|  | 30 | 2 | 100.0 |  |  | 1 | 100.0 |  |  |  |  |  |  |
| $\begin{aligned} & \text { 즘 } \\ & \text { च } \\ & \stackrel{0}{Z} \end{aligned}$ | 15 | 2 | 8.00 |  |  | 3 | 20.00 | 3 | 100.0 | 4 | 28.58 |  |  |
|  | 18 | 1 | 6.67 |  |  |  |  | 2 | 100.0 | 1 | 16.67 |  |  |
|  | 21 |  |  |  |  |  |  |  |  |  |  |  |  |
|  | 24 |  |  |  |  |  |  |  |  |  |  |  |  |
|  | 27 |  |  |  |  |  |  |  |  |  |  |  |  |
|  | 30 |  |  |  |  |  |  |  |  |  |  |  |  |

There are n-grams that occur in association rules related to all three ORF-s. These ngrams have maximal length 9, and as this correspond to AA n-grams of length 3 or shorter which, as shown earlier, do not have high precision in regions characterization (especially not satisfactory level of characterization for disordered regions) so no such n -grams are considered.

### 4.5.1.4 Association rules of inverse non complementary AA repeats

Inverse non-complementary repeat (in further text IN repeats) represents palindrome with a gap of arbitrary ( $\geq 1$ ) length between left and right components of the repeat. Left and right component can belong to different types of regions, so 9 different "double order levels" exist: DD, DO, DN, OD, OO, ON, ND, NO, NN ${ }^{8}$. Repeats characterize region type ' X ' if both components (left and right) fall into regions of such types, so high accuracy is reached in the research only for DD, OO and NN combinations. Also, in process of determination of association rules only left component of repeat (on Figure 23 "REPEAT_LEFT") and double order level combinations are considered because right component is unambiguously determined by the left one.

Association rules are determined for both sets of all IN repeats and statistically significant IN repeats. The similar parameters were used as in determining association

[^5]rules for n-grams: confidence>=51\%, support>=0.0005 and lift $>=1.05$ or lift $<=0.95$. Support threshold for association rules is increased to 0.0005 because, as the number of repeats is significantly lower than number of n-grams with the same length, using support equal to 0.0001 as in association rules for n-grams lead to plenty of association rules for small repeat lengths. The results obtained have similar form as in the case of ordinary n-grams, as illustrated on Figure $23^{9}$.


Figure 23. Association rules for IN repeats with length=10 produced by IBM Intelligent miner. Information about each rule includes rule and related support, confidence, lift, absolute support, number of items in rule body and rule head, group, rule body, rule head and weight mean.

Association rules are determined for all repeat lengths from 3 to 10 where term "repeat length" is related to length of either left or right component of repeat. Association rules are also determined for DisProt repeats. Number of rules found for different repeat lengths are shown in Table 18. There is one anomaly in the table: regardless the significantly higher number (about 20 times) of all repeats with length 3 in the used material (from NCBI) than in DisProt material, the number of association rules for this category of repeats is higher for DisProt material. The reason is very small absolute support (two) for DisProt repeats which corresponds to support 0.0005. For the same reason sets with smaller number of repeats (repeat length $>5$ in NCBI material and

[^6]repeat length $>3$ for DisProt material) produce majority of rules that occurs only once in the complete results (see Table 18).

Table 18. Number of discovered association rules for inverse non-complementary repeats where source material is originated from NCBI and DisProt. Results are shown for sets of all and statistically significant repeats. Also, for each set, number of association rules where repeat included in the body of the rule occurs more than 1 in complete material (column "absolute support>1") is shown. Repeat length is related to length of either left or right component of repeat.

|  | Repeats from NCBI material |  |  |  | Repeats from DisProt |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | All repeats |  | Statistically significant |  | All repeats |  | Statistically significant |  |
|  | All rules | Abs. support>1 | All rules | Abs. support>1 | All rules | Abs. support>1 | All rules | Abs. support>1 |
| 3 | 4186 | 4186 | 3566 | 3566 | 4645 | 4645 | 2925 | 2925 |
| 4 | 17175 | 17175 | 15130 | 15130 | 7342 | 2738 | 5360 | 2616 |
| 5 | 8691 | 8691 | 8021 | 8021 | 2421 | 582 | 2234 | 538 |
| 6 | 5057 | 2539 | 5011 | 2520 | 363 | 73 | 361 | 73 |
| 7 | 9369 | 3091 | 9328 | 3076 | 353 | 49 | 353 | 49 |
| 8 | 1471 | 521 | 1471 | 521 | 70 | 16 | 70 | 16 |
| 9 | 1902 | 584 | 1902 | 584 | 82 | 16 | 82 | 16 |
| 10 | 457 | 152 | 457 | 152 | 19 | 9 | 19 | 9 |

Analysis of rules obtained for all and statistically significant repeats produce the following results:

1. For smaller repeat length all rules have absolute support>1
2. If number of rules is equal for all repeats and statistically significant repeats than rules are identical
3. For larger repeat length the set of repeats have smaller cardinality and predefined support 0.0005 is equivalent to absolute support 1 with consequence that all repeats are taken into consideration and produce some rules. There are no guarantees that such rules with minimal possible absolute support are valid in general. Because those rules can not produce highly accurate results, they will not be taken into consideration.
4. If it is assumed that the probability of appearance each individual AAs is equal, the following filter can be applied on rules based on smaller repeat length: if support for rule is smaller than probability for repeat occurring (for trigrams 0.0125 , for tetragrams 0.000625 ) than this rule is ignored. Although this presumption does not hold in real life (because frequency of occurring is not the same for different AAs and depends on content of material) proposed filter is useful for decreasing number of rules with low probability. Rule is not
applicable on repeats with length longer than 4 for NCBI based material and longer than 3 for Disprot based material because probability of occurring specific repeats is lower than predefined support for association rules. When this filter is applied on repeats from NCBI based material number of rules for statistically significant repeats becomes larger than number of rules for all repeats ${ }^{10}$ ( 827 for all repeats and 869 for statistically significant repeats of length 3, and 12691 and 15130 for length 4). For DisProt material number of repeats decrease to 1245 (all repeats) and 1100 (statistically significant repeats).
5. An additional reduction in the number of rules that are considered in further analysis is achieved by using only those rules with double order level 'OO', 'DD' or 'NN', which are useful for region characterization. Percentage of these rules in total number of rules before and after applying filters (including probability filter and absolute support>1) are

|  | Material from NCBI |  | Material from DisProt |  |
| :---: | :---: | :---: | :---: | :---: |
|  | Before | After | Before | After |
| All repeats | $95.24 \%$ | $88.29 \%$ | $93.22 \%$ | $71.36 \%$ |
| Stat. signif. rep. | $95.44 \%$ | $89.44 \%$ | $91.80 \%$ | $76.71 \%$ |

Rules with longer repeats do not contain other order levels than 'OO','DD' or 'NN'. Numbers of rules after applying previous filters are shown in Table 19.
6. In general, rules for repeats with length $>3$ that do not belong to the set of statistically significant repeats have the following characteristics:
a. Higher support corresponds to lower confidence. Majority of such rules have double order level 'OO', confidence between 0.51 and 0.65 and lift near 0.95 and 1.05 . As this value of lift indicates that the rule body and the rule head appear almost as often together as expected, means that the occurrence of the rule body has almost no effect on the occurrence of the rule head, these rules will not be taken into account for determining characterization strings.

[^7]b. Majority of rules with high confidence have small support and very low absolute support (2 or 3).
7. Rules based on repeats from NCBI with length=3 that do not belong to the set of statistically significant repeats have the following characteristics:
a. All rules have double order level ' OO '
b. There are no rules with confidence $100 \%$. About $65 \%$ of rules have lift smaller than 0.95 and confidence below 61.5\%.
c. About $30 \%$ rules have confidence $>70 \%$ and lift $>1.08$. Repeats in these rules are potential characteristic sequences (for ordered regions). Majority of these repeats have palindromes as left and right components. Left components of these repeats are: AVI, CDC, CEC, CKC, CRC, CSC, CTC, ELG, FCF, FHF, FMF, FWF, HAH, HDH, HEH, HFH, HIH, HKH, HNH, HVH, HYH, IDA, IED, IWI, KVI, MFM, MIM, MVM, NWN, PCP, PWP, RTL, WLW, YCY, YHY, YMY
8. Rules based on repeats from DisProt with length=3 and support>0.0125 that do not belong to the set of statistically significant repeats, have the following characteristics:
a. All rules have double order level 'OO'
b. About $30 \%$ of rules include repeats that have palindromes as left and right components.
c. There are $\approx 10 \%$ rules with confidence $100 \%, \approx 17 \%$ of rules have lift smaller than 0.95 and confidence below $61.5 \%$.
d. About $78 \%$ rules have confidence $>70 \%$ and lift $>1.08$. As previously mentioned, there is no guarantee that protein regions in DisProt database that are not marked as disordered are ordered. Based on this premise, there is no guarantee that repeats in such rules can be used as strings that characterize ordered regions, and hence such repeats were not listed.

Based on the results of this analysis, the set of rules based on statistically significant repeats from NCBI material with previously described filters applied was used as a base for determining repeats that characterize protein regions. For verification of obtained results set of rules based on statistically significant repeats from DisProt material with applied same filters was used. Numbers of rules obtained after applying filers are shown in Table 19.

Table 19. Number of association rules based on repeats after applying filters

| Repeat <br> length | Repeat from NCBI material |  |  |  | Repeats from DisProt material Order level |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Order level |  |  |  |  |  |  |  |
|  | All | DD | OO | NN | All | DD | OO | NN |
| 3 | 869 | 78 | 791 | -- | 1032 | 31 | 1001 | -- |
| 4 | 13872 | 1898 | 11969 | 5 | 2375 | 218 | 2157 | -- |
| 5 | 7589 | 2230 | 5311 | 48 | 507 | 147 | 360 | -- |
| 6 | 2463 | 991 | 1427 | 45 | 68 | 28 | 40 | -- |
| 7 | 3066 | 1475 | 1454 | 137 | 49 | 28 | 21 | -- |
| 8 | 517 | 371 | 124 | 22 | 16 | 8 | 8 | -- |
| 9 | 584 | 442 | 124 | 18 | 16 | 10 | 6 | -- |
| 10 | 152 | 134 | 15 | 3 | 9 | 5 | 4 | -- |

Appendix Tables A20 -- A22 include left components of repeats that characterize disordered, ordered and borderline regions from NCBI and Tables A23-A24 include left components of repeats that characterize disordered and ordered regions from DisProt material respectively. Tables include first 100 repeats (if exists), ordered by confidence, lift and support, all in descending order. Although it seems that if some n-gram ' X ' characterize some region type ' Y ' that repeat with left or right component equal to ' X ' characterize region type ' Y ' (i.e. 'YY') this is not always true. For example, repeat with left/right components ATTTAA/AATTAA have order level 'OO' while both n-grams ATTTAA and AATTTA have order level 'D' in association rules. Of course, if left and right components of repeat in association rule related to n-grams have confidence $100 \%$ than both rules types characterize the same order level.

Results of comparison of order levels in association rules based on material from NCBI and DisProt are shown in Table 20. As in previous cases, results of comparison where repeats are predicted to be disorder related, but in DisProt they are order related, should be taken with reserve. These numbers in the Table 20 are marked yellow, while results of comparison where repeats are in disordered region in DisProt and ordered region in material from NCBI are marked red. As in previous comparison with DisProt, there are no disagree in order levels for longer repeats when order level in DisProt is equal to 'DD', i.e. method provide high accuracy for repeat length $\geq 7$. Again, as in previous cases, as left components of repeats that are not equal when order level in DisProt is equal to 'OO' for length $\geq 7$ are

[^8]which include only disorder promoting AAs, it can be supposed, with a high probability, that the characterization of the disorder regions is one hundred percent correct for repeats with length $\geq 7$.

Table 20. Numbers of equal/not equal order levels related to identical repeats in association rules. Source: materials from DisProt an NCBI.

| Order levels | Repeat length | Order level in association rules based on DisProt repeats |  |
| :---: | :---: | :---: | :---: |
|  |  | DD | OO |
| Equal | 3 | 7 | 357 |
|  | 4 | 75 | 624 |
|  | 5 | 72 | 109 |
|  | 6 | 14 | 5 |
|  | 7 | 16 | -- |
|  | 8 | 3 | -- |
|  | 9 | 5 | -- |
|  | 10 | 2 | -- |
| Not equal | 3 |  | 41 |
|  | 4 |  | 212 |
|  | 5 |  | 22 |
|  | 6 | - | 17 |
|  | 7 |  | 11 |
|  | 8 |  | 5 |
|  | 10 |  | 6 |
|  |  |  | 3 |

Some general characteristics related to repeats (material from NCBI) that characterize regions are:

1) Homorepeats of all amino acids except $Y$ characterize some type of region. In general, homorepeats of disorder promoting AAs characterize disordered regions and homorepeats of order promoting AAs characterize ordered regions. Exceptions are M , which characterizes ordered regions, and H and N , which characterize disordered regions. There is no overlapping or duplicate characterization - not even one amino acid characterizes different region type for different homorepeat length. Only homorepeats of amino acid A have lift smaller than 1 (more precisely smaller than 0.878 ). Characterizations of region types by homorepeats are very accurate. As illustration, found homorepeats, their lengths, lift and confidence of corresponding association rule are shown in Appendix table A25.
2) All rules with repeats whose length is 10 have confidence $100 \%$ and lift 1.0963, regardless support which varies between 8.895 and 0.110 . The only exception is rule with repeat AAAAAAAAAA with confidence $80 \%$, support 0.884 and lift 0.877 , from which can be concluded that amino acid A (which is small and hydrophobic) behaves little different than other disorder promoting AAs.
3) Majority of left and right components of repeats are palindromes itself (see Figure 24).


Order level/Repeat length

Figure 24. Percentage of palindromic left/right components of repeats that characterize regions

Left and right components that are not palindromes are

- tandem repeats, or
- combinations of smaller homorepeats or palindrome with some AAs

4) Tandem repeats are highly represented in repeats which characterize some region (see Table 21). It is interesting that almost all longer repeats that are not palindromes itself are tandem repeats (see Table 20), while shorter repeats that are neither palindromes nor tandem repeats also includes some sub-palindrome (length $\geq 3$ ) combined with other AAs. ${ }^{11}$
[^9]Table 21. Percentage of tandem repeats in set of all repeats and in nonpalindrome repeats

| Repeat <br> length | Tandem repeat percentage |  | Non-palindrome Tandem repeat percentage |  |  |  |
| :---: | ---: | ---: | ---: | ---: | ---: | ---: |
|  | DD | NN | OO | DD | NN | OO |
| 4 | 7,00 | 0,00 | 0,65 | 4,05 | 0,00 | 0,61 |
| 5 | 22,15 | 8,33 | 7,26 | 24,51 | 0,00 | 4,98 |
| 6 | 57,31 | 31,11 | 19,90 | 75,75 | 100,00 | 27,34 |
| 7 | 76,54 | 36,49 | 42,09 | 95,18 | 0,00 | 37,50 |
| 8 | 93,26 | 50,00 | 50,80 | 96,03 | 0,00 | 100,00 |
| 9 | 95,24 | 72,22 | 70,96 | 100,00 | 0,00 | 100,00 |
| 10 | 99,25 | 100,00 | 100,00 | 100,00 | 0,00 | 100,00 |

5) If repeat includes only order promoting AAs, it does not characterize disordered region, with exception of only 20 repeats:

- 7 homorepeats of AA Asparagine (length from 4 to 10 )
- 7 homorepeats of AA Histidine (length from 4 to 10)
- repeats with very small support/absolute support:
o NNNYNNN (abs. support=4),
o LHHHHL, HHNHH, INNNNN, HHYHH, HHLHH (abs. support=2)


### 4.5.2 Classification

Another method for discover characteristic n-grams can be applying tree classification method on available set of repeats to predict order/disorder class. Although the obtained model has very limited capabilities ${ }^{12}$ for correct prediction on previously unseen material it can be used for discovering $n$-gram sequences that characterize order/disorder regions (class in model). Due to a large number of n-grams/palindromes the model could not be constructed based on complete sets of n-grams/palindromes. Instead of that, the initial sets are divided by the association of the phyla. For each phylum, sets of n-grams/palindromes are divided into two parts, as described in chapter 3.2. Classification models are constructed using tree based algorithms SPRINT (Scalable PaRallelizable INduction of decision Trees) [32] for each phylum and

[^10]checked on corresponding test sets. Quality of each of classification models were between $82 \%$ and $96 \%$, while quality of applying constructed models on test data were between $68 \%$ and $85 \%$. Sets of n-grams and palindromes produced in models as characteristics of regions confirm previously obtained results from association rules mining. An example of characteristic n-grams obtained with classification is shown on figure 23.


Figure 23. N-grams from classification model for Anelloviridae phylum that characterize specific type of regions

## 5 Conclusion

Discovering characteristic sequences for ordered/disordered regions in proteins is very important. Intrinsic disorder of proteins are implicated in most important cellular processes such as: cell signaling, transcription and chromatin remodeling functions. On the other side, they are involved in a number of diseases, such as neurological, cardiovascular and malignant pathological states. Taking this in mind, studying structural and dynamical properties of intrinsically disordered proteins is of great importance for better understanding of their actions and developing new medicaments.

In this thesis a new method for determining sequences that characterize ordered/disordered regions with very high confidence is presented. Proposed method establish correspondence with amino acid n-grams to specific region type using n-gram (repeat) characteristics (mole fraction, fractional difference, z -score) and data mining techniques (association rules and classification) applied on both repeats and palindromes. Each of these characteristics/techniques produces n-grams sequences that characterize regions with very high percent of confidence. Sets of sequences produced with various techniques intersect in a very large degree and can be used as characterization sequences for specific region types. General principles that can be observed from the results are:

- type of characterized region depends on sequence (either repeat or palindrome) length
o shorter n-grams (length up to 6) more precisely characterize ordered regions
o longer n-grams (length 6 or longer) more precisely characterize disordered regions
- sequences that appear in intersection of results obtained by different methods (fractional characteristics, $z$-score, association rules) have almost 95\% confidence for characterization
- ordered regions are characterized with
o AAs patterns (VV, FF, WW, YY, LLL)
o almost all n-grams with patterns CC and II
o homorepeats of order/border promoting AAs with exception H and N
o tandem repeats of order promoting AAs
- disordered regions are characterized with
o homorepeats of various lengths of disorder/border promoting AAs with exception M , and their combination with some AA
o tandem repeats of disorder promoting AAs
o palindromes of disorder promoting AAs
o combinations of homorepeats of disorder/border promoting AAs and some (disorder/border promoting) AA (like PPPA, REEEE, TGGGGG, GAGGGGGS, RYGGGGGGG, etc.)
o border regions are characterized with some specific n-grams (HCP, ...) or pattern (PLL or YFY)

The proposed method is verified by compared obtained results with results obtained with applying identical methods on material from DisProt database. Results of this thesis show that exists significant correlation between ordered/disordered regions and specific n-grams which can be used for improvement of disorder prediction.

## References

[1] A. M. Lesk: Introduction to Bioinformatics, 3rd ed. Oxford University Press, 2008
[2] G.N.Ramachandran, C. Ramakrishnan, V. Sasisekharan: Stereochemistry of polypeptide chain configurations, Journal of Molecular Biology. 7: 95-9. (1963)
[3] G. H. Reginald, C. M. Grisham: Biochemistry, fourth Edition, Belmont, CA: Brooks/Cole, 2013
[4] A. J. Cozzone: Proteins: Fundamental Chemical Properties, Institute of Biology and Chemistry of Proteins, CNRS, Lyon, France, 2002.
[5] P. Tompa, A. Fersht: Structure and Function of Intrinsically Disordered Proteins. Boca Raton: Chapman and Hall/CRC Taylor and Francis Group; 2010.
[6] DisProt Database - Database of protein disorder http://www.disprot.org/
[7] V. N. Uversky, A. K. Dunker: Understanding protein non-folding, Biochim Biophys Acta - Proteins \& Proteomics 2010, 1804(6):1231-1264.
[8] D. Eliezer: Biophysical characterization of intrinsically disordered proteins, Current Opinion in Structural Biology 2009, 19:23-30
[9] M. Punta, I. Simon, Z. Dosztanyi: Prediction and Analysis of Intrinsically Disordered Proteins, In Owens J R (ed.), Structural proteomics: High-Troughput Methods, Methods in Molecular Biology, vol. 1261, SpringerScience+Business Media New York, 2015, pp. 35-59.
[10] Z. Dosztányi, B. Mészáros, I. Simon: Bioinformatical approaches to characterize intrinsically disordered/unstructured proteins, Briefings In Bioinformatics, Vol 11. No 2, 225-243, 2009.
[11] B. Xue, R. L. Dunbrack, R. W. Williams, A. K. Dunker and V. N. Uversky: PONDR-FIT: A Meta-Predictor of Intrinsically Disordered Amino Acids. Biochim

Biophys Acta 1804(4):996-1010, 2010.
[12] M. Lobanov and O. Galzitskaya. The Ising model for prediction of disordered residues from protein sequence alone. Phys. Biol. 8 (2011) 035004 (9pp).
[13] P. Romero, Z. Obradovic, C. Kissinger, J. E. Villafranca, and A. K. Dunker. Identifying Disordered Regions in Proteins from Amino Acid Sequence. Proceedings of the 1997 IEEE International Conference on Neural Networks. Part 4, pp90-95 ( 1997).
[14] J. Flint, V. R. Racaniello, G. F. Rall, AM Skalka, L. W. Enquis: Principles of Virology, Garland science, Taylor \& Francis Group, USA, (2015)
[15] N. Tokuriki, C. J. Oldfield, V. N. Uversky, I. N. Berezovsky, D. S. Tawfik: Do viral proteins possess unique biophysical features?, Trends in Biochemical Sciences, 34, 53-59, (2008))
[16] B. Xue, A. K. Dunker, V. N. Uversky: Orderly order in protein intrinsic disorder distribution: disorder in 3500 proteomes from viruses and the three domains of life, Journal of Biomolecular Structure and Dynamics, 30, 137-149, (2012).
[17] D. Tauritz: Application of n-Grams, Department of Computer Science University of Missouri-Rolla; 2002.
[18] A. Jelović, N. Mitić, S. Eshafah, M. Beljanski: Finding statistically significant repeats in nucleic acids and proteins, Journal of Computational Biology, DOI: 10.1089/cmb.2017.0046
[19] P. Woolf, C. Burge, A. Keating, M. Yaffe: Statistics and Probability Primer for Computational Biologists, Massachusetts Institute of Technology, 2004
[20] PN. Tan, M. Steinbach, V. Kumar: Introduction to Data Mining, Pearson Education, 2006
[21] M. Kantardzic: Data mining : concepts, models, methods, and algorithms, John Wiley \& Sons, 2011
[22] IBM SPSS Modeler 18.0 Algorithms Guide, IBM Corporation 2016.
[23] Z. Dosztányi, V. Csizmok, P. Tompa, I. Simon: IUPred: web server for the prediction of intrinsically unstructured regions of proteins based on estimated energy content, Bioinformatics 21:3433-3434, 2005.
[24] K. Peng, P. Radivojac, S. Vučetić, AK . Dunker, Z. Obradović: Length-dependent prediction of protein intrinsic disorder, BMC Bioinformatics 7:208, 1-17, 2006.
[25] M. Yu Lobanov, I. V. Sokolovskiy, O. V. Galzitskaya: IsUnstruct: prediction of the residue status to be ordered or disordered in the protein chain by a method based on the Ising model, Journal of Biomolecular Structure and Dynamics 2013, 31(10), pp. 1034-1043
[26] M. Ganapathiraju, D. Weisser, J. Klein-Seetharaman, R. Rosenfeld, J. Carbonell, R. Reddy: Comparative n-gram analysis of whole-genome sequences. HLT'02: Human Language Technologies Conference: 2002 San Diego; 2002.
[27] H. U. Osmanbeyoglu, M. K. Ganapathiraju: N-gram analysis of 970 microbial organisms reveals presence of biological language models, BMC Bioinformatics 2011, 12:12.
[28] M. Ganapathiraju, A. Mitchell, M. Thahir, K. Motwani, S. Ananthasubramanian: Suite of Tools for Statistical N-gram language modeling for pattern mining in whole genome sequences, Journal of Bioinformatics and Computational Biology, Dec;10(6) 2012.
[29] G. Pavlovic-Lazetic, N. Mitic, M. Beljanski: n-Gram characterization of genomic islands in bacterial genomes, Computer Methods and Programs in Biomedicine, (2009), vol. 93 No. 3, pp. 241-256
[30] M. Yu. Lobanov, O. V. Galzitskaya: Occurrence of disordered patterns and homorepeats in eukaryotic and bacterial proteomes, Mol. BioSyst., 2012,8, 327-337.
[31] IBM corporation: Intelligent miner
https://www.ibm.com/support/knowledgecenter/SSEPGG_10.5.0/com.ibm.im.overview .doc/c_im_benefits.html
[32] Dynamic Warehousing: Data Mining Made Easy, SG24-7418-00, IBM corporation, 2007, http://www.redbooks.ibm.com/redbooks/pdfs/sg247418.pdf
[33] IBM InfoSphere Warehouse: Visualizing mining models, IBM Corporation, 2008, SH12-6840-03

## Appendix

## Table A1. Amino acid codes

| Amino acid names | One letter code | Three letter code* |
| :--- | :---: | :---: |
| Alanine | A | Ala |
| Asparagine or aspartic acid | B | Asx |
| Cysteine | C | Cys |
| Aspartic acid | D | Asp |
| Glutamic acid | E | Glu |
| Phenylalanine | F | Phe |
| Glycine | G | Gly |
| Histidine | H | His |
| Isoleucine | I | Ile |
| Leucine or Isoleucine | J | Xle |
| Lysine | K | Lys |
| Leucine | L | Leu |
| Methionine | M | Met |
| Asparagine | N | Asn |
| Pyrrolysine | O | Pyl |
| Proline | P | Pro |
| Glutamine | Q | Gln |
| Arginine | R | Arg |
| Serine | S | Ser |
| Threonine | T | Thr |
| Selenocysteine | U | Sec |
| Valine | V | Val |
| Tryptophan | W | Trp |
| Unspecified or unknown | X | Xaa |
| Tyrosine | Y | Tyr |
| Glutamine or glutamic acid | Z | Glx |
| N-Formylmethionine |  | fMet |

* N-Formylmethionine has only four-letter code


## Table A2: Summary of disorder-prediction methods

| Xue, B., R. L. DunBrack, R.W. Williams, A.K. Dunker, and V. N. <br> Uversky (2010) "PONDR-Fit: A meta-predictor of intrinsically <br> disordered amino acids." Biochim. Biophys. Acta 1804(4):996-1010, <br> PMID: 20100603 | $\underline{\underline{\text { PIT }}}^{\text {TM }}$ |
| :--- | :--- |
| Linding R, Jensen LJ, Diella F, Bork P, Gibson TJ, Russell RB. <br> "Protein disorder prediction: implications for structural proteomics." <br> Structure. 2003;11(11):1453-9, PMID: 14604535 | $\underline{\text { DisEMBL }^{\text {TM }}}$ |
| Ward JJ, Sodhi JS, McGuffin LJ, Buxton BF, Jones DT. "Prediction <br> and functional analysis of native disorder in proteins from the three <br> kingdoms of life." J Mol Biol. 2004;337(3):635-45, PMID: <br> 15019783 | DISOPRED2 |
| MacCallum B. "Order/Disorder Prediction With Self Organising <br> Maps." CASP 6 meeting, Online paper | $\underline{\text { DRIPPRED }}$ |
| Cheng J, Sweredoski M, Baldi P. "Accurate Prediction of Protein <br> Disordered Regions by Mining Protein Structure Data" Data Mining <br> and Knowledge Discovery. 2005; 11(3):213-222, Online Paper | $\underline{\text { DISpro }}$ |
| Prilusky J, Felder CE, Zeev-Ben-Mordehai T, Rydberg EH, Man O, <br> Beckmann JS, Silman I, Sussman JL. "FoldIndex: a simple tool to <br> predict whether a given protein sequence is intrinsically unfolded." <br> Bioinformatics. 2005;21(16):3435-8, PMID: 15955783 | $\underline{\text { FoldIndex© }}$ |
| Linding R, Russell RB, Neduva V, Gibson TJ. "GlobPlot: Exploring <br> protein sequences for globularity and disorder." Nucleic Acids Res. <br> 2003;31(13):3701-8, PMID: 12824398 | $\underline{\text { GlobPlot 2 }}$ |
| Dosztanyi Z, Csizmok V, Tompa P, Simon I. "IUPred: web server <br> for the prediction of intrinsically unstructured regions of proteins <br> based on estimated energy content." Bioinformatics. <br> 2005;21(16):3433-4, PMID: 15955779 | IUPred |
| Romero P, Obradovic Z, Li X, Garner EC, Brown CJ, Dunker AK. <br> "Sequence complexity of disordered protein." Proteins. <br> 2001;42(1):38-48, PMID: 11093259 | PONDR |
| Coeytaux K, Poupon A. "Prediction of unfolded segments in a <br> protein sequence based on amino acid composition." Bioinformatics. <br> 2005;21(9):1891-900, PMID: 15657106 | $\underline{\text { PreLink }}$ |
| Yang ZR, Thomson R, McNeil P, Esnouf RM. "RONN: the bio-basis <br> function neural network technique applied to the detection of <br> natively disordered regions in proteins." Bioinformatics. <br> 2005;21(16):3369-76, PMID: 15947016 |  |
| RONN |  |


| Vullo A, Bortolami O, Pollastri G, Tosatto S. "Spritz: a server for the <br> prediction of intrinsically disordered regions in protein sequences <br> using kernel machines" Nucleic Acids Res. 2006;34(Webserver <br> Issue):W164-W168, PMID: 16844983 | SPRITZ |
| :--- | :--- |
| Garbuzynskiy SO, Lobanov MY, Galzitskaya OV. "To be folded or <br> to be unfolded?" Protein Sci. 2004;13(11):2871-77., PMID <br> 15498936 |  |
| Galzitskaya OV, Garbuzynskiy SO, Lobanov MY. "Prediction of <br> natively unfolded regions in protein chain." Mol Biol (Mosk). <br> 2006;40(2):341-8., PMID 16637275 | $\underline{\text { FoldUnfold }}$ |
| Vucetic S, Brown CJ, Dunker AK, Obradovic Z. "Flavors of protein <br> disorder." Proteins. 2003 Sep 1;52(4):573-84, PMID: 12910457 | $\underline{\text { VL2 }}$ |
| Obradovic Z, Peng K, Vucetic S, Radivojac P, Brown CJ, Dunker <br> AK. "Predicting intrinsic disorder from amino acid sequence." <br> Proteins. 2003;53 Suppl 6:566-72, PMID: 14579347 | $\underline{\text { VL3, VL3H, }}$ |
| Obradovic Z, Peng K, Vucetic S, Radivojac P, Dunker AK. <br> "Exploiting heterogeneous sequence properties improves prediction <br> of protein disorder." Proteins. 2005;61 Suppl 7:176-82, PMID: <br> 16187360 | $\underline{\text { VSL2 }}$ |
| M. Lobanov and O. Galzitskaya. " The Ising model for prediction of <br> disordered residues from protein sequence alone". Phys. Biol. 8 <br> (2011) 035004 (9pp). | $\underline{\text { IsUnstruct }}$ |
| Walsh,I., Martin,A.J., Di Domenico,T., and Tosatto, S.C. (2012) <br> ESpritz: accurate and fast prediction of protein disorder. <br> Bioinformatics., 28(4), 503-509. | ESpritz |

(partially reproduced from http://disorder.compbio.iupui.edu/predictors.php).

## Table A3: Distribution of proteins over phyla and classes



## Table A4. N -grams that occur only in disordered regions

For each length first 100 n -grams that appear only in disordered regions sorted according their mole fractions in descending order are presented, except for length 4 where only four such n-grams exist.

| N-gram length |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| HHHH | GGGGG | GGGGGG | GGGGGGG | SSSSSSSS | SSSSSSSSS | SSSSSSSSSS |
| SNAM | PPPPP | PPPPPP | PPPPPPP | GGGGGGGG | PPPPPPPPP | PEPEPEPEPE |
| GHMA | APAPA | TTTTTT | EEEEEEE | PPPPPPPP | PEPEPEPEP | EPEPEPEPEP |
| GSHM | PSPPP | PEPEPE | DDDDDD | EEEEEEEE | EPEPEPEPE | EEEEEEEEEE |
|  | NnNnN | EPEPEP | PEPEPEP | PEPEPEPE | EEEEEEEEE | KPAPKPAPKP |
|  | EEEED | GGGGGA | EPEPEPE | EPEPEPEP | PKPAPKPAP | PKPAPKPAPK |
|  | PPAPP | PKPAPK | PKPAPKP | DDDDDDDD | PAPKPAPKP | PPPPPPPPPPP |
|  | SSTSS | KPAPKP | TTTTTTT | PKPAPKPA | KPAPKPAPK | PAPKPAPKPA |
|  | KKKKK | AGGGGG | PPPSPPP | KPAPKPAP | GGGGGGGGG | APKPAPKPAP |
|  | DEEDE | PPSPPP | KPAPKPA | APKPAPKP | APKPAPKPA | DDDDDDDDD |
|  | PAPPP | APKPAP | PAPKPAP | PAPKPAPK | DDDDDDDDD | PSPPPPSPPP |
|  | KKSKK | PSPPPP | APKPAPK | TTtttttt | QQQQQQQQQ | PPSPPPPSPP |
|  | EEDDD | GGGGGS | QQQQQQQ | QGAKSSSD | PPPPSPPPP | PPPSPPPPSP |
|  | NSSSS | APAPAP | GGGGGGA | QQQQQQQQ | PSPPPPSPP | QQQQQQQQQQ |
|  | PPAAP | SGGGGG | PPSPPPP | PPPSPPPP | TTTTTTTTT | SPPPPSPPPP |
|  | KKEKK | PAPAPA | PPPPSPP | PPPPSPPP | PPSPPPPSP | GGGGGGGGGG |
|  | SKKKK | PPPPSP | MDSRTGE | RSRSRSRS | PPPSPPPPS | PPPPSPPPPS |
|  | ESSSS | NNNNNN | PAPAPAP | GGGGGGGA | SPPPPSPPP | RSRSRSRSRS |
|  | RRRGR | SRSRSR | GAKSSSD | PSPPPPPS | RSRSRSRSR | APAPAPAPAP |
|  | AAPPA | QGAKSS | QGAKSSS | PPSPPPPS | PAPAPAPAP | TTTTTTTTTT |
|  | GGGDD | SPPPPS | RSRSRSR | SPPPPSPP | SRSRSRSRS | PAPAPAPAPA |
|  | EEEEG | SDSDSD | AGGGGGG | AGGGGGGG | APAPAPAPA | SARGGQQTAN |
|  | ннннн | PQGPQG | GAGGGGG | APAPAPAP | KGDKGDKGD | SRSRSRSRSR |
|  | PPPPQ | DSDSDS | SRSRSRS | SRSRSRSR | NnNNNNNNN | PPSPPPSPPP |
|  | STTST | GGGGGR | GPQGPQG | GPQGPKGD | SARGGQQTA | NNNNNNNNNN |
|  | DSDEE | TGGGGG | PSPPPPS | PAPAPAPA | PSPPPSPPP | TTAATTTAAT |
|  | DEDSD | RGGGGG | NNNNNNN | NNNNNNNN | GGGGGGGGA | TAATTTAATT |
|  | DDDDK | GPQGPK | APAPAPA | KGDKGDKG | DSDSDSDSD | PPPSPPPSPP |
|  | DDDKD | PQGPKG | GGGGGGS | KGDKGDTG | PPPSPPPPP | DIVISTPASK |
|  | REEEE | SPPPSP | SPPPPSP | GAGGGGGG | PPSPPPSPP | AATTTAATTT |
|  | SPSPG | PSPPPS | SGGGGGG | SARGGQQT | RSARGGQQS | ADIVISTPAS |
|  | EKKKS | EEEEED | GPQGPKG | SDSDSDSD | TTAATTTAA | SARGGQQSAN |
|  | GGSRS | AAPAPA | GGGGGSG | SGGGGGGG | AATTTAATT | VISTPASKVR |
|  | SSSVD | PPPPPS | GGGGGAG | GGAGGGGG | RGGQQSAND | ARGGQQSAND |
|  | EAEED | APPPPP | GGGGAGG | SPPPSPPP | GAGGGGGGG | IVISTPASKV |
|  | PSPEP | SSSSSD | DSDSDSD | PPPSPPPPS | ATTTAATTT | RSARGGQQSA |
|  | NTERH | TTAATT | PQGPKGD | PSPPPSPP | DIVISTPAS | TTTAATTTAA |
|  | PQQQP | SPPPPP | PSPPPSP | DSDSDSDS | SDSDSDSDS | ATTTAATTTA |
|  | KKKAA | GGGGGY | SDSDSDS | PPSPPPSP | IVISTPASK | MSKRPADIVI |
|  | PAATS | SSSTSS | MSKRPAD | GGGGGGAG | ADIVISTPA | SDSDSDSDSD |
|  | TPEPP | QGPQGP | KGDKGDK | TNGIEPPR | SQLKGSSST | SKRPADIVIS |
|  | QQEEE | PPPPPA | DKGDKGD | GGGGGGGS | ARGGQQSAN | DSDSDSDSDS |
|  | KKTSS | MSKRPA | KGDKGDT | GGGGGAGG | TTTAATTTA | PADIVISTPA |
|  | PKPRP | SKRPAD | PPSPPPS | GPQGPQGP | SARGGQQSA | ELNPAPTSSP |
|  | RGEET | EEEEDE | DEDEDED | QGPKGDTG | VISTPASKV | RYGGGGGGGG |
|  | KPTPP | PKGDTG | TNGIEPP | TTAATTTA | AGGGGGGGG | NSTNGIEPPR |
|  | KRPPP | TGPQGP | PTPSPTP | TTTAATTT | SKRPADIVI | ISLGSGLSMS |
|  | APEDP | LPPPPP | GPAGPQG | ATTTAATT | MSKRPADIV | PADTPVSEIP |
|  | MEEEE | TPPPTP | SPPPSPP | SARGGQQS | ALRRRLERG | SSRASSRASS |
|  | THMPR | YGGGGG | GGGGSGG | GGQQSAND | MPKRDAPWR | SQLKGSSSTS |
|  | KKGKS | PPTPPP | GGSGGGG | RGGQQSAN | GGGGGGGAG | SILEEAQRLI |
|  | KSASS | SSSSGS | YGGGGGG | VISTPASK | LNPAPTSSP | ESILEEAQRL |
|  | QQPPQ | edeeee | SSSSSSD | IVISTPAS | SGGGGGGGG | ILEEAQRLIH |
|  | DSPPS | APAPAA | PTPPPTP | DIVISTPA | YGGGGGGGG | PPGPEEGEGP |
|  | PEPPS | GGGGGD | GGGGGGR | ARGGQQSA | SRASSRASS | NSGYRYGGGG |
|  | SSEKP | PGGGGG | TTAATTT | SPASMEGN | TGPQGPKGD | LEEAQRLIHG |
|  | DSPPP | PPAPPA | NGIEPPR | QLKGSSST | DKGDKGDTG | SGYRYGGGGG |
|  | NKGPE | NGIEPP | GSGGGGG | GGGGGGSG | EEQKQLTLF | SSQVSNSTNG |
|  | AQAQE | SPSPPP | GPEGPEG | EDEDEDED | QGPKGDKGD | GPPGPEEGEG |
|  | GPSSG | GGGGGV | TTTAATT | GPEGPEGP | SSRASSRAS | YRYGGGGGGG |
|  | SPEPP | PAGPQG | PQGPQGP | EGPEGPEG | STNGIEPPR | GYRYGGGGGG |
|  | SQPEE | SPSPSP | PPPPLPP | YGGGGGGG | GGAGGGGGS | DISLGSGLSM |
|  | LMPCE | GQQTAN | PPPPPPS | AGTSKVSR | APAAPAAPA | PCESSSQVSN |
|  | GPLGS | PPPTPP | SPPPPPP | SKRPADIV | TNGIEPPRG | KGDKGDKGDT |
|  | KRPGP | EDEEDE | DDEDDED | SSSSSSSD | GDGDGDGDG | GDGDGDGDGD |
|  | PKRPR | GGGGGN | GGGGGGY | QGPKGDKG | RYGGGGGGG | SSSQVSNSTN |
|  | VASMQ | PTPPPT | QPEESVG | DEDEDEDE | NSTNGIEPP | QLKGSSSTSS |
|  | KGPPY | PPPPAP | SSRASSR | AGGGGGSG | PAPVPKPAP | MPCESSSQVS |
|  | VKGPP | QGIQGP | PPPTPPP | ASSRASSR | ILEEAQRLI | SRASSRASSR |
|  | QQPQA | EDEDEE | QGPKGDT | GAGGGGGS | GAGGGGGSG | STNGIEPPRG |
|  | GVPRG | QPEESV | ASSSSSS | MPKRDAPW | ADTPVSEIP | CESSSQVSNS |
|  | QPRRR | EGPEGP | SSSSSSA | RASSRASS | RASSRASSR | GEGGEGGEGG |
|  | AHSTQ | SSSSDS | PSSSSSS | TSSSSSSS | PADTPVSEI | DGDGDGDGDG |
|  | EPRHH | PAPPPP | QPQPEES | GPTGPTGP | QLKGSSSTS | SNSTNGIEPP |
|  | ESPPP | APTSSP | TGGGGGG | LRRRLERG | ISLGSGLSM | ESSSQVSNST |


|  | PKPPE | PPPPPL | RGGGGGG | NPAPTSSP | DGDGDGDGD | GGEGGEGGEG |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | QQTQQ | SRASSR | ARGGQQS | ALRRRLER | DISLGSGLS | PSPPPSPPPS |
|  | DDQAS | PEGPEG | DDDDDDE | SRASSRAS | SLGSGLSMS | TDISLGSGLS |
|  | EPEEM | VGGGGG | GGQQSAN | PKRDAPWR | PGPEEGEGP | QTANDAAAEA |
|  | PQSPS | GPEGPE | GQQSAND | GPQGIQGP | SSQVSNSTN | GGAGGGGGSG |
|  | QPPRR | SSSSSE | RGGQQSA | GGGGGGGY | SILEEAQRL | AGGGGGSGRR |
|  | SQPSQ | TPPPPP | EDDEDDE | PAPVPKPA | GGGGGSGRR | SQVSNSTNGI |
|  | EMNRQ | NGGGGG | AGGGGGS | GDGDGDGD | GYRYGGGGG | GAGGGGGSGR |
|  | EQKES | RRSPSP | GPQGPAG | SSRASSRA | GPPGPEEGE | GGAGAGGGAG |
|  | GHMAS | GGGGGL | VISTPAS | DKGDKGDT | EEAQRLIHG | QVSNSTNGIE |
|  | MEGRE | AGPQGP | PVPKPAP | VQPQPEES | SQVSNSTNG | QSGTSARRAE |
|  | PASQP | DEEEED | LKGSSST | DEDDEDDE | SGYRYGGGG | EGGEGGEGGE |
|  | PSRPR | PPPPPT | PSPPPPP | PPPLPPPP | GAGAGGGAG | LMPCESSSQV |
|  | QPPEE | QQQQQP | IVISTPA | EEQKQLTL | ESILEEAQR | RHKLAEKRAR |
|  | SPPQP | PSPSPS | SPASMEG | TGPQGPKG | GEGGEGGEG | ATDISLGSGL |
|  | AQQQT | GGGGGT | PASMEGN | APTSSPTS | PPGPEEGEG | ALRRRLERGE |
|  | EPKKP | RSPSPR | PGGGGGG | APVPKPAP | NSGYRYGGG | PAAPAAPAAP |
|  | KGPEQ | DEEDEE | RRRSSGG | STNGIEPP | LEEAQRLIH | VSNSTNGIEP |
|  | QQQAS | EEDEDE | ENTERHT | GPAGPQGP | YRYGGGGGG | ASSRASSRAS |
|  | QREQM | GGGGGP | KRDAPWR | NGIEPPRG | NATNGIEPP | GNEMVLPAET |
|  | RYCRK | QPQPEE | KRPADIV | RYGGGGGG | CESSSQVSN | MVLPAETRPG |
|  | SPEPA | PPPPTP | PPLPPPP | PSSSSSSS | SNSTNGIEP | QATEFDSPFA |
|  | AGHQQ | TTTPTT | EGPEGPE | NSTNGIEP | TANDAAAEA | VLPAETRPGA |
|  | RQQQE | KKKKKK | GTSKVSR | ISLGSGLS | SSSQVSNST | NGAAAREQAT |
|  | RRHHH | EEEEDD | EEEEEED | GGEGGEGG | PCESSSQVS | TEFDSPFADR |

## Table A5. N-grams with positive disorder fractional difference

Table includes for each length first 100n-grams occurring both in disordered and ordered regions with positive disorder/order fractional difference sorted according mole fractions in descending orders, except for length one where 11 monograms exists.

| N -gram length |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| S | SS | SSS | GGGG | SSSSS | SSSSSS | AKSSSDV | AKSSSDVK | HPNIQGAKS | FHPNIQGAKS |
| E | EE | GGG | SSSS | EEEEE | DDDDD | KSSSDVK | PNIQGAKS | FHPNIQGAK | AHFHPNIQGA |
| A | AA | PPP | PPPP | QQQQQ | EEEEEE | NIQGAKS | HPNIQGAK | HFHPNIQGA | SAHFHPNIQG |
| K | KK | EEE | DDDD | PEPEP | PAPKPA | PNIQGAK | RSARGGQQ | GRSARGGQQ | DGRSARGGQQ |
| R | GG | RRR | EEEE | PPPSP | KGDKGD | HPNIQGA | FHPNIQGA | AHFHPNIQG | IDGRSARGGQ |
| P | AS | DDD | APAP | PAPKP | RSRSRS | SARGGQQ | HFHPNIQG | DGRSARGGQ | AVSQLKGSSS |
| D | RR | PAP | PAPA | PKPAP | AKSSSD | RSARGGQ | GRSARGGQ | TPASKVRRR | TPASKVRRRL |
| G | SA | APA | QQQQ | PPSPP | MDSRTG | GPKGDKG | SQLKGSSS | TAATTTAAT | SAVSQLKGSS |
| T | EA | KKK | PPPS | SPPPP | KSSSDV | ISTPASK | TPASKVRR | VSQLKGSSS | ISASAYNGND |
| Q | AE | SPS | PPAP | PPPPS | DSRTGE | GDKGDTG | PASKVRRR | AVSQLKGSS | SASAYNGNDT |
| M | KE | PSP | PSPP | KPAPK | GPQGPQ | TPASKVR | AATTTAAT | PASKVRRRL | PASKVRRRLN |
|  | SG | PSS | PEPE | SRSRS | IQGAKS | QLKGSSS | TAATTTAA | SAVSQLKGS | ISIRTFRELN |
|  | PP | PTP | PPSP | QGPQG | NIQGAK | SQLKGSS | VSQLKGSS | IIISTPASK | ASkVRrRLNF |
|  | SE | EED | SPPP | APKPA | PNIQGA | EDEDEDE | AVSQLKGS | ISASAYNGN | IEQSVISASA |
|  | PS | SSP | GPQG | PPPPA | DEDEDE | PASKVRR | KSYIDKDG | SASAYNGND | EQSVISASAY |
|  | SK | PPA | APPP | PPPTP | EDEDED | APAAPAA | ASkVRRRL | ASKVRRRLN | QSVISASAYN |
|  | EK | PPS | PAPP | RRRSS | SARGGQ | AAPAAPA | SAVSQLKG | ASAYNGNDT | SVISASAYNG |
|  | ES | APP | PSSS | PSPSP | ARGGQQ | GPQGIQG | GGAGAGGG | SIRTFRELN | VISASAYNGN |
|  | DE | KRK | EEED | QGPKG | TSSSSS | SDWSFLK | IIISTPAS | IEQSVISAS | RPMNRKPRMY |
|  | GS | SRS | EPEP | RSPSP | DEEEEE | SDVKSYI | IISTPASK | SVISASAYN | PMNRKPRMYR |
|  | DD | SES | PTPP | PPPPT | PAAPAA | GPTGPTG | GAGAGGGA | QSVISASAY | LSAVSQLKGS |
|  | SD | PEP | RRRS | QGAKS | ASSSSS | ASKVRRR | IEQSVISA | KSYIDKDGD | NLSAVSQLKG |
|  | ST | TPP | PPPA | MDSRT | PKGDKG | ATTTAAT | EQSVISAS | EQSVISASA | STHFHPNIQG |
|  | AP | SSE | SPSP | SPSPS | DDEDDE | KSYIDKD | VISASAYN | VISASAYNG | SSTWYPQPGQ |
|  | KS | RRS | PPTP | PQGPQ | PSSSSS | ASSRASS | SVISASAY | MNRKPRMYR | LNERTATETR |
|  | PA | APS | SSSP | SPSPP | DDEDED | GGRGGGG | QSVISASA | RPMNRKPRM | KLNERTATET |
|  | KA | EES | PAPK | PAPEP | DKGDKG | NDDDDDD | NRKPRMYR | PMNRKPRMY | EDIKGYKPHT |
|  | AK | ESS | SSPS | PPPPR | GDKGDK | AVSQLKG | TGPTGPTG | LSAVSQLKG | IEDIKGYKPH |
|  | AR | RKR | APKP | MSKRP | SSSSSA | GGGAGAG | PMNRKPRM | NLSAVSQLK | ANLSAVSQLK |
|  | SP | RSS | PSPS | PQGPK | GPKGDK | ALRRRLE | TTFKDSTG | LTASDWSFL | NGNIHVSKLP |
|  | ED | QQQ | KPAP | RRSSS | STPASK | SAVSQLK | MNRKPRMY | THFHPNIQG | LIAARGYVYT |
|  | TS | RSR | PKPA | PPRPP | ISTPAS | AGAGGGA | TEFDSPFA | KLNERTATE | EFGFDGGDSE |
|  | RA | SPP | PPPT | PQPQP | DKGDTG | DDDDDD | LSAVSQLK | DIKGYKPHT | AARGYVYTAA |
|  | SR | PKP | PRRR | QQQQP | GDKGDT | SSSSSAS | GDKGDTGA | NERTATETR | ENGNIHVSKL |
|  | KR | EPE | PAPS | PPQPP | PLPPPP | ESILEEA | AGGAGAGG | LNERTATET | AVLIAARGYV |
|  | ER | KPK | PSSP | QPQPQ | TPSPTP | IISTPAS | TTGLSKAK | SSTWYPQPG | VLIAARGYVY |
|  | RS | RRK | PPGP | PPPEP | PTPSPT | IIISTPA | GFDGGDSE | EDIKGYKPH | IAARGYVYTA |
|  | PE | PRP | PPSS | PQQQQ | LKGSSS | GDDDDDD | SSTWYPQP | DAEQRELLD | TVTITADVRD |



## Table A6. N -grams that appear only in ordered regions

For each length first 100 n -grams that appear only in ordered regions sorted according their mole fractions in descending order are presented, except for length 3 where only 10 such n-grams exist.

| N-gram length |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |  |
| WIC |  | IFII | YNVID | IKGGIP | LYMACTH | LYMACTHA | THASNPVYA |  |
| YCW | FINY | IKGGI | NVIDDV | YMACTHA | HASNPVYA | YMACTHASN | THASNPVASN |  |
| WCY | FVFL | VGKRF | YNVIDD | QIKGGIP | MACTHASN | HASNPVYAT | CTHASNPVYA |  |
| WYW | ILYV | ATLKI | YMACTH | ASNPVYA | NPVYATLK | SNPVYATLK | HASNPVYATL |  |
| CWF |  | FIII | ACTHA | LYMACT | ACTHASN | SNPVYATL | ASNPVYATL |  |
| FWW | LLLW | GKRFC | QIKGGI | SNPVYAT | ASNPVYAT | THRVGKRFC | NPVYATLKIR |  |


| CWY | YILV | LYMAC | SNPVYA | NPVYATL | THRVGKRF | NPVYATLKI | SNPVYATLKI |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| CYW | LYVY | MACTH | MACTHA | PVYATLK | HRVGKRFC | PVYATLKIR | TLKIRIYFYD |
| HWW | VLAC | YMACT | NPVYAT | YNVIDDV | VYATLKIR | LKIRIYFYD | VYATLKIRIY |
| CWW | YYVL | ALLLY | CTHASN | NHTENAL | PVYATLKI | TLKIRIYFY | ATLKIRIYFY |
|  | VLLC | KYENH | PVYATL | THRVGKR | KIRIYFYD | YATLKIRIY | YATLKIRIYF |
|  | IFLC | GPHNY | NHTENA | HRVGKRF | LKIRIYFY | VYATLKIRI | PVYATLKIRI |
|  | CVLV | RFFDL | htenal | WMDENIK | YATLKIRI | ATLKIRIYF | NHTENALLLY |
|  | FVIF | PHNYL | THRVGK | RVGKRFC | ATLKIRIY | YENHTENAL | HRVGKRFCVK |
|  | FIVF | IYFYD | SDVTRG | LGPHNYL | TLKIRIYF | KYENHTENA | THRVGKRFCV |
|  | TMWA | KIRIY | HRVGKR | YATLKIR | GKRFCVKS | NHTENALLL | RVGKRFCVKS |
|  | YAYI | TMWAR | RVGKRF | VYATLKI | YENHTENA | HRVGKRFCV | HTENALLLYM |
|  | VFYL | RIYFY | GPHNYL | IRIYFYD | Enhtenal | HTENALLLY | KYENHTENAL |
|  | VVIF | LLYMA | VGKRFC | KIRIYFY | ENALLLYM | RVGKRFCVK | TENALLLYMA |
|  | IVIF | PLYFK | LGPHNY | LKIRIYF | KYENHTEN | TENALLLYM | ENALLLYMAC |
|  | LFIY | LYFKI | WMDENI | TLKIRIY | TENALLLY | VGKRFCVKS | ALLLYMACTH |
|  | VLYY | RFCVK | mDENIK | ATLKIRI | NHTENALL | ENALLLYMA | NALLLYMACT |
|  | YAIY | KIWMD | YATLKI | KRFCVKS | HTENALLL | LLYMACTHA | LLLYMACTHA |
|  | LYYY | GKIWM | ATLKIR | GKRFCVK | RVGKRFCV | NALLLYMAC | GKIWMDENIK |
|  | YYVV | LLLYM | RIYFYD | VTGGQYA | VGKRFCVK | LLLYMACTH | HNYLCGHLDL |
|  | FAII | IWMDE | IRIYFY | Yenhten | LLYMACTH | ALLLYMACT | GGIPTIFLCN |
|  | LICL | NNVIR | KIRIYF | Enhtena | NALLLYMA | GKIWMDENI | KGGIPTIFLC |
|  | YYIL | NYLCG | LKIRIY | ENALLLY | GKIWMDEN | KIWMDENIK | KHFKEFMGAQ |
|  | VAYY | HNYLC | NPLYFK | KYENHTE | LLLYMACT | HNYLCGHLD | IKGGIPTIFL |
|  | WYVD | IFLCN | RFCVKS | HTENALL | ALLLYMAC | NYLCGHLDL | LKHFKEFMGA |
|  | LLWL | LCGHL | GKRFCV | NALLLYM | KIWMDENI | GGIPTIFLC | NYLCGHLDLS |
|  | YINF | TIFLC | KYENHT | TENALLL | IWMDENIK | GIPTIFLCN | GIPTIFLCNP |
|  | IYIV | YLCGH | KRFCVK | VGKRFCV | HNYLCGHL | KGGIPTIFL | WARSLGPHNY |
|  | YLVF | AQRDW | VTGGQY | GKIWMDE | NYLCGHLD | LKHFKEFMG | ARSLGPHNYL |
|  | YIYT | KNYFL | GKIWMD | LLYMACT | YLCGHLDL | KHFKEFMGA | LGPHNYLCGH |
|  | YLYF | YLKHF | NALLLY | ALLLYMA | GIPTIFLC | HFKEFMGAQ | YLCGHLDLSP |
|  | YFTF | HFKEF | ENALLL | KIWMDEN | GGIPTIFL | IKGGIPTIF | GPHNYLCGHL |
|  | YIGF | NYFLT | ALLLYM | LLLYMAC | IPTIFLCN | YLCGHLDLS | YGKPVQIKGG |
|  | IAWL | FMGAQ | YENHTE | IWMDENI | WARSLGPH | WARSLGPHN | KYGKPVQIKG |
|  | LVFY | FLCNP | ENHTEN | HNYLCGH | KHFKEFMG | GPHNYLCGH | PHNYLCGHLD |
|  | FVFI | FFDLV | TENALL | NYLCGHL | KGGIPTIF | ARSLGPHNY | KPVQIKGGIP |
|  | YGIF | CRELH | KIWMDE | YLCGHLD | LKHFKEFM | IPTIFLCNP | GKPVQIKGGI |
|  | YIAI | YFLTY | LLYMAC | NPLYFKI | IKGGIPTI | RSLGPHNYL | GKRFCVKSVY |
|  | YYVY | NHNLR | LLLYMA | LCGHLDL | FKEFMGAQ | LCGHLDLSP | QIKGGIPTIF |
|  | VWLA | FGQVF | IWMDEN | PTIFLCN | HFKEFMGA | LGPHNYLCG | LGKIWMDENI |
|  | LVCV | LGKIW | HNYLCG | GGIPTIF | LCGHLDLS | GKPVQIKGG | SLGPHNYLCG |
|  | YFVI | LLLLV | NYLCGH | IPTIFLC | VTGGQYAS | YGKPVQIKG | RSLGPHNYLC |
|  | LLWF | WYNVI | LKHFKE | GIPTIFL | GPHNYLCG | PHNYLCGHL | Yenhtenall |
|  | VCVL | VVYNH | PLYFKI | LKHFKEF | ARSLGPHN | KYGKPVQIK | ENHTENALLL |
|  | YYYL | FNHNL | LCGHLD | KGGIPTI | PHNYLCGH | KPVQIKGGI | WYNVIDDVDP |
|  | IYFV | VISIN | YLCGHL | KHFKEFM | RSLGPHNY | GKRFCVKSV | LCGHLDLSPK |
|  | LHYY | AWYNV | CGHLDL | ARSLGPH | PTIFLCNP | PVQIKGGIP | YNVIDDVDPH |
|  | RIYY | IQFEG | IFLCNP | KEFMGAQ | CGHLDLSP | LGKIWMDEN | YLKHFKEFMG |
|  | CIAL | LILLL | GAQRDW | WARSLGP | WYNVIDDV | AWYNVIDDV | DDVDPHYLKH |
|  | FIAY | QVFNM | KHFKEF | HFKEFMG | SLGPHNYL | QIKGGIPTI | VGKRFCVKSV |
|  | LIFY | VYNHQ | GGIPTI | IKGGIPT | NVIDDVDP | KRFCVKSVY | IDDVDPHYLK |
|  | CLAI | DPHYL | PTIFLC | FKEFMGA | LGPHNYLC | NVIDDVDPH | AWYNVIDDVD |
|  | ICAL | FLRVF | TIFLCN | CGHLDLS | KPVQIKGG | YNVIDDVDP | HFKEFMGAQR |
|  | LTWL | HVLIQ | EFMGAQ | TGGQYAS | KYGKPVQI | SLGPHNYLC | KEFMGAQRDW |
|  | LYYF | LHVLI | KGGIPT | IDDVDPH | GKPVQIKG | ENHTENALL | FKEFMGAQRD |
|  | DIIC | VLIQF | GIPTIF | PHNYLCG | VQIKGGIP | WYNVIDDVD | DVDPHYLKHF |
|  | FYVI | ICREL | HFKEFM | LGKIWMD | YGKPVQIK | CGHLDLSPK | VDPHYLKHFK |
|  | TFIY | AGKYE | RSLGPH | RSLGPHN | KRFCVKSV | YLKHFKEFM | PHYLKHFKEF |
|  | YIPI | VVVVV | KEFMGA | GPHNYLC | LGKIWMDE | DDVDPHYLK | GKTMWARSLG |
|  | LAWI | RCMLA | GGQYAS | GHLDLSP | PVQIKGGI | DVDPHYLKH | HYLKHFKEFM |
|  | FIYF | GFRCM | FKEFMG | TIFLCNP | RFCVKSVY | KEFMGAQRD | VIDDVDPHYL |
|  | LCVI | HTNSV | GHLDLS | SLGPHNY | QIKGGIPT | EFMGAQRDW | DPHYLKHFKE |
|  | LFCL | FRCML | DDVDPH | WYNVIDD | AWYNVIDD | IDDVDPHYL | NVIDDVDPHY |
|  | YLFV | LIIGL | YNHQEA | KYGKPVQ | VIDDVDPH | FKEFMGAQR | CGHLDLSPKV |
|  | FAVY | CGCSY | VQIKGG | PVQIKGG | YNVIDDVD | GKTMWARSL | GHLDLSPKVY |
|  | YYEI | ILSLI | KPVQIK | NVIDDVD | KNYFLTYP | VDPHYLKHF | TMWARSLGPH |
|  | VWVV | LLVVL | GKPVQI | KPVQIKG | YLKHFKEF | DPHYLKHFK | MWARSLGPHN |
|  | LYML | GCGKT | PHNYLC | NYFLTYP | GHLDLSPK | PHYLKHFKE | KTMWARSLGP |
|  | IGYF | DAWYN | SLGPHN | YGKPVQI | DVDPHYLK | HYLKHFKEF | LHVLIQFEGK |
|  | CFAL | IIILL | HLDLSP | VIDDVDP | KEFMGAQR | VIDDVDPHY | GKYENHTENA |
|  | FTYI | YVLGK | WYNVID | RFCVKSV | DDVDPHYL | KTMWARSLG | GFTHRGTHHC |
|  | FYLY | GITHR | LGKIWM | VQIKGGI | VDPHYLKH | GHLDLSPKV | ITHRVGKRFC |
|  | YFLF | NDAWY | KNYFLT | GKPVQIK | EFMGAQRD | QEAGKYENH | GITHRVGKRF |
|  | FYIV | VHGFR | YGKPVQ | FCVKSVY | IDDVDPHY | HLDLSPKVY | QEAGKYENHT |
|  | IWEI | FLLLL | VIDDVD | AWYNVID | FMGAQRDW | HVLIQFEGK | AGKYENHTEN |
|  | YLCD | HGFRC | YFLTYP | YLKHFKE | GKTMWARS | TMWARSLGP | EAGKYENHTE |
|  | CLGI | ILLVL | PVQIKG | KNYFLTY | KTMWARSL | MWARSLGPH | DAWYNVIDDV |
|  | LFYY | QIRFN | NLDRIF | LIQFEGK | PHYLKHFK | LHVLIQFEG | LDLSPKVYSN |
|  | LYLC | VLLLV | NYFLTY | HLDLSPK | DPHYLKHF | GKYENHTEN | HLDLSPKVYS |
|  | IIFF | ILALL | FNHNLR | FGQVFNM | TMWARSLG | GFTHRGTHH | NDAWYNVIDD |
|  | LFLC | VVLAL | YLKHFK | EFMGAQR | HYLKHFKE | FTHRGTHHC | GLTHRVGKRF |
|  | FTFY | ALGIH | AWYNVI | DDVDPHY | HLDLSPKV | GITHRVGKR | LTHRVGKRFC |
|  | YIAF | GDLIY | CVKSVY | DVDPHYL | LDLSPKVY | ITHRVGKRF | SNDAWYNVID |
|  | CGLI | ILILL | FGQVFN | VDPHYLK | VLIQFEGK | EAGKYENHT | YSNDAWYNVI |
|  | CVIA | LAVLG | GQVFNM | MGAQRDW | EAGKYENH | AGKYENHTE | HGFTHRGTHH |
|  | LCYL | GNIIG | LIQFEG | DPHYLKH | QEAGKYEN | LDLSPKVYS | ENIKTKNHTN |
|  | LIGW | NYVVY | IQFEGK | FMGAQRD | HVLIQFEG | DLSPKVYSN | KRFCVKSVYI |
|  | VVYC | VLLLA | VYNHQE | VVYNHQE | VVYNHQEA | NDAWYNVID | LVRDRRPYGT |


|  | WLAI | ALVIL | DVDPHY | KTMWARS | MWARSLGP | DAWYNVIDD | DENIKTKNHT |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
|  | LKCF | GKVMC | VDPHYL | TMWARSL | LHVLIQFE | LVRDRRPYG | WQSNCKYGKP |
|  | FLCA | IILLL | NNVIRA | PHYLKHF | GKYENHTE | TVKNDLRDR | WMDENIKTKN |
|  | VMFF | LLLLG | FMGAQR | HYLKHFK | ITHRVGKR | KNHTNSVMF | MDENIKTKNH |
|  | YFKY | AVLLV | PHYLKH | MWARSLG | FTHRGTHH | LTHRVGKRF | RGNGITHRVG |
|  | IDWI | ISLLL | DPHYLK | LDLSPKV | GITHRVGK | FWLVRDRRP | NGITHRVGKR |
|  | CFLT | LGVVA | MGAQRD | QEAGKYE | THRGTHHC | GLTHRVGKR | GNGITHRVGK |

## Table A7. N-grams with positive order fractional difference

Table includes for each length first 100 n-grams occurring both in disordered and ordered regions with positive order/disorder fractional difference sorted according their mole fractions in descending orders, except for length one where 9 monograms exists.

| N -gram length |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| L | LL | YII | LIVL | LLLLL | SRTGKT | GPCKVQS | CEGPCKVQ | CEGPCKVQS | GCEGPCKVQS |
| v | VL | YIY | LLLF | NVIDD | GPCKVQ | CEGPCKV | EGPCKVQS | GCEGPCKVQ | DNEPSTATVK |
| I | LV | YFY | DIIL | KYGKP | VYATLK | EGPCKVQ | GCEGPCKV | QSNTKYGKP | VPRGCEGPCK |
| N | LI | CVI | LVIL | ASNPV | PCKVQS | GCEGPCK | EGDSRTGK | FDNEPSTAT | RGCEGPCKVQ |
| F | VV | IIC | ILVL | VYATL | YGDTDS | GDSRTGK | SNTKYGKP | RGCEGPCKV | CEGPCKVQSY |
| Y | IL | YCL | YILN | PVYAT | WARSLG | EGDSRTG | QSNTKYGK | EGPCKVQSY | MFDNEPSTAT |
| H | VI | CYL | NLIV | YAtLK | VRDRRP | SNTKYGK | FDNEPSTA | MFDNEPSTA | PRGCEGPCKV |
| C | IV | FCV | VYVL | THASN | GFTHRG | NTKYGKP | MFDNEPST | GPCKVQSYE | EGPCKVQSYE |
| W | II | CVF | IGII | DVTRG | TGGQYA | QSNTKYG | NTKYGKPV | DVPRGCEGP | DVPRGCEGPC |
|  | IG | WLY | IIFL | NPLYF | DSRTGK | HRGTHHC | KEEALSQL | CKVQSYEQR | FDNEPSTATV |
|  | LF | LIW | VLAY | HTENA | GDSRTG | FDNEPST | QSYEQRHD | PCKVQSYEQ | PCKVQSYEQR |
|  | FL | YVC | VFID | WMDEN | KYGKPV | MFDNEPS | VQSYEQRH | SNTKYGKPV | GPCKVQSYEQ |
|  | YL | FWL | INFI | THRVG | IDDVDP | TKEEALS | RGTHHCSS | KVQSYEQRH | PDVPRGCEGP |
|  | LY | WIV | YKII | LGVIS | ARSLGP | RELHEDG | HRGTHHCS | VQSYEQRHD | QSNTKYGKPV |
|  | FV | CVY | IILV | MDENI | FCVKSV | TKYGKPV | KFLNQVNA | HRGTHHCSS | CKVQSYEQRH |
|  | VF | VWI | LYYL | LGPHN | GAGKST | QSYEQRH | DEYQLSHD | DVPKGCEGP | KVQSYEQRHD |
|  | FI | CFV | IVLF | KNHTN | TKYGKP | SYEQRHD | VPKGCEGP | RKFLNQVNA | PDVPKGCEGP |
|  | VY | YIC | IIIA | FCVKS | NTKYGK | GTHHCSS | RKFLNQVN | KEEALSQLQ | VGSGKSTGLP |
|  | YV | IYC | KYII | FTHRG | SNTKYG | RGTHHCS | SSYKEFLD | VEGDSRTGK | ESRRKFLNQV |
|  | YI | ICF | LYIL | INNVI | LDLSPK | LGIHSPS | VELEGVNG | ESRRKFLNQ | RRKFLNQVNA |
|  | GY | WFL | LFIL | QFEGK | QSNTKY | NGAGKST | ESRRKFLN | RDEYQLSHD | SRRKFLNQVN |
|  | FT | WLF | VVVF | YGKPV | NDLRDR | YgdTdSv | KVDGRTMK | LVAEVERLR | ELVAEVERLR |
|  | YA | LFW | GIII | LKHFK | RGTHHC | KVDGRTM | SRRKFLNQ | ELVAEVERL | HRDEYQLSHD |
|  | AY | CYI | NYIL | ENHTE | HRGTHH | SYKEFLD | SYKEFLDE | RRKFLNQVN | ESHRDEYQLS |
|  | YG | YCI | LIFL | ILLLL | SIVIEG | SSYKEFL | ELVAEVER | SRRKFLNQV | RHPNISQLST |
|  | IF | VCY | LIVF | IVIEG | SLTKEE | SSYKEYL | LVAEVERL | SSYKEFLDE | SHRDEYQLSH |
|  | IY | CYV | YFYD | GAQRD | ATVTGG | FLNQVNA | RHPNISQL | IGVVKPLAI | IESHRDEYQL |
|  | YT | WII | YALV | KEFMG | VKNDLR | KFLNQVN | VAEVERLR | ESHRDEYQL | IVEGDSRTGK |
|  | YN | YVW | VAII | LLLLI | GLTHRV | ELVAEVE | RRKFLNQV | HRDEYQLSH | NYIESHRDEY |
|  | TY | CFI | AYVV | CVKSV | ELHEDG | GAGFGAG | IGVVKPLA | RHPNISQLS | YIESHRDEYQ |
|  | NY | IIW | LLIF | HLDLS | VIEGDS | PNSSYKE | LSSFTNVP | SHRDEYQLS | IGVVKPLAIT |
|  | FF | WFI | NILY | LALLL | VYSNDA | RKFLNQV | GVVKPLAI | YIESHRDEY | PMYRKPRMYR |
|  | YF | VWY | VLLY | TVVDN | SNLDRI | ELEGVNG | ESHRDEYQ | FVKTLTGKT | MSAEVLDRTK |
|  | YY | FWT | VLYA | IRDLI | DENIKT | PTSSYKE | HRDEYQLS | IESHRDEYQ | RPMYRKPRMY |
|  | FY | WVF | LYIT | SNTKY | STVVDN | RRKFLNQ | SHRDEYQL | IVEGDSRTG | SAEVLDRTKQ |
|  | LC | YFC | YIIE | LLAVL | AKFKGK | TKNTFSL | YIESHRDE | NYIESHRDE | GIGVVKPLAI |
|  | WL | VYW | GYIL | LLVEL | LGRVGR | VDGRTMK | YNNRWVKD | GVVKPLAIT | MYRKPRMYRM |
|  | HI | IWI | IFLK | IIIDE | GKLKLS | VELEGVN | IESHRDEY | KTLTGKTIT | PNSSYKEFLD |
|  | CL | YCF | VIGF | IGAGI | SYEQRH | ESRRKFL | VKTLTGKT | MYRKPRMYR | SSYKEFLDEE |
|  | IH | FWV | IRIY | HYLKH | HQEAAK | KGTVKIE | NYIESHRD | PMYRKPRMY | GVVKPLAITN |
|  | vc | FYC | TFVL | ALVLA | YEQRHD | SRRKFLN | VVKPLAIT | AEVLDRTKQ | YKEFLDEEKN |
|  | AW | YWV | LIYL | MGAQR | ASNEQA | LVAEVER | KTLTGKTI | MSAEVLDRT | DNEPSTATIK |
|  | GW | FWI | TIVF | LKLST | NGVTLD | YKEFLDE | MYRKPRMY | SAEVLDRTK | EPSTATIKND |
|  | cV | FCY | LVLY | ALLLT | THHCSS | VAEVERL | TLTGKTIT | SYKEFLDEE | NEPSTATIKN |
|  | CG | WIY | YFLT | VTLAL | GTHHCS | KTLTGKT | YKEFLDEE | GIGVVKPLA | SYKEFLDEEK |
|  | LW | CFY | LYLF | HRGTH | LPATAD | LSSFTNV | AEVLDRTK | PNSSYKEFL | NSSYKEFLDE |
|  | CA | YIW | NVII | INNIK | LRVLAA | SSFTNVP | EALSQLQN | NEPSTATIK | SGIGVVKPLA |
|  | WA | CYF | RIYF | GTHHC | RKALGI | QSGLDFK | EVLDRTKQ | NSSYKEFLD | SSFTNVPDEM |
|  | AC | WYI | RILI | AAVLL | SHVGKV | RHPNISQ | MSAEVLDR | KEFLDEEKN | LSSFTNVPDE |
|  | DW | DWW | FVVV | LVALT | GVSSRG | GAGLGAG | SAEVLDRT | VVKPLAITN | NSGIGVVKPL |
|  | WV | FIW | LYMA | ALVAG | LKDPIP | NNRWVKD | EFLDEEKN | YKEFLDEEK | RKPRIYRTLR |
|  | VW | FWF | YIVD | LLALI | HENGEP | GVVKPLA | EPSTATIK | AKEAFHPMY | FVKTLTGKTI |
|  | IC | IWY | IILF | VNGVL | IQIKGG | IGVVKPL | GIGVVKPL | DNEPSTATI | VKTLTGKTIT |
|  | CD | VWC | IINY | LGLLL | LKAELR | AGKSLIQ | NSSYKEFL | EEALSQLQN | EEALSQLQNL |
|  | GC | WYF | IIFN | NHTNS | PAGTGK | DDIDDID | PNSSYKEF | EPSTATIKN | EGGQHLNVNV |
|  | HF | FFW | YLVN | INNII | STAKHS | ESHRDEY | ALSQLQNL | PSTATIKND | FLEKISIPRG |



## Table A8. $\mathbf{N}$-grams that appear only in border between disordered and ordered regions

For each length first 100 n -grams that appear only on border between ordered and disordered regions sorted according their mole fractions in descending order are presented, except for length 4 where only 1 such n-gram exists.

| N-gram length |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| MWCW | IWRFP | FYDSVT | YFYDSVT | YFYDSVTN | MEGNRPTFV | ASMEGNRPTF |
|  | GPAWY | NRPTFV | FYDSVTN | EGNRPTFV | SMEGNRPTF | SMEGNRPTFV |
|  | PAWYW | VVYKYE | EGNRPTF | MEGNRPTF | LYDALEAPA | LYDALEAPAD |
|  | HAWMP | VYKYEE | GNRPTFV | QVVYKYEE | GQVVYKYEE | IRIYFYDSIT |
|  | HQQLW | CHLKNP | QVVYKYE | LYDALEAP | IYFYDSITN | KNYGHPRENF |
|  | HWMEI | FYDSIT | VVYKYEE | YFYDSITN | KNYGHPREN | NKNYGHPREN |
|  | WMEIP | GHPREN | FYDSITN | YGHPRENF | NYGHPRENF | RIYFYDSITN |
|  | SWWRH | HPRENF | GHPRENF | GQVVYKYE | RIYFYDSIT | EGNRPTFVVQ |
|  | WADHG | ICHLKN | YFYDSIT | IYFYDSIT | EGNRPTFVV | GNRPTFVVQN |
|  | HGMPD | DLDYVG | YGHPREN | NYGHPREN | GNRPTFVVQ | IRIYFYDSVT |
|  | MVVFK | MKKIIL | ICHLKNP | VICHLKNP | NRPTFVVQN | MEGNRPTFVV |
|  | AEKTH | PTFVVQ | VICHLKN | GNRPTFVV | PTFVVQNET | NRPTFVVQNE |
|  | IYWGM | HNTRDG | NRPTFVV | IYFYDSVT | RIYFYDSVT | PTFVVQNETQ |
|  | MDEDH | WVTLGG | PTFVVQN | NRPTFVVQ | RPTFVVQNE | RPTFVVQNET |
|  | MDINW | YDSVQN | RPTFVVQ | PTFVVQNE | IYFYDSVTN | RIYFYDSVTN |
|  | MPFRD | HPNLRM | VYKYEEE | RPTFVVQN | QVVYKYEEE | WVTLGGAGGG |
|  | NCYDR | LYIPEQ | EFAPDAP | VVYKYEEE | KEFAPDAPL | DGKRVSPPRE |
|  | PWNIQ | PNLRML | FAPDAPL | EFAPDAPL | WVTLGGAGG | DREPDLYIPE |



## Table A9. N -grams with positive fractional difference on border between disordered and ordered regions

Table includes for each length first 100 n -grams that appear on border between ordered and disordered regions, and in ordered or disordered regions or both, but prefer border region, sorted according their mole fractions in descending orders, except for length two where 78 bigrams exists.

| N-gram length |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| EL | MIY | DWWE | YFYDS | YFYDSV | RIYFYDS | IRIYFYDS | KIRIYFYDS | LKIRIYFYDS |
| LS | NIY | KFQI | FYDSI | IYFYDS | IYFYDSV | RIYFYDSV | IRIYFYDSV | KIRIYFYDSV |
| SL | WVS | GHWL | RPTFV | YFYDSI | LVSPTRS | DLVSPTRS | FDLVSPTRS | KIRIYFYDSI |
| LE | MFY | PAWY | LYIPE | LVSPTR | IYFYDSI | RIYFYDSI | IRIYFYDSI | VSPTRSAHFH |
| LK | MYY | GGHW | VYKYE | YDSVTN | TGELITA | RTGELITA | DSRTGELIT | FFDLVSPTRS |
| KL | WVK | AWYW | YKYEE | IKFNLY | VSPTRSA | SRTGELIT | DLVSPTRSA | MDSRTGELIT |
| IE | WEV | HWLG | HLKNP | NDTEGL | GQPSTVV | LVSPTRSA | LVSPTRSAH | FDLVSPTRSA |
| KI | WIE | DSHW | YDSIT | DTEGLL | SRTGELI | VSPTRSAH | VSPTRSAHF | DLVSPTRSAH |
| IK | HII | FVLQ | DLDYV | TEGLLK | DLVSPTR | SGQPSTVV | SRTGELITA | LVSPTRSAHF |
| SV | KIC | YVVI | WDPLV | YDSITN | NSGQPST | DSRTGELI | GQPSTVVDN | DSRTGELITA |
| EI | VWE | FHEM | GQLGI | MWDPLV | YDALEAP | GQPSTVVD | MDSRTGELI | SGQPSTVVDN |
| VS | MWL | HWTF | PTFVV | PSTVVD | DALEAPA | FDLVSPTR | SGQPSTVVD | GNNSGQPSTV |
| VE | WIK | TIYI | DLYIP | RGPAGW | QPSTVVD | NNSGQPST | GNNSGQPST | NNSGQPSTVV |
| DL | WFK | WNLH | GHPRE | NMFDNE | DTEGLLK | NSGQPSTV | NNSGQPSTV | KSYIDKDGDT |
| RL | WVD | WRHK | WVTLG | RPWGLE | GNDTEGL | YDALEAPA | NSGQPSTVV | NSGQPSTVVD |
| SI | QLW | CWGL | QAYIN | APDAPL | NDTEGLL | DALEAPAD | YDALEAPAD | YDALEAPADT |
| LR | AVW | TKMW | YDSVQ | SDAIDL | NGNDTEG | PLLNEFPE | DALEAPADT | DALEAPADTP |
| EV | YWE | KSCY | YHNTR | YDNEPS | NMFDNEP | ALEAPADT | DPLLNEFPE | WDPLLNEFPE |
| KV | CYD | WMEI | YIPEQ | GQLGIL | SSDVKSY | QPSTVVDN | ALEAPADTP | CCCPHCPRHK |
| IS | WKV | KMWQ | EPEFI | RPTFVV | TEGLLKE | SSDVKSYI | CCPHCPRHK | KSSSDVKSYI |
| VK | WIP | KQFW | INLVM | WDPLVN | MEGNRPT | LEAPADTP | SSSDVKSYI | RFFDLVSPTR |
| LP | MIW | SFWV | MFKKW | YKYEEE | PSTVVDN | DTEGLLKE | FFDLVSPTR | AYNGNDTEGL |
| PL | CIN | WWRH | RHLID | AISIRK | SRGPAGW | GNDTEGLL | AYNGNDTEG | GNDTEGLLKE |
| QL | HTW | FFEL | EKYYL | DTLVEL | FNMFDNE | NDTEGLLK | GNDTEGLLK | GQPSTVVDNT |
| RI | WID | GMWM | FVRPP | DAIDLI | TLSDAID | NGNDTEGL | NDTEGLLKE | NGNDTEGLLK |
| PV | WIR | MISW | HCTQV | FAPDAP | YDNEPST | YNGNDTEG | NGNDTEGLL | SAYNGNDTEG |
| VR | IWS | WNIQ | KYYLY | YGHPRE | MWDPLVN | FNMFDNEP | YNGNDTEGL | YNGNDTEGLL |
| RV | CEF | AWFA | LINLV | YHNTRD | SDAIDLI | NMFDNEPS | VFNMFDNEP | DPLLNEFPET |
| FS | KWI | CMVK | WRFPS | DLYIPE | DAISIRK | SSSDVKSY | FNMFDNEPS | FNMFDNEPST |
| LQ | WYE | ELWG | ASYAF | FDLDYV | DAIDLIN | SMEGNRPT | NMFDNEPST | NMFDNEPSTA |
| SF | CIR | FYYK | ATIFD | FYDSVQ | EAPADTP | VFNMFDNE | PLLNEFPET | VFNMFDNEPS |
| IR | MCI | HILA | GSPIW | GGRVKP | LSDAIDL | MTLSDAID | KSSSDVKSY | PLLNEFPETV |
| VP | MIC | NPQW | LIPSC | PDLYIP | GGRVKPL | YDNEPSTA | ASMEGNRPT | AKSSSDVKSY |
| IP | WVN | QVHC | LRGYN | PREVRI | NYGHPRE | SDAIDLIN | LRAVLTEAL | HFHPNIQGAK |
| FK | CYE | WWGG | MFEIT | REVRIV | RFDSQTK | LSDAIDLI | QPSTVVDNT | IDKDGDTLEW |
| EF | WWR | WYLS | MPPTK | VSPPRE | YFYDSVQ | TLGGAGGG | YDNEPSTAT | SYIDKDGDTL |
| FE | EWY | YWPA | NLIEL | YYHNTR | DKPIPLS | TLSDAIDL | LEAPADTPV | ALEAPADTPV |
| KY | ICN | FCLH | RRIRM | DKPIPL | GDKPIPL | GGRVKPLP | DTEGLLKEI | ELRAVLTEAL |
| QI | WWE | IFGV | VYVKG | GDKPIP | NQMFKKW | KNYGHPRE | LSDAIDLIN | KGNNSGQPST |
| YS | WIN | KWWQ | YLPTK | LINLVM | RSNMIRH | LGGAGGGG | MTLSDAIDL | PASMEGNRPT |
| PI | KMW | VVFM | AWYWT | QAYINA | HCTQVPI | LRAVLTEA | TLGGAGGGG | QPCCCPHCPR |
| KF | WTM | CEMQ | DVKTF | QMFKKW | KPIPLSG | RFDSQTKE | TLSDAIDLI | YDNEPSTATV |
| MI | GHW | EPFW | GHWLG | KPIPLS | PRSNMIR | DKPIPLSG | LGGAGGGGG | LEAPADTPVS |
| SY | YWG | FIHR | KLPIV | PRSNMI | RAIRRRR | GDKPIPLS | NKNYGHPRE | VKSYIDKDGD |
| EY | HWL | GWMK | KLYAN | RAIRRR | WRDPSTP | PRSNMIRH | RFDSQTKER | DTEGLLKEIE |
| YK | SWY | IWNG | LDYIG | RSNMIR | WSRPWGL | TEGLLKEI | VTLGGAGGG | FDSQTKERLT |
| QV | CHN | LMVI | MFKVY | DPLVNE | KNPEKGK | YDAISIRK | DKPIPLSGI | MTLSDAIDLI |
| YE | MCY | MCGI | NVVVD | HCTQVP | RHLIDTS | YNQMFKKW | ELRAVLTEA | NDTEGLLKEI |
| VQ | DFC | THAW | SHGIA | PLYSGS | RRHLIDT | HCTQVPIK | GDKPIPLSG | TLGGAGGGGG |
| IQ | CFR | TKCF | SKLYA | RHLIDT | RRRRVDL | KPIPLSGI | TEGLLKEIE | TLSDAIDLIN |
| ML | LHW | VFIG | SRDPY | WVSGWS | RWVSGWS | WRDPSTPT | HCTQVPIKV | LGGAGGGGGS |
| PF | AWY | VQYI | WKDGE | APGEGK | WVSGWSE | EAPADTPV | KPIPLSGIK | RFDSQTKERL |
| FR | IWK | VTWL | WLEET | ARKEYL | DPLVNEF | GRRHLIDT | TYNQMFKKW | RNKNYGHPRE |
| FP | MCW | YYHK | ADLKW | ARRFYD | GRRHLID | RNSTLSAL | EAPADTPVS | VTLGGAGGGG |
| PY | WGY | LFYF | ESQNY | IEIKPK | PLVNEFP | RRHLIDTS | GRRHLIDTS | DKPIPLSGIK |
| QF | MVW | NCYD | EYFYE | PKEKYY | RLINLVM | RWVSGWSE | RNSTLSALM | GDKPIPLSGI |
| MF | WWQ | PKHW | GGHWL | PLVNEF | RPRSNMI | WSRPWGLE | RPRSNMIRH | TEGLLKEIED |
| YP | MFW | WTQM | HWLGI | RETRNS | TANDDVE | DPLVNEFP | MWDPLVNEF | KPIPLSGIKG |
| YQ | MMW | GLFM | IGPAW | RRHLID | WDPLVNE | DRLINLVM | QPCCCPHCP | QPSTVVDNTL |
| QY | CYP | HHCG | IPSCA | SRPWGL | EKYYLYR | MWDPLVNE | RGSWQKKKL | EAPADTPVSE |
| MV | DWW | IWNA | IQNNK | YEEEQE | ITGEKYP | RPRSNMIR | RWVSGWSEA | HCTQVPIKVQ |
| FQ | FFC | MYYY | KWLAA | EKYYLY | PKEKYYL | VWRDPSTP | TVWRDPSTP | RNSTLSALMP |
| KH | MWW | TWGW | KYLPT | GEKYPE | TGEKYPE | WDPLVNEF | VWRDPSTPT | ETVWRDPSTP |
| DM | WYK | WIKT | LWDTV | ITGEKY | DERLNKM | WVSGWSEA | WDPLVNEFP | MWDPLVNEFP |
| HE | YWP | WVTL | MSGEW | KEKYYL | KEFAPDA | ITGEKYPE | IRNSTLSAL | RGSWQKKKLR |
| WE | IWQ | WWKN | NGGHW | KKVEYK | LEGPLYS | IYFYDSVQ | MMTANDDVE | STPASKVRRR |
| WS | WWG | YCNS | PDINE | KYNAKK | LIRLGIR | MTANDDVE | VKSYIDKDG | TVWRDPSTPT |
| MY | HLC | YWGM | PHPIV | KYYLYR | NAKKVEY | DERLNKML | DERLNKMLK | VRGSWQKKKL |
| HK | NFW | AGWW | PIWER | RFPSVE | PRHMEVF | GLIRLGIR | KSVGITGQL | DVKSYIDKDG |
| SW | WWK | AVWA | PLHWP | TGEKYP | SRPWGLE | NPASAEAI | RIYFYDSVQ | GIRNSTLSAL |
| WK | MHW | AYLI | QLWDT | VMVKEE | VSGWSEA | SPRHMEVF | SVGITGQLT | IMMTANDDVE |
| WR | MCC | CHCS | QQAYI | VSGWSE | YNAKKVE | SVGITGQL | GLEYEEQKQ | IRNSTLSALM |


| CP | MYW | CYGC | RIRMP | YNAKKV | AWYWTVA | AWYWTVAR | HGLEYEEQK | DERLNKMLKG |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| PW | CFQ | CYKE | RLAFV | AKKVEY | GLEYEEQ | GLEYEEQK | KQHGLEYEE | IRIYFYDSVQ |
| WP | LCM | DYHH | RLITM | ASYAFG | GRLEAAT | HGLEYEEQ | LEYEEQKQL | KSVGITGQLT |
| MW | WMM | FIWE | RRFYD | ATIFDI | GVRQNTS | KLAAEKAA | QHGLEYEEQ | SVGITGQLTG |
| WQ | KWC | GCYE | RRHVT | EFAPDA | HGLEYEE | KQHGLEYE | ADLKWAGIG | VKSVGITGQL |
| MC | PWW | HEFF | SIEYG | IRLGIR | KLAAEKA | LEYEEQKQ | AWYWTVARP | GLEYEEQKQL |
|  | WCS | IWDP | SPLHW | NAKKVE | LEYEEQK | QHGLEYEE | DPHPIVRDL | HGLEYEEQKQ |
|  | YWH | LCVK | SQTLI | PIEIRP | QHGLEYE | RETRNSSF | GRIEAATSS | KQHGLEYEEQ |
|  | WVH | MCVL | TIFGI | RHMEVF | RETRNSS | TLEGPLYS | LADLKWAGI | LEYEEQKQLT |
|  | CCK | MEWV | VIGMQ | SEEIKV | TFDNSPG | ADLKWAGI | NQMLSSLLV | QHGLEYEEQK |
|  | DWC | MNFW | VYQLR | SGLRGY | AAKVIAD | DLKWAGIG | NSTLSALMP | DPHPIVRDLY |
|  | FWQ | PPYW | WDTVM | YIPEQT | ADLKWAG | DPHPIVRD | PHPIVRDLY | LADLKWAGIG |
|  | HNW | RCHC | WLAAE | YLPTKR | DLKWAGI | GRIEAATS | PLGLTDPHP | NSTLSALMPC |
|  | SWC | VYFK | WMPPT | AWYWTV | DRLINLV | LADLKWAG | PPLGLTDPH | PPLGLTDPHP |
|  | CWE | WCIR | YFYEE | DKNEVI | EEALAWA | LARKLAAE | RAFESGDFA | RAFESGDFAR |
|  | FWP | WITL | ADDQW | EEYAAA | GGSFELA | LGLTDPHP | STLSALMPC | STLSALMPCE |
|  | MWC | WVTS | DQWVP | GNLLDV | GLTDPHP | NQMLSSLL | WQKKKLREV | SWQKKKLREV |
|  | WGC | YCSQ | DSHWT | GVRQNT | GRIEAAT | NSTLSALM | AATASPASM | AWYWTVARPD |
|  | WWS | CLMK | GASCA | ISIRKP | LADLKWA | PHPIVRDL | AGEDGLTYR | DDSHWTFSSD |
|  | CHC | LCFQ | HRYQI | KNAENF | LLYAVSN | PLGLTDPH | ARWVSGWSE | DENGNIHVSK |
|  | WCE | LWNQ | LSCEY | LEYEEQ | LRRLADE | PRAWSRPW | DENGNIHVS | DSHWTFSSDL |
|  | CWI | LWYI | MKAIC | LTSLGG | NQMLSSL | PSTVVDNT | GALPGEVVG | KVVSHLPGVV |
|  | HCM | MCYG | PEFIT | MQQQAY | NSNLGQL | QKKKLREV | GFSGCEHRS | LADENGNIHV |
|  | WCK | NIIY | QWVPD | PEEALA | PHPIVRD | RAFESGDF | GLNKVVSHL | LPGVVHEMRS |
|  | CWD | QVYI | RKAFL | VDRVER | PRAWSRP | RIEAATSS | PGVVHEMRS | LSRVTDATTS |
|  | WKW | SWIL | SCEYS | WKDGEL | QQQAYIN | RSRSYIKL | SHLPGVVHE | RLSRVTDATT |
|  | CWM | WDRN | SKVIL | WRDPST | RAWSRPW | STLSALMP | SRVTDATTS | RSGLNKVVSH |
|  | CWH | WLIG | YQIKD | YKLDLD | RIEAATS | TRAFESGD | VVHEMRSEA | YYGRSGLNKV |

## Table A10. Characteristic n-grams in ordered regions by zscore values

N -grams presented in the table have abs(z-score) $>2.58$ in ordered and abs(z-score) $<1.65$ in ordered regions. Table includes for each length first 100 n -grams sorted according z-score values in descending order.

| N-gram length |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| INN | GCPW | CRELH | NALLLY | KGSGKSM | INVIDDVE | SHASNPVYQ | KVTGGQYASN |
| NII | YMAC | HASNP | HLDLSP | DGDTDSY | HNVIDDVR | THASNPVYA | TVTGGQYASK |
| WAR | CEGP | HNYLC | RTGKTM | FSGKSTE | VNVIDDVI | WYNVIDDVD | LHLHVLIQFK |
| DDV | CPWD | HRGTH | ENALLL | EASNPVG | VGPCKVQD | GPHNYLCGH | YWLVRDRRPN |
| DTI | EGPC | TIFLC | CVAEAW | QASNPVH | GGKTMWAV | SLGPHNYLS | TYDLIRDLIA |
| AMA | WMDE | FLTYP | CNIDLH | FASNPVW | EYATLKID | ALGPHNYLS | LWARSLGPHI |
| NIG | WQSN | HGFTH | YGKPVQ | DGTGKSG | LRTGKTMN | SLGPHNYLC | STHFAKFKGR |
| FIN | MACT | MKIDH | ATLKIR | HASNPVY | SKYENHTF | ALGPHNYLC | MSTAKHSVDV |
| IFN | CKVQ | HTENA | NLDRIF | RLCNPGW | ATGGQYAQ | EYNVIDDVA | FDRINVRRLF |
| NYI | GNHD | YLCGH | YATLKI | CLCNPGF | DNALLLYN | VATNIIENG | MRADVKEFEQ |
| VYI | PHNY | HRVGK | HDDLVM | QAAAVAI | RNALLLYF | GAQRDWQSN | FRADVKEFEA |
| VNG | KTMW | MACTH | ALLLYM | GHASNPQ | YALLLYMF | PVQIKGGIP | NQVPINATGH |
| ITV | CTHA | DVDPH | MDENIK | TVRDRRF | YPHLHVLF | TSLYPSIIR | ISDVTRGNGI |
| FTL | GPCK | KYENH | CAIIAW | RVGKRFC | EKYENHTV | ASLYPSIIR | GSSYKEFLDK |
| DPR | ACTH | HGFRC | RLEAIC | DGPCKVP | NLKHFKEN | YLRVLAALK | ISDVTRGNGL |
| KKF | DECH | RDRRP | SGHLDF | DWARSLR | AKYGKPVE | YFWRPEEVS | VLPTSAGKSA |
| ITI | IWMD | QIKGG | CISDVT | GCKVQSK | GIPTIFLC | LENALMLYS | NLPTSAGKSL |
| GEV | DRRP | NVIDD | HRVGKR | TCKVQSN | KVQIKGGF | SSLYPSIII | LGGDFLTSLV |
| FGP | NHQE | PVYAT | CVKSVY | RDVTRGR | CKPVQIKR | EGTGKTTLS | RCVSDVTRGS |
| QQL | TMWA | KGGIP | WKELIG | HVGKRFG | GGKPVQIA | FVGSGKSTY | CGYSQGAIVC |
| GDV | PHLH | CTHAS | HGSTIM | QGTGKTW | NVIDDVDP | FGLMVWCII | FLVRDRRPVD |
| ISS | YNHQ | QFEGK | HGSTVM | HLAAAGW | QGTGKTTY | IGGDFLTSF | DYSPDTLGYE |
| LDD | GHLD | FMGAQ | NVIDDV | RIDDVDF | MAGTGKTV | FGLPATADL | NEQALVKRFW |
| SLI | YYCW | FKEFM | GNGITH | VYNVIDW | NKNDLRDG | WLVRDRRPY | GDPFWYEDDV |
| DDL | TPLH | FTHRG | IKGGIP | PVYATLK | EGPCKVQS | ERIDANLLN | YDFASLYPSN |
| IYN | CKIC | HVLIQ | QIKGGI | YthRVGE | DFFDLVSA | KRIDANLLD | VSDVTRGNGI |
| DNI | CRTC | QSNTK | QLRRAW | QIYFYDG | ENALLLYM | ASLYPSIIQ | LWARSLGPHN |
| FAP | CIPC | GKRFC | KNDLRD | YMDENIR | FAAAAAAH | MSNLCTEIS | YRFFDLVSPS |
| VDG | MWAR | KIRIY | WTKTVW | TENALLL | RPSTATVG | NNKKFIKIL | IPRRHGKTWI |
| VVG | HTEN | NHTEN | QIGRVP | TPVYATN | TENALLLY | TAGFGAGFT | VPRRHGKTWF |
| IHS | CDKC | GKTMW | ELIGAQ | NPVYATL | NALLLYMA | CPGSGKSTW | VPRRHGKTWI |
| WRL | THAS | QVFNM | FDLVSP | YATLKIR | GNLRKALY | IYDKYNDVY | VSDVTRGNGL |
| NIL | LYMA | YATLK | MLAIKY | IVTGGQT | LRGCEGPA | QYDKYNDVN | FGPAGTGKTS |
| RRL | CAVC | MDENI | YGALGN | PVTGGQQ | WEPSTATK | GPTSAGKST | KQAIELLPDF |


| VFI | YCWM | LGKIW | WALKNA | QVYATLE | YNVIRAVY | TIHSRSYTH | VTDIAGYAGV |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| CPW | DWQS | SLGPH | CGVAAC | FLKIRIR | GKPVQIKG | VLTEGDSAS | RVGIAVDTGT |
| KMI | MVWC | THASN | HITNAH | SKRFCVV | NKYGKPVR | ANPFLRPEF | LGKTTVVAIF |
| NTG | HASN | SNPVY | CGHLDL | LRTGKTG | AGSGKSTC | ANPFLRPEL | LAGLPATADK |
| HNL | NHTN | NHNLR | SDVTRG | DRIYFYQ | IHTNSVMR | FDAIVQALK | KSCSQGGIRG |
| LNF | DEAH | CNPGP | WKALSH | VKYGKPT | FGTGKTTF | SLSICNAHV | VEGPIDSLFL |
| IKF | GKIW | YNVID | VGKRFC | PFCVKSK | YGKPVQIK | IGFKTRYGM | FLPEKTLGWQ |
| RVW | MDEN | QRDWQ | HAIAQC | HGKTTLF | QLRKALGH | IFLTYPQCD | DKQGARWTGR |
| IMV | RDWQ | IRIYF | QEAGKY | MGKTTLY | AQEAGKYL | GIELLPDFY | KVSDAAPYIF |
| WGP | CAYC | GDTDS | FPETVH | EINNVIH | KIRIYFYD | WDTETTGLP | QVSDAAPYII |
| KPA | KRFC | KYGKP | IKLKNH | GENALLG | PSTATVKN | KGARWAGEA | DYETAVREFI |
| WNG | RFCV | SDVTR | HFKEFM | FYATLKK | VCISDVTF | GEMTVAGKK | ASLYPSIIRA |
| AVP | CGCS | IFLCN | GIPTIF | FKHFKEE | IVRDRRPT | RGARWAGES | NEMDAGIYYA |
| KCA | PMYR | CSLTK | WARVAT | CRFCVKR | GPSTATVD | YFLTYPQCS | DEIIDNSVDE |
| NGV | DPPY | RIYFY | YNVIDD | PLGVISW | VGYSQGAE | FNKITKGGL | LPTLYFSADM |
| VFD | RPMY | SNDAW | CSLTKE | PYNHQEP | LTRGNGIL | MIFLAMLVI | RFLRGQLALV |
| IHD | HLHV | RVGKR | CVSDVT | QAAAIGY | TGKTMWAR | GTLFLTEGK | KSPKWLNDLI |
| WRP | YRKP | HDISH | NIKTKN | HRVGKRF | LKLKRLRG | MIHSRSYTY | SAGLPATADF |
| WLR | RIYF | ENHTE | HIGDLM | AENHTET | ERGNGITL | MIHSRSYTH | FDLNSLYPHL |
| WPI | PAGT | LLLYM | ASNPVY | YGTGKTN | NIIENGVT | IAEVERLRS | NKPGDDFQLG |
| ITS | LCNP | LYMAC | WARALG | RTGKTMW | LGQVFNMV | GFGAGFGAG | MIDLPPLGGT |
| CTH | CAIC | HDKRM | HTNSVM | GGIPTIF | IEALSQLG | LRVLAALSR | WLVRDRRPVT |
| LNS | YFYD | WYNVI | QINVAH | RLLLYMH | TLKNHTNL | RRVLAALSR | TVSGAVPGQM |
| KRY | CVVC | TLKIR | ELLPDF | CDSRTGF | ASINNVIY | KKALGIHKA | SVSGAVPGQI |
| RGW | LLYM | RGTHH | YLKLKH | YNLDRIV | SRVLAALD | KSIELAQDS | GGGDIYHNTT |
| LYN | NCKY | PVQIK | HVTGGH | DGTGKTD | VIENGVTD | RKALGIHKC | EKQGARWTGM |
| PGW | WTFP | EGDSR | YVSFAC | DNYLCGT | RIRFYVST | YSIELAQDL | PHLHVLIQFE |
| AIN | HRVG | NDAWY | WDIEIC | GKRFCVK | ASTATVKS | SPTGSGKSL | ACNLGHINLS |
| FSN | NCRC | PHYLK | WDLGGM | IRDRRPN | IKLKNHTN | VIGLHHVTG | EIARMYGVTR |
| LLS | CNLC | KNHTN | SRTGKT | DKYGKPF | YKLKRLRS | IIGLHHVTA | ATGNAAIEEA |
| LVG | DTDS | THRVG | CALINM | FGTGKSE | NENGVTLI | EVNRFIIYA | SFDKQGARWT |
| HGC | GFTH | FLCNP | RVGKRF | ACTHASN | FAEVERLA | TENALLLYM | DIARMYGVTP |
| YAS | KNHT | TGGQY | CVIGLH | WDDVDPE | GKVMCISD | SGMYASALN | PCNLGHINLA |
| RAL | WPWP | RSLGP | NIVAAH | VLCNPGE | GQYASKEQ | STPNGLNHY | THVVYNHQEQ |
| FTN | PVQI | MFFLV | WLVGEH | WPVQIKS | FNGAGKSF | GDFLTSLIN | LADRIADRIA |
| IVW | NHTE | CAYFW | CSLAAD | WMDENIK | RSLGPHNY | ITLFKEIRR | WYDPLAQSFI |
| GTD | WGHP | YLKHF | PIDSLF | CAARAAH | HGLPATAE | PTIGIGHLI | VFCIMLGTGM |
| ISI | SNPV | VVYNH | TLKIRI | FAGKSTE | QSYEQRHD | WNISPETII | IFCIMLGTGT |
| IYL | WYMW | GFRCM | MIIATY | GCVKSVS | FGSLKAEG | AVAIFLAHY | SSHQYGGTTL |
| GFN | FRCM | PLYFK | KAELRP | ECVKSVE | QRLGRVGR | VVAIFLAHF | EPIAYNATPN |
| RRS | CAFC | EQRHD | EAGKYE | PTIFLCN | HAAAAVAM | DAELNAILA | NFLRGQLALI |
| YGP | NYLC | HFKEF | WNGSLK | RALAAGM | ALRDRYQN | AFKTRYGIC | TEATDTSFVL |
| LVA | HKCF | AQRDW | FFLAAW | SEFMGAM | AGAGKSTS | WDPLAQSFL | LKPGDDFQLA |
| DRF | VFNM | SPKVY | FNIASY | AVYATLP | GETVHGFS | RDLECGCSA | REKIHGTNFS |
| NVG | WYVD | IYFYD | ENIKTK | VIDDVDP | FAKFKGKL | RALDNLLDY | QRLRDHGEYM |
| YSN | PLPW | CGHLD | MLAVKY | VGDSRTF | SNTKYGKP | KAAELRNFA | LLAHVGYPRL |
| LHV | FLTY | IKGGI | SNPVYA | FLCNPGP | KATNIIEW | SKEQALVKR | WVVEFDPNIP |
| EYV | FWKH | CMKID | PIAGLE | YQRDWQL | VATNIIEN | LEINREVVD | PAGTGKTTLT |
| NNL | GDIV | PTIFL | HLEAIF | WYLKHFY | KSVYVLGK | LSGIKGQIG | EDLNSLYPHI |
| KIA | ENHT | YFLTY | MVTAPC | DPHYLKH | IQFDSSLY | AGTGKTTLT | ARIFGGAWEQ |
| IIK | GTHH | NPVYA | YPAGTW | NFMGAQI | CVSDVTRG | RIHSPSRVA | GRIFGGAWER |
| GYN | CWEC | PHNYL | WLAGGW | FEQALVA | VDLIRDLQ | CNPFLRPEL | IADRIADRIT |
| SAI | YPQC | GKIWM | SDRRPQ | ARSLGPH | IKLKRLRF | IFLTYPQCS | WVGIAVDTGN |
| GND | LWFM | YNHQE | WDLTNC | GRELHEF | DVDPHYLK | ILTEGDSAA | LHGEDPHPFE |
| YNN | DPHY | WARSL | CGTALC | HNYLCGH | FSLKDPIP | HTKQAIELS | AFIQDIYDKI |
| IIS | EPWH | DRKPH | HVLIQF | NKRFCVG | KVCVDDFN | DDIDDIDDI | GGIRGGSATC |
| QTI | CECG | LCNPG | WDLDKD | CGHLDLS | NIDLHYFS | RTKQAIELL | HMQATLPGGT |
| NYG | FNMF | NMFDN | HASNPV | GHRVGKS | SLKDPIPW | PGAGKSTMM | FHGEDPHPFA |
| ESF | GARW | FCVKS | IRIYFY | TLVRDRH | PIPWKLYY | AQFDSSLTG | NGIRGGSATV |
| GHL | HQEA | DVTRG | YGDTDS | HFKEFMG | IVRGLLCT | SWWRNYAHA | SVNRFIIYSE |
| YIV | VWCI | VGKRF | HTENAL | QWMDENS | TPTRQFSS | VLFGKPFRS | VGSGKSTGLP |
| IKQ | AWYN | YMACT | LVRDRR | TLKIRIY | LVRDRRPT | FWTAKKRYA | KPKSIGVATT |
| IFV | CQIC | VTGGQ | PVYATL | KGGIPTI | LFRAPTVD | SQFDSSLTP | EVVFKHDYEE |
| IYK | PLYF | DWQSN | CTHASN | YTGGQYV | EKTLTGKV | IKGGIPTIF | ANTDCDGDKK |
| DYV | PTIF | GPHNY | THRVGK | KHGFTHA | VYATLKIR | VQIKGGIPT | AICNAHIPGN |
| YIK | IYFY | LGPHN | NPVYAT | NYFLTYP | AAKFKGKK | QVFNMFDNE | HPWMSPAGYR |

## Table A11. Characteristic n-grams in disordered regions by z-score values

N -grams presented in the table have abs(z-score) $>2.58$ in disordered and abs(z-score)<1.65 in disordered regions. Table includes for each length first 100 n-grams sorted according $z$-score values in descending order.

| N-gram length |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| DDD | SSSS | WALKC | KSSSDV | HAPAPAH | YSSSSSSI | WGGGGGGGF | PAGGGGGGGR |
| QQQ | WSFL | CRKRW | WKKKGW | EPPPSPF | IPKPAPKA | AAPKPAPKK | FSSSSSSSSY |
| DED | GIQG | sssss | ISASAY | PGDKGDM | WRRRRRRW | GPEPEPEPH | RAAAAAAAAI |
| HHH | PSPP | PKPAP | AKSSSD | FGGGAGH | FAGGGGGE | QGGGGGGSC | FEEEEEEEES |
| MSP | PQGP | CDGSC | SPPPPS | WDEDEDW | CPAPAPAC | VKPAPKPAV | FGGGGGGGAS |
| PPE | NnNN | IQGAK | YPLSPY | MGGGGSQ | GKPAPKPI | NSSSSSSSSN | SGGGGGGGAL |
| PDP | MWDP | WEREW | MPAPPC | IAKSSSI | SQGPKGDV | DAPAPAPAD | PTTAATTTAV |
| EAE | EPEP | EeEEE | HGGRGM | KSARGGF | TGGGGGRI | NGGGGGGAD | VPPPPPSPPPL |
| MNI | QGPQ | HLVEF | CASGAC | LGPEGPF | LGGGGGGK | VPSPPPPSPH | YGGGGGGGGA |
| SPS | MDSR | FTKRH | CESSSQ | FGGSSSP | RPEPEPEQ | AEPEPEPET | ASSSSSSSDE |
| KEK | DWSF | SSDVK | YAELKF | HGGGSGL | QDEDEDDY | KGGGGGGGW | LPPSPPPSPL |
| WYC | GEQG | RPADI | WSGGSI | VAPKPAR | NPSPPPPY | VAPAPAPAK | SDEDEDEDER |
| SES | HHHH | GPTGP | IGGGAE | KGAGAGM | KPEPEPES | KPKPAPKPK | RAGGGGGSGV |
| EME | DKGD | KGPPY | CRERAN | MSSDVKV | PPEPEPEK | DRSRSRSRD | VGGGGGGGAG |
| GGP | APKP | WSPPF | MKGDKP | MPAPAAD | KGGGGGYR | SPSPPPPSP | EPAAPAAPAP |
| PNP | WLNC | CGPEF | PRELNF | TQGPQGF | HGGGGGYQ | AGGGGGSGR | STNGIEPPRG |
| MGG | ISMC | QSAND | SSSSSS | HPAAAPD | LGGGSGGP | PGGGGGGGK | AGAGGGGGGR |
| DSE | PTGP | LVTTF | ISTPAS | IEDEEDV | SAAPAPAQ | DSRSRSRST | TEEEEEEEEI |
| ESE | DDED | GAKSS | HPKGDH | QGSGGGK | PSDSDSDF | GSRSRSRSQ | PSPPPPSPPT |
| MAN | GDQG | WAVQW | QGAKSS | HGGGGGY | QSGGSGGP | NSGGGGGGR | HGPAGPQGPR |
| MKP | SDSD | SARGG | YSLEEF | VSGSGSQ | ANNNNNNL | TDSDSDSDT | QSSSSSSSCT |
| QKQ | PKPA | GGGGG | CEDDDK | HGGNGGK | VSDDDDDF | PPSPPPPSP | AGGGGGSGGV |
| PHP | GERG | CGDDC | CKRLRC | TAKSSSK | GQLKGSSQ | VDDEDDEDQ | GSDSDSDSDG |
| MGL | AHFH | TTDPW | YAARAC | NAGGGAQ | SQLKGSSS | KPSPPPSPK | VGPKGDTGAD |
| MKG | WCCW | HEQDW | NPASAE | KEDDEDH | TTGGGGGP | EAATTTAAL | MAAAAAAAAE |
| PQL | LMPC | FASFH | CKEKVH | KPAPKPA | DIQGAKSA | AQQQQQQQM | vssssssssw |
| MDG | GLQG | SDVKS | CAPLPM | WSSSGGW | GPAAPAAS | EPTPPPTPE | AEEEEEEEER |
| MNP | WYPQ | TTTTT | GEEQKF | HEPEPET | HAAPAAAC | VGGGGGGAQ | SPPPSPPPPPT |
| NPA | PALP | AKSSS | FDSDDM | IGPEGPL | NPAAPAAG | PPAPAPAPP | QQSANDAYAE |
| DDP | GSTG | PSPPP | HAGTPN | GAKSSSD | RSSSSSCN | SSRSRSRSA | RPMNRKPRMY |
| GQG | GKDG | PPPSP | mbeeef | YGGAGGD | VPSPPPPY | AGGGGGGRV | NAAPAPAAPE |
| RMR | EDDE | PPSPP | WPRPAM | KDDDGDT | QSSSSDSM | GGAGGGGGS | ATNGIEPPRG |
| PPK | VKSY | CTGKW | DPKGDF | GAPKPAT | PGIEPPRE | KTtTTTTTK | AGGSGGSGGA |
| EYE | IWDQ | FMKKW | MEEKKF | GEEAEEM | DSPPPSPR | MGGGAGGAV | IGGGGGGSGH |
| SKS | PSSP | YSGKW | NDAAAE | RGGAGAN | FGGGGGYG | AGGRGGGGK | APAAPAAPAA |
| LAP | DSDD | VMGGH | HGGGDN | IRGGQQP | YGGGGGYA | GPKPAPKPS | RYGGGGGGGG |
| MPP | GHMA | CAPGF | FEAAAD | TEEDDDM | VPPTPPPV | TPSPPPSPI | YGGGGGGSRF |
| DTD | QHIS | YASDC | CSSTSC | NHPNIQP | QGGGGGGQ | YSDDDDDDS | KPGGGGNGGH |
| ENE | WAPW | QTAND | MGTGGQ | PPKPAPQ | ERGGGGGY | RYGGGGGGG | TSSSSSSSDG |
| FGF | YWFW | MVASM | YEEVEH | QAEAKAH | RGSGGGGN | ASSSSSSCV | CQSANDAYAE |
| TTG | DNDD | YQRVC | EPKGDE | EGGTGGP | LDDDEDDN | LSSSSGSSL | STTTPTTTTA |
| MGP | MKTY | APAPA | MRSSSP | GPQGPQG | TGGGGGAV | KAAPAAPAK | YAGGGGGGGL |
| MKS | PLFQ | FTALM | VELADH | PSRSRSH | TDDEDDDL | PAPAAPAPK | RGGGGGAGGA |
| PQI | PPGP | TERHT | RGDKGF | REEDDDN | PDEDEDEY | GGAGAGGGA | SEGDRRRVRI |
| MAD | FYHY | GPVGP | MPPLPK | TRRARRN | EPPPPTPD | QSDSDSDSE | TGGKGGNGGS |
| NQN | GTGG | ANDAY | GSGLSM | DGPEGPD | KPPPPTPN | SPSPPPPSL | VDGKDGKDGV |
| ESG | MGNL | WSFLK | FAATPC | QSSGGGY | KGGGGGPF | PPPPSPPPP | GDRRRVRIEV |
| MEA | NYGH | IVIST | MDSRTG | HTTAATL | TGGGGGPM | PDDDDDEDP | QGPQGPKGDG |
| GSL | GAKS | YIVKY | YPLPAM | FAGAGAN | RAAAPAAN | AGGSGGGGK | VSEGDRRRVR |
| GMI | DEED | YGLGW | YAREQT | EAAAAGR | FGGGGSSC | SGGGGGGAV | GAPAPAPAPS |
| TGP | GAPG | PAAAP | FIEKLI | MRRRRGE | DAAPAPAS | APAPAPAPA | KRRQKREDER |
| PPF | QGPK | YRQEW | STPASK | QGNGGGC | SGGAGAGL | PGRGGGGGC | RELLDLARQQ |
| EMQ | KGDT | WARAF | HPPPER | GSRTGEG | ADDGDDDW | PDSDSDSDV | TLTQQEQQAQ |
| PVE | PTPP | YGADY | WASTGH | SSRTGEK | AAAAEALR | ESSSSSSSA | EGPQGPQGPE |
| MNS | GSSG | NGIEP | SKRPAD | VPAPKPR | AAGAGGGM | KYGGGGGGS | ASSSSSSSSD |
| MAG | PVGP | GPPGP | NSPTPY | DAAAEAL | TAGGGGGR | ISGGGGGGP | PGEDVNSLVI |
| VMP | HLMC | QLKGS | EGDTGW | NSSSSTP | LAGGAGGK | RPAPAAPAR | SGGKGGNGGA |
| MDF | VMKW | FVENM | FLDELW | SPLPPPW | KPPAPPAE | FGGGGGGGA | AGATGPQGPM |
| MTN | YLVT | YLPFW | FSKSPW | DAKSSSA | TTPPTPPE | NESWASRSE | GDKGDKGDKG |
| GGR | SIRT | CDSSC | GERLEV | YDGDGDL | PGGSSGGA | VPGGGGNGA | NLAAARASTQ |
| TMA | DGDD | CKITC | EPEPEP | EPPAPPE | YFGGGGGW | SAGGGAGAD | QEAPEWAPPK |
| MEP | SSTW | SQLKG | YGGAGT | EPKPAPM | QGAKSSSD | GGAGAGGGG | KGGNGGSSPS |
| SLS | GEEG | WVRPI | FSSGSV | RATTTAR | DAAKKAAN | SAPAAPAAK | VRDALAGKRA |
| RYH | KKKK | YTAQF | MSGTTL | NSPSPSN | IGGGGSGE | VGGGGGGGA | NGGNGGSSPT |
| WFC | DGGD | QGPKG | NTASDF | QSSSDSY | TAEKAAEN | SLGSGLSMS | QGLGTEAPSN |


| TES | PNSP | CNTAH | YADADM | WEPEPER | QSANDAYA | AGPQGPKGE | DPAPAPAPPK |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| MPK | QHFA | FVSDC | FAPRTW | KDGRSAI | KAAPAPAK | KGKDGKDGC | EGPGGPPGPE |
| NGR | QPQQ | YGVLC | YLPSLW | PAGGGGK | PAATTTAI | KPSPSPSPK | HKSGKNKGQP |
| YPY | PSEP | WDDIF | YPRRRY | LGPTGPE | DAAAEALN | DNPASAEAI | LMPCESSSQV |
| DMA | PEVP | HGPRN | CNRRRI | KRRAARI | DGGAGGGD | GEGEGPGGE | GEEEEEEEEG |
| TPL | PGAP | CQQQC | YPARPF | NPPTPPM | QSSSSGSP | APAPAPAPP | FMPCESSSQI |
| DPQ | DSDS | MVKPW | QKLKKY | GDEEDEL | STGGGGGR | APKPAPKPA | AATRAVTAAG |
| PIG | QDVQ | HGFQM | FTPSSH | FAERAAS | VAAAAEAN | VGPVGPQGS | PGGGSGGGGA |
| EHE | YHAY | WNESH | CTAPAY | PPPSPPP | FAPAAPAS | ERTATETRR | AKGDTGAQGE |
| APY | IVIS | SSSDV | GPPQPM | MPPPPTA | QPAPPPPV | PAPAAPAPA | AGTPLRRYPL |
| HYP | PAIP | NNNNN | WTSKPH | PKPAPKP | HRSPSPRK | PPPPSPPPS | GGVSGNPRAN |
| QLR | QPQP | FNVPQ | GPNIQC | KAKSSST | KDEEEEEK | AAPAPAPAA | MSGLLDDGAN |
| IKG | MPCE | HALDH | LPNIQN | YAAATAE | SSGGGGGL | AGGGGGGGN | RGVSGNPRAD |
| GTS | DDSD | MSKRP | KGGGGK | PEPEPEP | TGGSGGSA | TGGGGGGVD | SGTPLRRYPM |
| SHS | DDGD | NTERH | QESDDH | GPQGPKG | NSRSRSRG | KGPQGPQGQ | TAAAPAPSKG |
| RHH | DDDP | FGFGV | NERLEI | PAPAPAP | SNPAPTSE | EEGEGPGGP | FKGAKGDKGE |
| GEL | NECY | FPDFH | LPGPGC | PAGGAGN | APPLPPPA | DGAGAGGAD | PAAAPAPSKP |
| TPP | SRYC | QGIQG | CTTTAL | NDEDEEY | YSGGGGGK | NGRGGGGGV | EPAPKPKPAA |
| RPQ | PAVP | HDTNM | SPDPDT | YLSSSSE | TRSSSPSV | LRGGGGGGK | PGPEGPQGPA |
| GYE | PSKP | VRGSW | YESLPC | GPSPSPG | RGGGGGYV | ATDLRGSGG | APGGGGNGGD |
| KTT | PVEP | YSDQM | YLERQH | NPPQPPG | RAPAPAAR | YGGGGGGGG | GGGGGGGGAG |
| IAG | TTST | PAAQW | CPSGSH | SAAAKAV | FRSSSSSP | KGAKGDKGE | AGGGGGGGRR |
| MGD | EEDE | MGSLI | FVESEW | AEDEDEV | MSSASSAT | KGAKGDKGD | DSSSSSSSGE |
| IRA | RLIH | FDEPH | GAKSSS | LSSSSEN | VRSSSSSC | TYGGGGGGA | AGGGGGGGVR |
| DYE | PSLP | HMSHH | MAAGPW | LGGGSGP | ELNPAPTS | YYGGGGGGA | QENTERHTAG |
| MRA | QMIA | QINGW | HSPSPG | CGGGAGS | LRRRLERG | APKPAPKPK | EEEEEEEEEE |
| SLG | GEPG | FGPHM | FQAPAR | GPPPAPR | CESSSQVS | SGPAGPQGA | VKGDKGDPGN |
| TAI | APAP | HSTQV | PKEQEP | IKSSSDM | ALRRRLER | SDPREEQVS | LSDEQLEALL |
| PEE | PLQP | YDESY | ISPADY | MGRRRSH | DLSDEELR | KPAAAPAPS | GAGGGGGSGR |
| LNP | WDPL | FQMRF | FELQEP | IEDDDDM | PAAPAPAA | SDDEDEDEI | TGPKGDKGDN |
| DID | PTEP | RSPSP | NATAAM | VRRGRRE | IIISTPAS | PGGGGNGGH | GEGEEGGGEG |
| GDP | RGGQ | YLVTT | KPAPKP | TSTPASD | TTTAATTT | VGPTGPTGD | KDLTESQKEK |
| IQG | LFQD | YVRVH | PKPAPK | NAELEAR | ILEEAQRL | AGGGGASSG | NSKFSEKKKS |
| RTA | KVFI | TASDW | RSRSRS | MTGGSGP | ESSSQVSN | GDAAAAAAP | ESSYLDARHK |
| ATR | GVQG | PPPPP | PEPEPE | GAEAERY | SSQVSNST | DARAAAAAP | SKVGRFTVMT |

## Table A12. Characteristic n-grams in ordered regions produced by combination of z-score, fractional difference and mole fractions

N -grams presented in the table have mole fractions $>1 \mathrm{E}-6$, abs(z-score) $>2.58$ in ordered and abs(zscore) $<1.65$ in disordered regions. Table includes for each length first 100 n -grams sorted according fractional difference in ordered regions in descending order.

| N-gram length |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| SAL | DALA | AALAR | PCKVQS | EGPCKVQ | VPRGCEGP | DNEPSTATV | SFDQVPEELE |
| LDD | ARAL | ALAAA | ARSLGP | QSNTKYG | PRGCEGPC | EGPCKVQSY | VGSGKSTGLP |
| STL | LSLS | LAALA | SRTGKT | HASNPVY | CKVQSYEQ | RGCEGPCKV | RKPRIYRTLR |
| VDE | LADA | AALAG | YGDTDS | QIKGGIP | EGPCKVQS | LVAEVERLR | NEPSTATIKN |
| LSN | LDAA | GAVAA | VRDRRP | THASNPV | EGDSRTGK | SRRKFLNQV | NYIESHRDEY |
| VKA | LLEK | LAALS | IKGGIP | ACTHASN | QSYEQRHD | SSYKEFLDE | FDNEPSTATV |
| NLK | ELLD | AAGLA | NVIDDV | ASNPVYA | VQSYEQRH | THASNPVYA | EGDSRTGKTM |
| EID | RALA | AALGG | YNVIDD | MACTHAS | SNTKYGKP | VYATLKIRI | FKEFMGAQRD |
| TLK | VDAA | AGAAV | HASNPV | SNPVYAT | THASNPVY | TENALLLYM | HYLKHFKEFM |
| LNS | ELLA | LRKAL | ASNPVY | NPVYATL | CTHASNPV | EGDSRTGKT | VIDDVDPHYL |
| RAL | RRLL | AAALL | YMACTH | CTHASNP | NPVYATLK | GAQRDWQSN | MWARSLGPHN |
| DDV | LLKE | VVAAA | LYMACT | PVYATLK | VYATLKIR | IKGGIPTIF | PHLHVLIQFE |
| ALD | EALL | AVAAG | QIKGGI | YNVIDDV | KIRIYFYD | GPHNYLCGH | DFGQVFNMFD |
| TTV | VEAL | GTGKS | SNPVYA | NHTENAL | TGKTMWAR | PVQIKGGIP | KVTGGQYASN |
| TIE | ALLS | AAVGA | MACTHA | HRVGKRF | LKIRIYFY | SLGPHNYLC | MDFGQVFNMF |
| DIS | LGAA | ALALA | THASNP | THRVGKR | TLKIRIYF | ENHTENALL | QSNCKYGKPV |
| DIE | GAAV | LLAAL | ACTHAS | RVGKRFC | YATLKIRI | WYNVIDDVD | TVTGGQYASK |
| VAR | TLTA | ALAVA | NPVYAT | WMDENIK | SRTGKTMW | CGHLDLSPK | ERIQRLGRVG |
| GEV | ADAV | AALAV | CTHASN | LGPHNYL | ENALLLYM | MGAQRDWQS | VKSVYILGKI |
| RLG | RALG | ARSLG | PVYATL | YATLKIR | TENALLLY | HLHVLIQFE | NHVVYNHQEA |
| EIA | AALG | LAAGL | NHTENA | VYATLKI | HTENALLL | EGPCKVQSF | FDRINVRRLF |
| TIK | ALGG | RDRRP | HTENAL | LVRDRRP | VGKRFCVK | STATVKNDL | KLKNHTNSVM |
| LAA | VATA | AGTGK | THRVGK | GKTMWAR | NALLLYMA | MFFLVRDRR | LSTAKHSVDI |


| LNA | VGAA | AAVLA | HRVGKR | IRIYFYD | GIPTIFLC | VATNIIENG | TKYGKPIQIK |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| GLD | SLGL | GDTDS | SDVTRG | TGKTMWA | KGGIPTIF | QVFNMFDNE | NLNSNLDRIF |
| AAV | KELI | AVALA | GKTMWA | KIRIYFY | LKHFKEFM | VKSVYILGK | ISDVTRGNGI |
| VEG | AGAL | SDVTR | GPHNYL | RTGKTMW | DSRTGKTM | WLVRDRRPY | FRCMLAIKYL |
| LNN | KLIE | GALAL | RVGKRF | LKIRIYF | VTGGQYAS | KVTGGQYAS | QIKGGIPSIV |
| LTA | VSAL | VTGGQ | LGPHNY | TLKIRIY | CGHLDLSP | VQIKGGIPT | SETIHSRSYT |
| ITS | LEAI | RVGKR | VGKRFC | ATLKIRI | RSLGPHNY | IWMDENIKT | ALEAIRFYVS |
| TLD | GALA | AILAA | WMDENI | GKRFCVK | WYNVIDDV | GVISINNVI | IRDLISVIRA |
| NNL | VAAG | TENAL | MDENIK | KRFCVKS | NVIDDVDP | NTKYGKPVQ | NLPTSAGKSL |
| SAI | GLGA | LVRDR | YATLKI | SRTGKTM | GKPVQIKG | VTRGNGITH | RVNNYVVYNQ |
| AGL | DELV | QIKGG | ATLKIR | VTGGQYA | YGKPVQIK | TVTGGQYAS | LKRLRFKGTV |
| NIK | RLLA | TLKIR | LVRDRR | Yenhten | PVQIKGGI | HVVYNHQEA | FRCMLAVKYL |
| VTA | LDAV | TGGQY | KTMWAR | ENALLLY | YNVIDDVD | RDRYQVLRK | ERIVSILEWD |
| LGD | AVGA | ENALL | RIYFYD | HTENALL | YLKHFKEF | SCMKIDHCV | AVGSGKSTGL |
| LTN | AALL | WARSL | TGKTMW | NALLLYM | DVDPHYLK | YGTPMDFGQ | TYSPDTLGYD |
| TVA | LLGG | GGQYA | IRIYFY | TENALLL | DFGQVFNM | ERIQRLGRV | KQAIELLPDF |
| VNA | SLLT | KHFKE | KIRIYF | GKIWMDE | HYLKHFKE | IQIKGGIPT | KQLSFFWRPE |
| TLN | AGVA | LLLLL | LKIRIY | VGKRFCV | QEAGKYEN | SVYVLGKIW | LGKTTVVAIF |
| SGI | ALTG | NVIDD | RTGKTM | LLYMACT | VVYNHQEA | KLSTAKHSV | NLSRQLGKTT |
| DTV | GTLA | PVYAT | TLKIRI | ALLLYMA | PHLHVLIQ | DRINVRRLF | FLVRDRRPVD |
| GDV | DLIK | KYGKP | NPLYFK | KIWMDEN | MWARSLGP | KGKLKLSTA | KSCSQGGIRG |
| PLL | LLAA | VYATL | RFCVKS | DSRTGKT | GFTHRGTH | FKGKLKLST | TSLYPSIIRQ |
| AGV | ADLV | YATLK | GKRFCV | LLLYMAC | GKYENHTE | PETVHGFRC | LEGLRQKGWS |
| VSV | ELLL | THASN | KRFCVK | IWMDENI | LVRDRRPY | IPFRAPTVK | ALTPLRGSDP |
| AID | LAAV | DVTRG | KYENHT | HNYLCGH | RFFDLVSP | QELRVLAAL | FRADVKEFEA |
| NAV | DDVL | HTENA | VTGGQY | NYLCGHL | NHTNSVMF | NHGFTHRGT | GYDLIRDLIS |
| LLS | VLDA | THRVG | GKIWMD | NPLYFKI | ATVKNDLR | YFLTYPQCS | ILADGDDAGM |
| VVK | IAAG | LGVIS | NALLLY | YLCGHLD | VHGFRCML | KNDLRDRFQ | KQIKSRYGDK |
| VVS | LALA | MDENI | ALLLYM | LCGHLDL | HGFRCMLA | VQIKGGIPS | QQVPINATGS |
| LLQ | ALAL | LGPHN | ENALLL | IFLCNPG | TATVKNDL | APTVKILSK | SDPKNFQPVM |
| VDG | LVAA | KNHTN | YENHTE | GGIPTIF | CSLTKEEA | NVIRAVRFA | TSGSGMGKST |
| GVT | VELL | FCVKS | ENHTEN | GIPTIFL | SLTKEEAL | FASLYPSII | NEMDAGIYYA |
| VGT | LSLL | FTHRG | TENALL | IPTIFLC | CISDVTRG | TIHSRSYTH | DEIIDNSVDE |
| VLS | AVAL | INNVI | KIWMDE | LKHFKEF | CVSDVTRG | YVVYNHQEA | IHSRSYTHIM |
| LVS | DALV | QFEGK | LLYMAC | PTIFLCN | FGQVFNMF | APKDFVLQF | PGPNSSYKEF |
| SVL | LLSL | YGKPV | LLLYMA | QRDWQSN | FFLVRDRR | CMLAVKYLQ | MGGDFLTSLI |
| VLE | VLAG | YNVID | IWMDEN | ARSLGPH | IKTKNHTN | HFIVATNII | SEKGVSWAAE |
| IDT | LVSL | VIDDV | HNYLCG | KEFMGAQ | IVIEGDSR | VLCNPGEGA | VPRRHGKTWF |
| NIT | NALL | IKGGI | NYLCGH | KGGIPTI | LTKEEALS | VKNDLRDRF | WADNAVSFTA |
| DNI | LLLK | KGGIP | PHLHVL | KHFKEFM | ISINNVIR | GEMTVAGKK | GDFARPNLFE |
| VNG | ALGI | VGKRF | LKHFKE | WARSLGP | DENIKTKN | IRAVRFATD | LLVLKNNKGV |
| DTI | ALGV | FLTYP | LCGHLD | GAQRDWQ | MDENIKTK | GAGFGAGFG | NEQALVKRFW |
| INN | TLAL | LCNPG | PLYFKI | HFKEFMG | TRGNGITH | LPTSAGKSL | SLPIAGLEDI |
| RLV | GDVV | SNPVY | YLCGHL | AQRDWQS | KSVYILGK | EGRGQDYHA | IGKVMCISDV |
| NGV | AGVL | HASNP | CGHLDL | IKGGIPT | GNGITHRV | VNNYVVYNQ | LSLPIAGLED |
| GTI | VLGA | ATLKI | IFLCNP | CGHLDLS | IKLKNHTN | GFGAGFGAG | LSSSFDQVPE |
| VLR | GVVA | NPVYA | FLCNPG | FKEFMGA | ATNIIENG | YFLTYPKCS | REKIHGTNFS |
| TGI | LRLL | ACTHA | GAQRDW | TGGQYAS | NIIENGVT | ELRVLAALS | VPTLYFSADS |
| LFS | LSVL | GKRFC | KHFKEF | IDDVDPH | VATNIIEN | ERIVSILEW | YLDNLGVISI |
| ALL | LVLS | LYMAC | GGIPTI | PHNYLCG | VLQFHNLN | GDFLTSLIN | ISKRAGIGIN |
| SVI | VVAG | DWQSN | QRDWQS | GHLDLSP | LGVISINN | LNFQVWTTS | TPYLRLPIHD |
| VIE | VAGV | MACTH | IPTIFL | GPHNYLC | PKVYSNDA | RGARWAGES | FVLQFHNLNS |
| SLI | AGIA | YMACT | EFMGAQ | LGKIWMD | TNIIENGV | RKALGIHKC | IHAELNAILF |
| TVV | IAGL | CTHAS | GIPTIF | RSLGPHN | VISINNVI | YSIELAQDL | INESGLYSLI |
| ISI | LLAL | ALLLY | KGGIPT | WYNVIDD | GKVMCISD | INSLYGALG | LLAHVGYPRL |
| IVS | ALVG | KYENH | PTIFLC | KYGKPVQ | NIKLKNHT | NGLMVWCIE | RVTAEEIRYV |
| ITV | LVGA | HRVGK | RDWQSN | KPVQIKG | VYNHQEAG | VAFDMRGQQ | VSDVTRGNGL |
| VVA | LLTL | NHTEN | TIFLCN | NVIDDVD | GQYASKEQ | ALGPHNYLS | YGLNLHYIPP |
| VVN | VVVD | GHLDL | AQRDWQ | NYFLTYP | QYASKEQA | FLGLPFNIA | YGVFSTGISV |
| GVV | LTLL | GPHNY | GGQYAS | PVQIKGG | IIENGVTL | KICRELHED | LQTIGRVLRK |
| LIR | VALL | RFFDL | HFKEFM | VIDDVDP | KLKNHTNS | KICRELHEN | MGFKTRYGIG |
| LVA | GDIV | GKTMW | KEFMGA | YGKPVQI | VTRGNGIT | LPFNIASYA | PVSPMGCRSF |
| NVL | LVAL | NALLL | RSLGPH | RFCVKSV | QRLGRVGR | RALDNLLDY | QLIMKSKLPY |
| SII | LIAL | PHNYL | FKEFMG | FCVKSVY | INNVIRAV | SKEQALVKK | WKHFQTAVKS |
| VGV | LVLD | IYFYD | GHLDLS | HGFTHRG | YVLGKIWM | TSAGKSLIQ | WVVEFDPNIP |
| IIS | LVGL | KIRIY | DDVDPH | AWYNVID | MCISDVTR | TTLFLTEGD | ANTDCDGDKK |
| VVG | AVLV | KTMWA | YNHQEA | YLKHFKE | YDLIRDLI | WLAIQPVIS | DIARMYGVTP |
| LIA | IALL | IRIYF | VQIKGG | KNYFLTY | DLRDRYQV | AGFGAGFGA | EDLLIRVNEY |
| LIN | LAIL | LKIRI | KPVQIK | LIQFEGK | ENIKLKNH | AIELLPDFL | EGMATSIAEL |
| IIK | LVLL | KRFCV | GKPVQI | WQSNTKY | RRPYGTPM | LSGIKGQIG | GAKEAFHPMY |
| LVG | LLVL | TMWAR | PHNYLC | FGQVFNM | DENIKLKN | LYQSCHILQ | GPAGTGKTTL |
| NIL | VLLL | RIYFY | SLGPHN | FLCNPGP | NDLRDRYQ | NIFLAMLVN | IGRTWIQITW |
| VIA | LLLV | LLYMA | HLDLSP | HLDLSPK | SFFSLKDP | SKRYLYQDN | NGPAGTGKTT |
| GVI | LLIL | GIPTI | LGKIWM | DDVDPHY | YLSGHLDF | CGMYASALT | PAGTGKTTLT |
| GIV | LLLI | PLYFK | WYNVID | DVDPHYL | FFSLKDPI | TSLYPSIIR | PCNLGHINLA |
| IVG | LILL | YENHT | KNYFLT | EFMGAQR | NIDLHYFS | VLQFHNLNA | RVAHIHVVNG |
| NII | ILLL | RFCVK | YGKPVQ | VDPHYLK | SLKDPIPW | VNNYVVYNH | SINNVIRAVD |

# Table A13. Characteristic n-grams in disordered regions produced by combination of $z$-score, fractional difference and mole fractions 

N -grams presented in the table have mole fractions>1E-6, abs(z-score)>2.58 in disordered and abs(zscore) $<1.65$ in ordered regions. Table includes for each length first 100 n -grams sorted according fractional difference in disordered regions in descending order.

| N-gram length |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| QQQ | PSPP | GGGGG | GGGGGG | SSSSSSS | GGGGGGGG | PEPEPEPEP | SSSSSSSSSS |
| PPR | SSSS | PPPPP | PPPPPP | PPPPPPP | PPPPPPPP | EPEPEPEPE | EEEEEEEEEE |
| PPQ | EPEP | APAPA | TTTTTT | EEEEEEE | Eeeeeeee | EEEEEEEEE | HPNIQGAKSS |
| SPS | PQGP | PSPPP | PEPEPE | DDDDDD | PEPEPEPE | PKPAPKPAP | PSPPPPSPPP |
| TPP | PTPP | NNNNN | EPEPEP | PEPEPEP | EPEPEPEP | PAPKPAPKP | SPPPPSPPPP |
| PPK | PAPP | EEEED | QQQQQQ | EPEPEPE | KPAPKPAP | KPAPKPAPK | GGGGGGGGGG |
| DDD | APAP | PPAPP | GGGGGA | PKPAPKP | APKPAPKP | APKPAPKPA | PPSPPPSPPP |
| PPE | APKP | SSTSS | PKPAPK | TTTTTTT | PAPKPAPK | DDDDDDDDD | AATTTAATTT |
| SES | PKPA | KKKKK | KPAPKP | PPPSPPP | QGAKSSSD | QQQQQQQQQ | TTTAATTTAA |
| PDP | PPPA | DEEDE | AGGGGG | KPAPKPA | GAKSSSDV | PPPPSPPPP | VISTPASKVR |
| QKQ | KPAP | PAPPP | PPSPPP | PAPKPAP | PPPSPPPP | PSPPPPSPP | ATTTAATTTA |
| ESE | PPPR | KKSKK | PPPSPP | APKPAPK | QQQQQQQQ | PPSPPPPSP | RYGGGGGGGG |
| PEE | APPP | EEDDD | APKPAP | QQQQQQQ | NIQGAKSS | PAPAPAPAP | GDGDGDGDGD |
| RQQ | PPGP | PSPSP | PSPPPP | GGGGGGA | KSSSDVKS | APAPAPAPA | STNGIEPPRG |
| PNP | SPPS | PKPAP | APAPAP | PPSPPPP | GGGGGGGA | PSPPPSPPP | APAAPAAPAA |
| AQQ | PSSP | SPPPP | PAPAPA | PPPPSPP | PSPPPPSP | DSDSDSDSD | PADIIISTPA |
| PSA | PAPA | RSPSP | PPPPSP | MDSRTGE | PPSPPPPS | PPSPPPSPP | SLGSGLSMSG |
| SST | PPPT | PPPSP | NNNNNN | PAPAPAP | SPPPPSPP | TTAATTTAA | TDISLGSGLS |
| MEE | KKKK | QGPQG | SRSRSR | GAKSSSD | AGGGGGGG | ADIVISTPA | AGGGGGSGRR |
| EAE | SPSS | QGPKG | QGAKSS | QGAKSSS | APAPAPAP | ATTTAATTT | GAGGGGGSGR |
| SKS | EEDE | SPSPS | QGPKGD | RSRSRSR | ISTPASKV | TTTAATTTA | LMPCESSSQV |
| KEK | DEEE | PPSPP | GAKSSS | SRSRSRS | MSKRPADI | SKRPADIVI | PAAPAAPAAP |
| DED | SDSD | PQGPQ | SPPPPS | GPQGPQG | RGGQQTAN | YGGGGGGGG | AGGGGGGGGG |
| SPT | PPPL | EEEEE | SDSDSD | IQGAKSS | SPPPSPPP | NDAAAEALN | AAPAAPAAPA |
| RGP | KSSS | SSSSS | PQGPQG | PSPPPPS | PSPPPSPP | TGPQGPKGD | TSSSSSSSSS |
| REQ | EDDE | RRRSS | SSDVKS | SSSDVKS | TTAATTTA | GDGDGDGDG | GAGGGAGAGG |
| PTS | DDED | PPPPT | DSDSDS | NNNNNNN | TTTAATTT | GGAGGGGGS | APAAPAPAAP |
| DSE | RKRK | PPPAP | TGGGGG | APAPAPA | ATTTAATT | RYGGGGGGG | GGGGGGGGGY |
| MAD | SSST | PSPTP | GPQGPK | SPPPPSP | PAAPAAPA | STNGIEPPR | PAAPAPAAPA |
| GGS | EERK | GPPGP | PQGPKG | GPQGPKG | APAAPAAP | GAGGGGGSG | IISTPASKVR |
| AKK | DSDS | GAKSS | SPPPSP | SSDVKSY | EDEDEDED | DGDGDGDGD | GDRRRVRIEV |
| PGS | PSTS | DEEEE | PSPPPS | GGGGAGG | ELNPAPTS | DISLGSGLS | KPGGGGNGGH |
| NSS | SDSS | SDSDS | DEDDED | DSDSDSD | GGGGGGSG | SLGSGLSMS | KRRQKREDER |
| ASQ | KRKR | AKSSS | PPPPPS | PSPPPSP | GPEGPEGP | GAGAGGGAG | RELLDLARQQ |
| GGP | DEED | SSSPS | PPPLPP | SDSDSDS | AGGGGGSG | MPCESSSQV | SEGDRRRVRI |
| NPS | TPTP | PLPPP | TTAATT | GDKGDKG | ASSRASSR | PAAPAAPAA | TLTQQEQQAQ |
| ASR | SSES | GPTGP | PPPPLP | MSKRPAD | DGDGDGDG | SPPPSPPPS | VSEGDRRRVR |
| RPT | EEDD | TPPPP | SPPPPP | GPKGDTG | GAGGGGGS | AAPAAPAAP | AINALRRRLE |
| TES | SDSE | PTPSP | GGGGGY | DKGDKGD | ALRRRLER | ADIIISTPA | GDKGDKGDKG |
| TTP | TSSS | PAPTP | SSSTSS | STPASKV | LNPAPTSS | GGAGAGGGA | LTPSDWSFLK |
| PPL | DDDE | TGPQG | QGPQGP | SKRPADI | LRRRLERG | AGGGGGSGR | NLAAARASTQ |
| SQT | KKSK | PPPLP | PAAPAP | DEDEDED | NPAPTSSP | ATDISLGSG | ESARAVREGQ |
| DDP | ASSS | DSSSS | PPPPPA | TNGIEPP | DAAAEALN | GGGGGGGG | QEAPEWAPPK |
| SET | SSEE | EDEDE | MSKRPA | PTPSPTP | AAPAAPAA | SSSTPPSIK | EGPGGPPGPE |
| NSP | APTP | EEDEE | EEEEDE | GPAGPQG | GGGGGGGY | SSSSTPPSI | FTSSDLAFLK |
| TGP | SSTP | EDDED | GPKGDT | GGQQTAN | NALRRRLE | TATDISLGS | IPKEQARIDL |
| ENE | SSTS | EEEDE | SKRPAD | RGGQQTA | NDAAAEAL | DEDDEDDED | QGLGTEAPSN |
| GRS | EKKE | SSSSP | PKGDTG | GGSGGGG | PAPVPKPA | GAGGGAGAG | RLNKMLKGEK |
| AKE | KKEE | RRRSR | KRPADI | YGGGGGG | SSRASSRA | NSSSSTPPS | FQTTGLSKAK |
| GES | DSDD | RSSSS | LPPPPP | GQQTAND | APTSSPTS | SPPPSPPPP | KGGISQQPDI |
| GDP | DDSD | APPAP | EDDDDD | PTPPPTP | DEDDEDDE | AAPAPAAPA | PEESVGDTQM |
| QSG | KKRK | EDDDE | TPPPTP | TTAATTT | DKGDKGDT | APAAPAPAA | PKPAPVPKPA |
| RGG | SAPS | GPVGP | YGGGGG | GSGGGGG | EEQKQLTL | GAGGAGAGG | VRDALAGKRA |
| ADP | SSGS | DDEEE | EdEEEE | NGIEPPR | PPPLPPPP | HSTQVPIKV | ANLPTTHMPR |
| KRN | ESEE | STSSS | SSSSGS | GPEGPEG | NSTNGIEP | PAAPAPAAP | GGGGGGGGGA |
| SKT | KKEK | DDEDD | SSSSSS | PAAPAAP | RYGGGGGG | PQPQPQPQP | YEKKPRSVSQ |
| KST | SGSS | SSSDV | exeeee | PPPPLPP | DISLGSGL | AGAGGGAGA | YGGGGGGGGA |
| ESG | SSGG | EDDEE | PAPKPA | PQGPQGP | DTPVSEIP | AGGGAGAGG | PPPRHPGRRS |
| GDS | KSKK | TTTTT | RSRSRS | DDEDDED | GGEGGEGG | APAPAAPAP | AKGDKGEPGQ |
| DAE | ERRR | QGIQG | AKSSSD | QPEESVG | GGGGGSGR | DNPASAEAI | FKGAKGDKGE |
| GGR | EAEE | DEDDE | MDSRTG | SSRASSR | ILEEAQRL | GGGGGGSGG | GEGEEGGGEG |
| GAS | EKEK | EDEDD | GPQGPQ | ADIVIST | ISLGSGLS | QPQPQPQPQ | GGGGGGGAGY |
| ERG | RRGR | SEEEE | IQGAKS | PPPTPPP | SSSSSSSC | RTATETRRG | HKSGKNKGQP |
| NNS | EAER | APAAP | NIQGAK | DEDDEDD | GEGGEGGE | GGGGGGAGG | KDLTESQKEK |
| NNP | SAPA | DDDED | EDEDED | NPASAEA | SLGSGLSM | ERTATETRR | NSKFSEKKKS |
| TKA | SSDD | SSSAS | ARGGQQ | RELNPAP | AGAGGGAG | GGGAGAGGG | PELPSLDDID |
| SAG | nnns | SSSST | DDEDDE | QPQPEES | APAAPAPA | GGSGGGGGG | PSDWSFLKGI |
| STG | GGSS | DSDDD | PSSSSS | TSSSSSS | EEAQRLIH | LNKMLKGEK | ELRTERLERI |
| SGK | SSNS | SSDVK | DKGDKG | GGQQSAN | EGGEGGEG | LPTTHMPRQ | ESSYLDARHK |
| ANP | SGGG | TSSSS | GDKGDK | GQQSAND | GGGGSGRR | APPAPAPAP | GAGAGGGGGG |


| NPT | RRRA | PAAAP | SSSSSA | RGGQQSA | GPEEGEGP | DEEYYEEDR | LNENANKDSR |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| TAE | RRAR | ASSSS | STPASK | AGGGGGS | GPPGPEEG | DRAKANLAA | NGGNGGSSPT |
| NPA | EKAE | DEDDD | ISTPAS | EDDEDDE | PPPPLPPP | DRRRVRIEV | RGVSGNPRAD |
| GTS | EEGE | DDEDE | DKGDTG | GPQGPAG | QVSNSTNG | EDDEDDEDD | SGTPLRRYPM |
| ATR | PGGG | SSSSG | KSSSDV | IVISTPA | SGYRYGGG | ITPSGAVDD | AKFHSPKSPM |
| SLS | GSSG | PPAPA | PLPPPP | LKGSSST | SSQVSNST | KPGGGGNGG | ENDKTMFEKF |
| TDD | EARE | DDSDD | TPSPTP | PSPPPPP | TANDAAAE | KPLTQEHAD | LSDEQLEALL |
| SLP | ASTS | PAPVP | DDDDDD | PVPKPAP | TPSPTPSP | MGLIPTAPL | QGPQGIQGPQ |
| TAQ | STTS | IQGAK | PTPSPT | VKSYIDK | CESSSQVS | MPSESSSVV | SGAPEMSPAS |
| ATT | DDGD | GGSSS | PNIQGA | PASMEGN | ESSSQVSN | PGGGGNGGH | ADGGGDPEDI |
| RTA | STST | SGGGG | DEDEDE | PPPLPPP | KGSSSTSS | PTPPPTPPP | AGGAGAGGAG |
| ELR | DGDD | DGDDD | SARGGQ | RRRSSGG | QTANDAAA | QQEQQAQLD | LKQIQFKRSK |
| AGR | GASS | SSGGG | PAAPAA | SPASMEG | SSSQVSNS | QRELLDLAR | PAAAPAPSKP |
| QLR | RGGG | SDVKS | GDKGDT | ASMEGNR | GSGLSMSG | SDPREEQVS | SSSSSSSSGS |
| KTT | AKAK | GGGNG | EDDEDD | TPASKVR | GTSARRAE | ATNGIEPPR | EKAEKAAEKK |
| DEN | KAKA | GGGGN | DDDEDD | APAAPAA | YEEQKQLT | SRLIKASTS | GPQGSPGLNG |
| DTD | LSSS | SARGG | APAAPA | AAPAAPA | PAAPAPAA | VPEVPEVPE | SGGAGGTTSI |
| PVE | EARA | GSSGG | GFGSTG | RSARGGQ | PELPSLDD | KRPPPRHPG | TGGKGGNGGS |
| NGS | TSSA | GSGGG | SSSASS | SDWSFLK | QSGTSARR | KTLAELEAE | TTNNNSTNND |
| VPE | TTST | AASSS | HPNIQG | ASKVRRR | SGTSARRA | LSTPSLPPA | VGGGGGGGAG |
| DLE | STTT | GGRGG | PAAAPA | ATTTAAT | APAPAAPA | PEVPEVPEV | AAAPAAVAAD |
| DTT | VSSS | DDDGD | GSGGGG | SDVKSYI | SQLKGSSS | PKPKPAPKP | RPMNRKPRMY |
| ETG | REAA | NGGGG | AAPAAP | TAATTTA | GRSARGGQ | QRQAPQGAQ | KLNERTATET |
| TGR | EVEE | LSSSS | GGGGAG | GGSGGSG | AATTTAAT | TLAELEAEA | MSGLLDDGAN |
| ELA | GSTG | SSLSS | GGSGGS | GAGAGGG | TAATTTAA | ASAYNGNDT | KEGIPPDQQR |
| ALE | GSAA | AAAEA | SGGSGG | GGAGAGG | IIISTPAS | EGDRRRVRI | QVPIKVQHRL |
| TTG | GTGG | GGTGG | GGGAGG | AATTTAA | GAGAGGGA | AGSAAGSAA | GQHISIRTFR |
| AQL | GAAA | AGAGG | AGAGGG | SIRTFRE | QSANDAYA | GESWASRST | CQSANDAYAE |
| LAP | AAAV | AKAAA | GGGAGA | GGAGGAG | MSDVVERA | SANDAYAEA | KVRRRLNFDS |
| DVE | AAAL | AGAAA | GGAGAG | GQHISIR | RPMNRKPR | VSEGDRRRV | VRRRLNFDSP |

## Table A14. Characteristic n-grams in disordered regions produced by association rules

N -grams presented in the table belong to the body of association rules with head ORDER_LEVEL=' $\mathrm{D}^{\prime}$. Parameters used in mining are confidence $>=51 \%$, support $>=0.0001$ and lift $>=1.05$ or lift $<=0.95$. Except for n-gram with length two where only one rule exists, table includes for each length first 100 n grams, sorted according lift and confidence, both in descending order.

| N-gram length |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| PP | PPP | GHMA | AAPPA | AAPAPA | AAPAPAA | ADTPVSEI | ADIVISTPA | ADIVISTPAS |
|  | QQQ | GSHM | APAPA | AAPPPP | ADTPVSE | AGGGGGGG | ADTPVSEIP | APAPAPAPAP |
|  | PSP | HHHH | APEDP | AGPQGP | AGAGGGG | AGGGGGSG | AGGGGGGGG | APKPAPKPAP |
|  | SPP | SNAM | DDDDK | APKPAP | AGGGGGG | AGTSKVSR | ALRRRLERG | ARGGQQSAND |
|  | PAP | PPPP | DDDKD | APPAPP | AGGGGGS | ALRRRLER | APAAPAAPA | DDDDDDDDD |
|  | SSS | PSPP | DEDSD | APPPPP | AGTSKVS | APAPAPAP | APAPAPAPA | DIVISTPASK |
|  | PQP | QQQQ | DEEDE | APTPPP | ANDAAAE | APKPAPKP | APKPAPKPA | DSDSDSDSDS |
|  | PKP | QPQP | DSDEE | AQQQQQ | APKPAPK | APTSSPTS | ARGGQQSAN | EEEEEEEEEE |
|  | PPS | EPEP | DSPPS | AQRLIH | APPPPPP | ARGGQQSA | DDDDDDDDD | ELNPAPTSSP |
|  | QQP | SSSS | EAEED | ASMEGN | APTSSPT | DDDDDDD | DGDGDGDGD | EPEPEPEPEP |
|  | PEP | PPPS | EEDDD | CESSSQ | ASGGGGG | DEDDEDDE | DIVISTPAS | ESILEEAQRL |
|  | QPP | EEEE | EEEED | DDDDDS | ASSSSSS | DEDEDEDE | DKGDKGDTG | GGGGGGGGGG |
|  | RPP | PPSP | EEEEG | DDDDGD | DDDDDD | DIVISTPA | DSDSDSDSD | GYRYGGGGGG |
|  | PPR | SPSP | EKKKS | DEEDEK | DDEDEDD | DKGDKGDT | EEAQRLIHG | ILEEAQRLIH |
|  | EEE | PQPQ | ESSSS | DEEEED | DSSSSSS | DSDSDSDS | EEEEEEEEE | IVISTPASKV |
|  | EPP | QGPQ | GGGDD | DSDSDS | DTPVSEI | DTPVSEIP | EEQKQLTLF | KPAPKPAPKP |
|  | QPQ | PPQQ | GGGGG | DTGPQG | EAQRLIH | EAQRLIHG | EPEPEPEPE | LEEAQRLIHG |
|  | PPQ | SPPP | GGSRS | EDEDEE | EDDEDDE | EDEDEDED | ESILEEAQR | MSKRPADIVI |
|  | PRP | PQGP | HHHHH | EDEEDE | EDEEDEE | EEEEEEEE | GAGAGGGAG | NSGYRYGGGG |
|  | PQQ | QPQQ | KKEKK | EEEEDD | EDEEEEE | EEQKQLTL | GAGGGGGGG | NSTNGIEPPR |
|  | PPA | PPPQ | KKGKS | EESVGD | EEEEEEE | EGPEGPEG | GAGGGGGSG | PADIVISTPA |
|  | GPP | PQPP | KKKAA | ENTERH | EEQKQLT | EPEPEPEP | GDGDGDGDG | PADTPVSEIP |
|  | SSP | PEPP | KKKKK | GGGGGG | EGPEGPE | GAGGGGGG | GEGGEGGEG | PAPAPAPAPA |
|  | APP | PPQP | KKSKK | GGGGGL | ENTERHT | GAGGGGGS | GGAGGGGGS | PAPKPAPKPA |
|  | SPS | RPPP | KKTSS | GGGGGN | EPEPEPE | GDGDGDGD | GGGGGGGAG | PEPEPEPEPE |
|  | MSK | PSPS | KPTPP | GGGGGQ | GDTGPQG | GGAGGGGG | GGGGGGGGA | PKPAPKPAPK |
|  | PTP | PTPP | KRPPP | GGGGGS | GGGGGAS | GGEGGEGG | GGGGGGGGG | PPPPPPPPPPP |
|  | PGP | QQQP | KSASS | GGGGGV | GGGGGGA | GGGGGAGG | GGGGGSGRR | PPPPSPPPPS |
|  | KPP | GPQG | MEEEE | GPEGPE | GGGGGGG | GGGGGGAG | GYRYGGGGG | PPPSPPPPSP |
|  | TPP | QPPQ | NNNNN | GPPGPE | GGGGGGR | GGGGGGGA | ILEEAQRLI | PPPSPPPPSP |
|  | EPE | PAPK | NSSSS | GPVGPQ | GGGGGGV | GGGGGGGG | IVISTPASK | PPSPPPPSPP |
|  | PSS | PQQQ | NTERH | GQQSAN | GGGGGGY | GGGGGGGS | KGDKGDKGD | PPSPPPSPPP |
|  | RRR | PAPP | PAATS | GRRRSS | GGGGGSG | GGGGGGG | KPAPKPAPK | PSPPPPSPPP |
|  | PPK | QQPP | PAPPP | GYRYGG | GGGGNGG | GGGGGGSG | LEEAQRLIH | QQQQQQQQQQ |
|  | EEP | QQEE | PEPPS | KPAPAP | GGGGSGG | GGGGGSGG | LNPAPTSSP | RSARGGQQSA |
|  | PRR | QPPP | PKPRP | KQLTLF | GIEPPRG | GGGGGSGR | MPKRDAPWR | RSRSRSRSRS |


|  | QGP | QQPQ | PPAAP | KRDAPW | GPAGPQG | GGQQSAND | MSKRPADIV | RYGGGGGGGGG |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | RPR | PPAP | PPAPP | LPPPPP | GPEGPQG | GPAGPQGP | NATNGIEPP | SARGGQQSAN |
|  | MSS | PPPK | PPPPP | MNETEL | GPQGLQG | GPEGPEGP | NSGYRYGGG | SARGGQQTAN |
|  | MKK | PKPP | PPPPQ | MRSSSP | GPQGPKG | GPQGPKGD | NSTNGIEPP | SDSDSDSDSD |
|  | PPT | QSQP | PQQQP | MSKRPA | GPQGPQG | GPQGPQGP | PADTPVSEI | SGYRYGGGGG |
|  | QQE | GGGG | PSPEP | NATNGI | GPTGPQG | ILEEAQRL | PAPAPAPAP | SILEEAQRLI |
|  | QEE | MWDP | PSPPP | NGGGGG | GPVGPQG | IVISTPAS | PAPKPAPKP | SKRPADIVIS |
|  | SQP | APAP | QQEEE | NKNYGH | GQQSAND | KGDKGDKG | PEPEPEPEP | SPPPPSPPPP |
|  | QQR | KPAP | QQPPQ | NNNNNN | GSGGGGG | KGDKGDTG | PKPAPKPAP | SRSRSRSRSR |
|  | DDD | PGPP | REEEE | NPASAE | GSSSSSS | KPAPKPAP | PPPPPPPPP | SSQVSNSTNG |
|  | MSE | PEPE | RGEET | NTERHT | GTSKVSR | LEEAQRLI | PPPPSPPPP | ssssssssss |
|  | GPQ | GPPP | RRRGR | PASMEG | KGDKGDK | LRRRLERG | PPPSPPPPS | TTTTTTTTTT |
|  | RRS | DDDD | SKKKK | PEGPQG | KGDKGDT | MPKRDAPW | PPPSPPPSP | VISTPASKVR |
|  | QQS | PRPP | SPSPG | PERGSG | KPAPKPA | NATNGIEP | PPSPPPPSP | YRYGGGGGGG |
|  | QEQ | PPPA | SSEKP | PKGDTG | KRDAPWR | NGIEPPRG | PPSPPPSPP | KRPADIVIST |
|  | KPK | PKPA | SSSVD | PKPAPK | KRPADIV | NPAPTSSP | PSPPPPSPP | RPADIVISTP |
|  | SEP | PSQP | SSTSS | PKPKPA | LRRRLER | NSTNGIEP | PSPPPSPPP | DIIISTPASK |
|  | QSP | PQQP | STTST | PPAPPP | MDSRTGE | PADTPVSE | QGPKGDKGD | GRSARGGQQS |
|  | EQE | PPPR | THMPR | PPPAAP | MPKRDAP | PAPAPAPA | QQQQQQQQQ | IQGAKSSSDV |
|  | RRP | PPGP | TPEPP | PPPPPQ | NGIEPPR | PAPKPAPK | RGGQQSAND | ARGGQQTAND |
|  | PKK | SSSP | PSPSP | PPPPPR | NSTNGIE | PEPEPEPE | RSARGGQQS | AVSQLKGSSS |
|  | GGG | APKP | PQPQP | PPPPPPV | PAAPAPA | PKPAPKPA | RSRSRSRSR | NALRRRLERG |
|  | KRP | QGPK | PKPAP | PPPPSP | PAGPQGP | PKRDAPWR | RYGGGGGGG | NIQGAKSSSD |
|  | EPS | PPRR | QGPQG | PPQPQP | PAPTSSP | PPPLPPPP | SARGGQQSA | RELNPAPTSS |
|  | MSN | APPP | PEPEP | PQGPAG | PEPEPEP | PPPPPPPPP | SARGGQQTA | GDKGDKGDTG |
|  | MSD | QPEE | RPPPP | PQGPQG | PPPPLPP | PPPPPPPPS | SDSDSDSDS | RSARGGQQTA |
|  | RPS | PAPA | KPAPK | PSPPPP | PPPPPPPA | PPPPSPPP | SGGGGGGGG | GAKSSSDVKS |
|  | RSP | PPSS | PPPSP | PSPSPS | PPPTPPP | PPPSPPPP | SGYRYGGGG | ISASAYNGND |
|  | EPQ | SPQP | QPQQQ | PSPTPP | PPSPPPP | PPPSPPPS | SILEEAQRL | SASAYNGNDT |
|  | EQQ | PKPK | QQPQQ | PSPTPS | PQGIQGP | PPSPPPPS | SKRPADIVI | PNIQGAKSSS |
|  | KKK | RSPP | PEPPK | PTPPPT | PSPPPPP | PPSPPPSP | SPPPPSPPP | KRPADIIIST |
|  | EEQ | PSSP | TERHT | QGAKSS | PSPPPSP | PSPPPPSP | SQVSNSTNG | RPADIIISTP |
|  | SPQ | SPPS | PQGPQ | QGIQGP | PSPSPSP | PSPPPSPP | SRSRSRSRS | QGAKSSSDVK |
|  | SQQ | PPTP | SPPPP | QGPQGP | PSPTPSP | PSSSSSSS | SSQVSNSTN | SAVSQLKGSS |
|  | PPE | KRPR | PPPPS | QKQLTL | PVPKPAP | QGAKSSSD | sssssssss | HPNIQGAKSS |
|  | SRS | SSPP | QPQPE | QPEESV | QGAKSSS | QGPKGDKG | STNGIEPPR | GRSARGGQQT |
|  | QPS | PPEP | SPSPP | QPQPEE | QGPKGDT | QGPKGDTG | TGPQGPKGD | RGGQQTANDA |
|  | MTT | PPPT | RPPSP | QQQQQP | QPQPEES | QGPQGPQG | TNGIEPPRG | SPASMEGNRP |
|  | QSS | RRRS | MSKRP | RDAPWR | QQQQQQQ | QQQQQQQQ | TTTTTTTTT | ISTPASKVRR |
|  | RKR | PNPP | PPPPR | RGGGGG | RGGGGGG | RGGQQSAN | VISTPASKV | PQPGQHISIR |
|  | MPP | QQQR | PQGPK | RGRGRG | RGGQQSA | RNKNYGHP | YGGGGGGGG | SAHFHPNIQG |
|  | SES | PKPS | NPPPP | RNKNYG | RNKNYGH | RSRSRSRS | YRYGGGGGG | AHFHPNIQGA |
|  | EPK | SPTP | RRSSS | RPADIV | SDDDDD | RYGGGGGG | ARGGQQTAN | QPGQHISIRT |
|  | MSQ | EPPP | QQQRQ | RPGRPR | SGGGGGG | SARGGQQS | KRPADIVIS | HFHPNIQGAK |
|  | EES | PEEP | PPEPE | RSSSPS | SPASMEG | SARGGQQT | PADIVISTP | AKSSSDVKSY |
|  | SPR | RRPP | QQQQP | SDSDSD | SPPPPPP | SDSDSDSD | RPADIVIST | SSSDVKSYID |
|  | ESS | EEED | PPQPQ | SNSSSS | SPPPSPP | SGGGGGGG | IQGAKSSSD | FHPNIQGAKS |
|  | PSR | KPPP | EEEEE | SPRRRR | SPSPPPP | SKRPADIV | ELNPAPTSS | SDVKSYIDKD |
|  | APA | QPAP | QPQPQ | SPSPPP | SSGGGGG | SNSTNGIE | DIIISTPAS | SSDVKSYIDK |
|  | EED | HPPP | RQQQQ | SPSPSP | SSSSGSS | SPASMEGN | IIISTPASK | PGQHISIRTF |
|  | RSR | PKPQ | PPPEP | SPTPPP | SSSSSDS | SPPPPSPP | QGAKSSSDV | DGRSARGGQQ |
|  | KRK | SPPQ | PPKPK | SPTPSP | SSSSSSA | SPPPSPPP | VSQLKGSSS | STPASKVRRR |
|  | QPR | SSPS | PQQQQ | SSSQVS | SSSSSSD | SRSRSRSR | SPASMEGNR | ISIRTFRELN |
|  | APS | PPKK | QQEQQ | SSSSSE | STNGIEP | SSSSSSSC | RELNPAPTS | QHISIRTFRE |
|  | QPA | QPRP | RPPPR | SSTPPS | STSSSSS | SSSSSSSD | AVSQLKGSS | HISIRTFREL |
|  | KKP | QRQQ | PQQPP | STPPSI | TGGGGGG | SSSSSSSS | NIQGAKSSS | GQHISIRTFR |
|  | SSE | QPPR | QQQQR | TPAPAP | TGPQGPK | STNGIEPP | GDKGDKGDT | DVKSYIDKDG |
|  | RSS | AQQQ | SSSSS | TPPPPP | TNGIEPP | TGPQGPKG | RSARGGQQT | RSAHFHPNIQ |
|  | PDP | PEPS | PAPEP | TPPSIK | TPSPTPS | TNGIEPPR | RGGQQTAND | KSSSDVKSSYI |
|  | SSQ | PSSS | RQQRE | TSETNA | TPVSEIP | TSSSSSSS | AKSSSDVKS | MYRMYRSPDV |
|  | QRQ | QPQA | PPPTP | TSSPTS | TSSPTST | TTTTTTTT | ISASAYNGN | TQVPIKVQHR |
|  | SRP | QEEP | EPEAP | TTAATT | TTTTTTT | VISTPASK | SASAYNGND | ADIVISTPAS |
|  | PSQ | PRRR | QQQQQ | TTTTTT | VISTPAS | VQPQPEES | ASAYNGNDT | APAPAPAPAP |
|  | KRR | PSQQ | QPPQP | YGGGGG | VQPQPEE | YGGGGGGG | RPADIIIST | APKPAPKPAP |

## Table A15. Characteristic n-grams in ordered regions produced by association rules

N -grams presented in the table belong to the body of association rules with head ORDER_LEVEL=' $\mathrm{O}^{\prime}$. Parameters used in mining are confidence $>=51 \%$, support $>=0.0001$ and lift $>=1.05$ or lift $<=0.95$. Except for n-gram with length two where only one rule exists, table includes for each length first 100 n -grams, sorted according lift and confidence, both in descending order.

| N-gram length |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| WW | CWF | ACDW | AFGVL | AAKYEN | AAELRNF | ADGSQFDS | AFDICGVQP | AGPSKHFKSN |
| WC | CWW | ADWW | AFVGL | ADGSQF | ADSDAFT | AGKYENHT | ALEAIRFYV | AKYENHTENA |
| CW | CWY | AWCV | AGKYE | AEDPYI | ALDNLLD | ALDNLLDY | AVKSCSQGG | AQDLRAVHGM |
| CF | CYW | AWNY | AIFLA | AMSRRY | ALLFTWR | ARVATGRE | CAITHIDYG | AWCLMLISRG |
| WI | FWW | AWVC | AIIGI | ASYALL | ALLLYMA | ASGLADAL | CMLAIKYLQ | CGHLDLSPKV |
| YW | HWW | AYAC | AQAFF | CAWCLM | ARVATGR | ASSPDAVR | CMLAVKYLQ | CRELHENGEP |
| IC | WCY | CAVY | ATAYL | CDADGS | ASYGVFS | AVSQDQTK | CNIDLHYFS | CVSDVTRGNG |
| CI | WIC | CCEY | AVGYV | CLVWDI | ATNIIEN | AWYNVIDD | CSTLKDLIE | DADGSQFDSS |
| CY | WYW | CCIY | AVYEV | CNIDLH | AVEDLVN | CAWCLMLI | CVIEYRQQV | DAWYNVIDDV |
| FW | YCW | CDHI | AYTVL | CVIEYR | CRELHED | CMLAIKYL | CVKSVYVLG | DFASLYPSII |
| IW | YWI | CELF | DIAVG | DAAPYI | CVIEYRQ | DFASLYPS | DADGSQFDS | DLHYFSSSFF |
| WY | IWY | CFDI | DLLYI | DLECGC | DGSQFDS | DHCVIEYR | DFLQPGIVE | DRRPYGTPMD |
| YC | CLW | CFVL | DNWID | DLTSLY | DKRMTDN | DIPFRAPT | DFVLQFHNL | EMTVAGKKFF |
| WF | FYW | CGHI | DYGLY | DNLLDY | DNSLFEI | DKYNDVNR | DLRDRYQVM | FASLYPSIIQ |
| FC | CFW | CICD | EIKDY | DSIAWL | DSDAFTQ | DLIRDLIS | EAAKYENHT | FCVKSVYILG |
| VW | WWF | CLMT | EYPLK | EEYTRL | DVVVDFG | DLPCGCSY | ELFGARIHS | FDELFGARIH |
| LW | WWY | CNAF | FDINN | EFMGAQ | EATDTSF | EAGKYENH | ELPRILVDH | FDRINVRRLF |
| CC | WVC | CNCF | FFLAL | EGKYQC | ECLPNVC | EFLRETWT | ETSLWTLPD | FLVRDRRPVD |
| WV | FCW | CYYT | FGLIA | EGVFSL | ERIQRLG | EFMGAQRD | ETYCAITHI | FQPVMGFKTR |
| VC | YFW | DYCV | FLLGV | FGARIH | EWRVYLG | ELPRILVD | FASLYPSII | FWLVRDRRPY |
| WL | CWV | EWCG | FTDFP | FHNLNS | FDEILEG | FCVKSVYI | FHNLNANLD | GAIDLPPLGG |
| YF | CVW | FFMF | FTLAV | FIENET | FQVWTTS | FFSLKDPI | FLGLPFNIA | GARIHSHGNL |
| YI | WWI | FFWG | FYGLR | FIVATN | FYAKVTG | FGARIHSH | FRCMLAVKY | GFRCMLAVKY |
| CV | WMF | FHLC | FYSGL | FTLEKS | GCLVWDI | FIIASRNV | GFRCMLAVK | GGDFLTSLIN |
| LC | CYL | FIDC | GAYYG | GKTTVV | GCSTLKD | FLRETWTR | GGQYASKEQ | GKEFLRETWT |
| CL | LFW | FIHC | GDIVY | GNGLTH | GDFARPN | FRCMLAVK | GLPFNIASY | GKIWMDENIK |
| YY | WIY | FIMV | GFPAV | GRGQDY | GDTDSVF | FSLKDPIP | GPVAFSHFD | GKTMWARALG |
| FY | CYI | FIWL | GIVNV | HFKEFM | GKYENHT | FSSSFFSL | GQIYKHACA | HDKRMTDNES |
| FI | WII | FKLC | GQLIA | HGMDAD | GPLCKGD | FVHPVGFG | GRETCAWCL | HNLNSNLDRI |
| IY | WFW | FTIC | GRVTL | HLDFNS | GPPDTGK | GFRCMLAV | GSQFDSSLT | HTNSVMFWLV |
| IF | WQW | FWLF | GSLLI | HNLRKA | GPVAFSH | GHLDLSPK | GVISINNVI | HYLKHFKEFM |
| VY | CYC | FYDC | GYAVI | HTFDEL | GQVFNMF | GKEFLRET | GYSQGAIVT | ICRELHENGE |
| YV | YWF | FYGC | HLLAF | HTNTVM | GQVFNMY | GKSLGLCS | HTNSVMFFL | IIENGVTLDI |
| II | WCV | FYVC | IAAVI | IARGDS | HGEMTVA | GKTTVVAI | ILVYVASYN | IKICRELHED |
| FV | WFI | FYVM | IAMAL | IGIGHL | IENGTSP | GLPFNIAS | IVYFAETYC | IPSIVLCNPG |
| FF | WMC | GCYL | IECNG | ILEWDR | IKGGIPS | GVGPLCKG | KFKGKLKLS | IVKPFPFLAD |
| YL | YWC | GIWF | IFINY | IPFRAP | IKGLGSL | HACATGSG | KIWMDENIK | IYKHACATGS |
| LY | YVW | HLCI | IFNNG | ISKNAL | ILGKIWM | HDDLVMSL | KNDLRDRFQ | KALGIHKCFL |
| VF | CCW | HLWA | IIASR | IVHFKE | IPFRAPT | HNYLCGHL | KNHTNTVMF | KNFQPVMGFK |
| HW | IIC | HWLL | IIDIS | KELAPK | IQFEGKF | INAKNYFL | KPVQIKGGI | KNHTNSVMFW |
| CM | YFC | HWVL | IITSL | KGKLKL | IQFEGKY | INVRRLFN | KRYLYQDNE | KSVYILGKIW |
| WH | IIW | HYYF | ILGYA | KIWMDE | IQRLGRV | IPSIVLCN | KSCSQGGIR | KTKNHTNTVM |
| FL | VYW | ICTI | ILIGI | KTLITG | KILSKQF | IRCNIDLH | LADALVILA | KVQSFESRHD |
| IV | VWI | ILWY | INIFL | LAADIA | KIWMDEN | ITHRVGKR | LCGHLDLSP | LARYAFDFYE |
| VI | FYC | IWFV | IVCLL | LGGVYS | KPVQIKG | IVSILEWD | LCGPVAFSH | LGYDLIRDLI |
| CH | CMW | IWIL | IVTLV | LGIILL | KQLSFFW | KEFLRETW | LEAIRFYVS | LHENGEPHLH |
| LI | FWC | IYWE | KHNLV | LGNDLR | KSVYILG | KFIKICRE | LFVNILRLE | LINSLYGALG |
| LF | ICW | KCYG | KIAYT | LGQQLS | LCGPVAF | KGKLKLST | LGPHNYLSG | LKHWKELIGA |
| IL | VWC | LCHF | KPDFV | LGVVPV | LCSLAAD | KIRIYFYD | LKLSTAKHS | LLLYMACTHA |
| AW | CFI | LFCL | LAIFL | LGYTDA | LEGVNGE | LAELCGPV | LNSFTLEKS | LVYVASYNEV |
| HC | WYF | LQWW | LGLVN | LHVLIQ | LETSLWT | LGKTTVVA | MDENIKTKN | LWTLPDNPLD |
| WM | HFW | LWCA | LIRLF | LIAAAP | LKIRIYF | LNSNLDRI | MTDNESLQA | MTDNESLQAS |
| WT | WAW | LWFM | LITTM | LIQFEG | LKNHTNS | LPTPIMAG | NAKNYFLTY | MWARSLGPHN |
| TW | VFW | LWHT | LLIDI | LKTGMY | LLIRVNE | LRDRYQVM | NALEAIRFY | NALLLYMACT |
| CT | WLY | LWIY | LLLDI | LLALLA | LLTVGHP | LRRLGAPI | NFQPVMGFK | NCKYGKPVQI |
| WA | YIC | LWVV | LLLGI | LLVGYG | LLVYYCW | LSGIKGQI | NNYVVYNQQ | NDLRDRYQVM |
| CN | YCF | LYIC | LLLIV | LMLISR | LMLISRG | LTYPKCSL | NPGEGASYK | NIKLKNHTNS |
| WN | CCI | LYNC | LLTTF | LPCGCS | LNSNLDR | LVRDRRPY | NVIRAVRFA | NIKTKNHTNS |
| NW | CYF | NFTC | LRLIV | LPIHDE | LPRILVD | MKIDHCVI | PETVHGFRC | NLPTSAGKSL |
| GW | IFW | NWYI | LSVII | LSFDVT | LRDRFQV | NDAWYNVI | QEAGKYENH | NLSRQLGKTT |
| VL | IWW | NYLW | LTLVG | LSHDLT | LSGIKGQ | NLNANLDR | QPGIVEWNK | NNVIRAVRFA |
| AC | YCL | PQLW | LVFLA | LYNGGP | LSNALYG | NYKYQYDK | RFWKVNNHV | PFRAPTVKIL |
| LV | FIW | PWKF | LVNRA | MALPPC | MCISDVT | PFLADNSP | RGNGITHRV | PHNYLCGHLD |
| CA | CVI | QVGC | LVYAL | MLYMAC | MEGGGVG | PGIVEWNK | RIQRLGRVG | PKDYVLQFHN |
| YM | WLC | RWAW | NLIYQ | NDVNRW | MKIDHCV | PKNFQPVM | RLMEGGGVG | PKVYSNDAWY |
| GC | WCI | RWFP | NLMIL | NHNLRK | MTDNESL | PNIMMNNN | RPYGTPMDF | PVQIKGGIPT |
| NC | WHY | RWGW | QIVAL | NLHYIP | NDAWYNV | PTIFLCNP | SAFDRINVR | PYVALTPLRG |
| HY | YCI | TVCH | REGWA | NLPRIA | NHQEAAK | PVQIKGGI | SLQASWTFP | QIKGGIPTIF |
| WG | CYV | TWAI | RIVAI | NPVYAT | NIFLAML | QFEGKFQC | SNPVYATLK | QLGKTTVVAI |


| YH | FFW | VCVL | RLVRV | PKCSLT | NISPETI | QKDWQSNC | SVYVLGKIW | QQEAGKYENH |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| CG | YIW | VFWD | RVGIV | QIMAHF | NLDRIFT | RAPTVKIL | TGGQYASNE | RLETSLWTLP |
| TC | YWV | VHCL | RVLAW | QRTHFA | NSNLDRI | RCMLAVKY | THFAKFKGK | RQQVPINATG |
| VV | WIW | VLWH | RVNNY | QWAYDN | NSVMFFL | RCNIDLHY | TKYGKPIQI | RRIGVGHLGV |
| HF | WCW | VSMC | SFPNI | RDSIAW | NWRNIVK | RDDIVYFA | TTTPNGLNH | SLQASWTFPI |
| CD | WCC | VTMC | TDTSF | RFKGTV | PIPWKLY | REWRVYLG | TVVDNTLMV | SPKVYSNDAW |
| FM | CWC | VYIM | TGKIY | RHGKTW | PSIVLCN | RGQQKRFA | TYPQCSLTK | TAVKSCSQGG |
| FH | WIV | VYWG | VAIVV | RTHFAK | PVISSGR | RIGVGHLG | VAGKKFFLC | TGRETCAWCL |
| CK | IYW | WACL | VALLY | RVFKTQ | PWKLYYR | RINVRRLF | VEIHDKRMT | TKNHTNSVMF |
| QW | CIY | WFFN | VDLFY | SLGYIG | PYVALTP | RLETSLWT | VHGFRCMLA | TKNHTNTVMF |
| YT | ICC | WICK | VIIII | SNPVYA | QFPSTAS | RVNNYVVY | VKNDLRDRF | TMWARALGPH |
| YA | CFY | WISI | VITLG | SRSCMK | RETCAWC | RYQVLRKW | VLGKIWMDE | TVAGKKFFLC |
| LL | WHC | WIVY | VLAFL | SVIVEI | RGTGLTH | SLARYAFD | VLIQFEGKY | TVKNDLRDRY |
| YG | FWV | WKAC | VLTCL | TFVSFD | RKALGIH | SLPTPIMA | VMCISDVTR | VDIPFRAPTV |
| IH | VWY | WLWR | VTRDI | VGHPYF | RTFTAAP | SNDAWYNV | VSDVTRGNG | VEIHDKRMTD |
| GY | YLW | WMIN | VVCTN | VIDRNE | SFFHGEM | SYSLKEKE | VWGPSAPDA | VIDDVDPHYL |
| WD | VCY | WQQC | VVDGI | VISSGR | SHQYGGT | TLGYDLIR | VYKKAQAFD | VISINNVIRA |
| KC | WLW | WSFI | VVGLL | VLCNPG | SQFDSSL | TRGNGLTH | VYVASYNEV | VKNDLRDRFQ |
| KW | YYW | WWYA | WLQRA | VSEGIH | SRDEGLH | TYPKCSLT | VYVLGKIWM | VNNHVVYNHQ |
| YN | VIW | WYKF | WSGKE | VVVVDR | SSAVEDL | VGIAVDTG | WCIENGTSP | VNNYVVYNQQ |
| TY | VYC | WYLF | WYQRS | WIVIHA | TAGYTPF | VGKSLGLC | WMDENIKTK | VQIKGGIPTI |
| IG | ICY | YCFG | YDMYR | WSGKEF | TDNESLQ | VKRFWKVN | WSGKEFLRE | VYNHQEAGKY |
| QC | YLC | YCTF | YDVIK | YCAITH | TFVSFDL | VKSVYILG | YCDADGSQF | WARVATGRET |
| HI | FWF | YFAI | YFIRL | YDLIRD | VGKVMCI | VLCNPGEG | YFLTYPQCS | WTFPIRCNID |
| DC | WLF | YFKW | YIKKY | YGNDLR | VKILSKQ | WARVATGR | YGKPVQIKG | WYNVIDDVDP |
| RW | FWI | YINW | YISDI | YHAKRF | VKSCSQG | YCAITHID | YKHACATGS | YASKEQALVK |
| DW | WTC | YLCF | YITDI | YIDQYA | VKSVYIL | YGTPMDFG | YLCGHLDLS | YCAITHIDYG |
| AY | LWW | YMYY | YIVEL | YIKICR | WLAIQPV | YIKGLGSL | YMACTHASN | YCDADGSQFD |
| FT | WFY | YNWA | YKHAC | YPTASA | WMDENIK | YPQCSLTK | YNHQEAGKY | YFLTYPKCSL |
| NY | WIF | YRWD | YLDFA | YVDSRI | YAKVTGG | YQSCHILQ | YQDNERVAH | YGTPMDFGQV |
| IM | WDC | YWIT | YNVLR | YYIDLE | YTLGQQL | YRQQVPIN | YYFHGHIVP | YQYDKYNDVN |

## Table A16. Characteristic n-grams in border regions produced by association rules

N -grams presented in the table belong to the body of discovered association rules with head ORDER_LEVEL=' $N$ '. Parameters used in mining are confidence $>=51 \%$, support $>=0.0001$ and lift $>=1.05$ or lift $<=0.95$. N-grams in table are sorted according lift and confidence, both in descending order.

| N-gram length |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 5 | 6 | 7 | 8 | 9 | 10 |
| VYKYE | CHLKNP | EGNRPTF | EGNRPTFV | GQVVYKYEE | ASMEGNRPTF |
| WDPLV | FYDSIT | FYDSITN | GQVVYKYE | IYFYDSITN | IRIYFYDSIT |
| NRPTF | FYDSVT | FYDSVTN | IYFYDSIT | KNYGHPREN | KNYGHPRENF |
| CHLKN | GHPREN | GHPRENF | LYDALEAP | LYDALEAPA | LYDALEAPAD |
| YKYEE | HPRENF | GNRPTFV | MEGNRPTF | MEGNRPTFV | NKNYGHPREN |
| FYDSQ | ICHLKN | ICHLKNP | NYGHPREN | NYGHPRENF | RIYFYDSITN |
| HPREN | NRPTFV | QVVYKYE | QVVYKYEE | RIYFYDSIT | SMEGNRPTFV |
| PRENF | VVYKYE | VICHLKN | VICHLKNP | SMEGNRPTF | KIRIYFYDSV |
| HLKNP | VYKYEE | VVYKYEE | YFYDSITN | IRIYFYDSV | AYNGNDTEGL |
| WDPLL | YDSVTN | YFYDSIT | YFYDSVTN | AYNGNDTEG | GNDTEGLLKE |
| FYDSV | IKFNLY | YFYDSVT | YGHPRENF | GNDTEGLLK | NGNDTEGLLK |
| DPLLN | YDSITN | YghPren | RIYFYDSV | ndTEGLLKE | SAYNGNDTEG |
| MKKII | GNRPTF | IYFYDSV | DTEGLLKE | NGNDTEGLL | YNGNDTEGLL |
| RPTFV | YFYDSV | DTEGLLK | GNDTEGLL | Yngndtegl | VSPTRSAHFH |
| LDYVG | NDTEGL | GNDTEGL | NDTEGLLK | MWDPLLNEF | MWDPLLNEFP |
| YDSIT | MWDPLV | NDTEGLL | NGNDTEGL | WDPLLNEFP | WDPLLNEFPE |
| FYDSI | SRGPAG | NGNDTEG | YNGNDTEG | DPLLNEFPE | KIRIYFYDSI |
| KFNLY | PLLNEF | GSKSEAL | DPLLNEFP | IRIYFYDSI | DPLLNEFPET |
| DLDYV | WDPLLN | MWDPLLN | MWDPLLNE | PLLNEFPET | PLLNEFPETV |
| IKFNL | TEGLLK | DPLLNEF | WDPLLNEF | YDALEAPAD | YDALEAPADT |
| CGGGR | YFYDSI | PLLNEFP | PLLNEFPE | WGEFQIDGR | WGEFQIDGRS |
| MKKLL | IKFNIY | WDPLLNE | RIYFYDSI | DALEAPADT | GEFQIDGRSA |
| DSVTN | DPLLNE | IYFYDSI | YDALEAPA | GEFQIDGRS | DALEAPADTP |
| PLYSG | DALEAP | TEGLLKE | DALEAPAD | YIDKDGDTL | LEWGEFQIDG |
| SRGPA | MWDPLL | YDALEAP | GEFQIDGR | EFQIDGRSA | SPTRSAHFHP |
| KFNIY | SKSEAL | DALEAPA | ALEAPADT | SPTRSAHFH | EFQIDGRSAR |
| LYIPE | DTEGLL | ALEAPAD | EFQIDGRS | ALEAPADTP | EWGEFQIDGR |
| YFYDS | ALEAPA | EFQIDGR | FQIDGRSA | CCPHCPR HK | CCCPHCPRHK |
| YDSVT | EFQIDG | FQIDGRS | WGEFQIDG | FQIDGRSAR | FQIDGRSARG |
| NDTEG | GELITA | GEFQIDG | RTGELITA | DKDGDTLEW | DKDGDTLEWG |
| KNPEK | FQIDGR | TGELITA | KDGDTLEW | LEWGEFQID | KSYIDKDGDT |
| EFQID | AFNYIE | DGDTLEW | PTRSAHFH | EWGEFQIDG | KDGDTLEWGE |
| QQRLI | TGELIT | DKDGDTL | DGDTLEWG | DGDTLEWGE | PTRSAHFHPN |


| DTEGL | VSPTRS | KDGDTLE | SRTGELIT | PTRSAHFHP | DGDTLEWGEF |
| :---: | :---: | :---: | :---: | :---: | :---: |
| LKNPE | KDGDTL | RTGELIT | CCPHCPRH | KDGDTLEWG | DSRTGELITA |
| MKKLI | GEFQID | AFNYIES | EWGEFQID | SYIDKDGDT | MDSRTGELIT |
| CPRHK | GNDTEG | LVSPTRS | LVSPTRSA | SRTGELITA | TRSAHFHPNI |
| DSITN | RTGELI | VSPTRSA | VSPTRSAH | DSRTGELIT | DLVSPTRSAH |
| DPLVN | EGNRPT | WGEFQID | DLVSPTRS | CCCPHCPRH | FFDLVSPTRS |
| MWDPL | DGDTLE | CCPHCPR | TRSAHFHP | LVSPTRSAH | LVSPTRSAHF |
| AFNYI | QIDGRS | MEGNRPT | CCCPHCPR | TRSAHFHPN | FDLVSPTRSA |
| ALEAP | LGGAGG | TRSAHFH | LEAPADTP | DLVSPTRSA | PCCCPHCPRH |
| QIDGR | LEAPAD | SRTGELI | CPHCPRHK | VSPTRSAHF | QIDGRSARGG |
| FQIDG | SPTRSA | LEAPADT | DSRTGELI | FDLVSPTRS | LKIRIYFYDS |
| GPLYS | CCPHCP | QIDGRSA | QIDGRSAR | PCCCPHCPR | IDGRSARGGQ |
| PHCPR | CPHCPR | SPTRSAH | SPTRSAHF | MDSRTGELI | ASKVRRRLNF |
| HCPRH | DKDGDT | IDKDGDT | YIDKDGDT | QIDGRSARG | GNNSGQPSTV |
| PLLNE | IYFYDS | PHCPRHK | PCCCPHCP | KIRIYFYDS | SGQPSTVVDN |
| SPTRS | HCPRHK | CPHCPRH | IRIYFYDS | IDGRSARGG | NNSGQPSTVV |
| IEAAT | PHCPRH | CCCPHCP | IDGRSARG | GNNSGQPST | NSGQPSTVVD |
| FDSQT | IDGRSA | RIYFYDS | SGQPSTVV | SGQPSTVVD | PASKVRRRLN |
| SYIEK | SGQPST | PTRSAHF | NNSGQPST | NNSGQPSTV | TPASKVRRRL |
| CPHCP | PTRSAH | IDGRSAR | NSGQPSTV | NSGQPSTVV | GQPSTVVDNT |
| AVSNS | GQPSTV | GQPSTVV | GQPSTVVD | GQPSTVVDN |  |
| GNRPT | TRSAHF | SGQPSTV | ASKVRRRL | ASKVRRRLN |  |
|  | LVSPTR | NSGQPST | FDLVSPTR | PASKVRRRL |  |
|  | QPSTVV | DLVSPTR | QPSTVVDN |  |  |
|  |  |  |  |  |  |

## Table A17. Characteristic n-grams in disordered regions produced by combination of z -score, fractional difference, mole fractions and association rules

N -grams presented in the table characterize disordered regions by association rules, and have abs(zscore) $>2.58$ in disordered and abs(z-score) $<1.65$ in ordered regions, mole fractions $>1 \mathrm{E}-7$ and positive fractional difference in disordered regions. Table includes, for each $n$-gram length, (at most) first 100 n grams sorted according lift, confidence, and support, all in descending order.

| N -gram length |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| QQQ | HHHH | GGGGG | GGGGGG | PPPPPPP | GGGGGGGG | PEPEPEPEP | SSSSSSSSSS |
| PPR | SNAM | PPPPP | PPPPPP | EEEEEEE | PPPPPPPP | EPEPEPEPE | Eteeeeeeee |
| PPQ | GHMA | APAPA | TTTTTT | DDDDDDD | EEEEEEEE | EEEEEEEEE | PSPPPPPSPPP |
| SPS | GSHM | PSPPP | PEPEPE | PEPEPEP | PEPEPEPE | PKPAPKPAP | SPPPPSPPPP |
| TPP | PSPP | NnNnN | EPEPEP | EPEPEPE | EPEPEPEP | PAPKPAPKP | GGGGGGGGGG |
| PPK | QPQP | EEEED | GGGGGA | PKPAPKP | KPAPKPAP | KPAPKPAPK | PPSPPPSPPP |
| DDD | EPEP | PPAPP | PKPAPK | TTTTTTT | APKPAPKP | APKPAPKPA | VISTPASKVR |
| PPE | SSSS | SSTSS | KPAPKP | PPPSPPP | PAPKPAPK | DDDDDDDDD | RYGGGGGGGG |
| MPP | PQPQ | KKKKK | AGGGGG | KPAPKPA | QGAKSSSD | QQQQQQQQQ | HPNIQGAKSS |
| SES | QGPQ | DEEDE | PPSPPP | PAPKPAP | QQQQQQQQ | PPPPSPPPP | GQHISIRTFR |
| PDP | PQGP | PAPPP | APKPAP | APKPAPK | PPPSPPPP | PSPPPPSPP |  |
| HHH | QPQQ | KKSKK | PSPPPP | QQQQQQQ | GGGGGGGA | PPSPPPPSP |  |
| QKQ | PQPP | EEDDD | APAPAP | GGGGGGA | PSPPPPSP | PAPAPAPAP |  |
| ESE | PPQP | PPAAP | PAPAPA | PPSPPPP | PPSPPPPS | APAPAPAPA |  |
| PEE | PTPP | KKEKK | PPPPSP | PPPPSPP | SPPPPSPP | PSPPPSPPP |  |
|  | QPPQ | SKKKK | NNNNNN | MDSRTGE | AGGGGGGG | DSDSDSDSD |  |
|  | PQQQ | ESSSS | SRSRSR | PAPAPAP | APAPAPAP | PPSPPPSPP |  |
|  | PAPP | AAPPA | QGAKSS | GAKSSSD | SPPPSPPP | ADIVISTPA |  |
|  | QPPP | RRRGR | SPPPPS | QGAKSSS | PSPPPSPP | SKRPADIVI |  |
|  | MWDP | GGGDD | SDSDSD | RSRSRSR | GPEGPEGP | YGGGGGGGG |  |
|  | APAP | HHHHH | PQGPQG | SRSRSRS | GGGGGGSG | TGPQGPKGD |  |
|  | KPAP | PPPPQ | DSDSDS | GPQGPQG | EDEDEDED | GDGDGDGDG |  |
|  | PGPP | STTST | TGGGGG | PSPPPPS | AGGGGGSG | GGAGGGGGS |  |
|  | PPPA | DEDSD | GPQGPK | APAPAPA | GAGGGGGS | STNGIEPPR |  |
|  | PKPA | DDDKD | PQGPKG | SPPPPSP | ALRRRLER | RYGGGGGGG |  |
|  | PPPR | REEEE | SPPPSP | GPQGPKG | NPAPTSSP | GAGGGGGSG |  |
|  | PPGP | EKKKS | PSPPPS | GGGGAGG | LRRRLERG | DGDGDGDGD |  |
|  | APKP | GGSRS | PPPPPPS | DSDSDSD | GGGGGGGY | GAGAGGGAG |  |
|  | QGPK | SSSVD | TTAATT | PSPPPSP | DKGDKGDT | ASAYNGNDT |  |
|  | APPP | EAEED | SPPPPP | SDSDSDS | APTSSPTS | NDAAAEALN |  |
|  | PAPA | PSPEP | GGGGGY | MSKRPAD | DEDDEDDE |  |  |
|  | PSSP | NTERH | SSSTSS | DKGDKGD | EEQKQLTL |  |  |



# Table A18. Characteristic n-grams in ordered regions produced by combination of z-score, fractional difference, mole fractions and association rules 

N -grams presented in the table characterize ordered regions by association rules, and have abs(zscore) $>2.58$ in ordered and abs(z-score) $<1.65$ in disordered regions, mole fractions $>1 \mathrm{E}-7$ and positive fractional difference in ordered regions. Table includes, for each n-gram length, (at most) first 100 n grams sorted according lift, confidence, and support, all in descending order.

| N-gram length |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| WIC | IFII | YNVID | IKGGIP | QIKGGIP | NPVYATLK | THASNPVYA | FKEFMGAQRD |
| YCW | LLLW | IKGGI | NVIDDV | ACTHASN | VYATLKIR | VYATLKIRI | HYLKHFKEFM |
| WYW | VLAC | VGKRF | YNVIDD | ASNPVYA | KIRIYFYD | TENALLLYM | VIDDVDPHYL |
| CLW | IFLC | ATLKI | YMACTH | SNPVYAT | LKIRIYFY | IKGGIPTIF | MWARSLGPHN |
| WWF | CVLV | ACTHA | LYMACT | NPVYATL | TLKIRIYF | GPHNYLCGH | KVTGGQYASN |
| WWY | FVIF | GKRFC | QIKGGI | PVYATLK | YATLKIRI | PVQIKGGIP | QSNCKYGKPV |
| FCW | FIVF | LYMAC | SNPVYA | YNVIDDV | ENALLLYM | SLGPHNYLC | TVTGGQYASK |
| YFW | TMWA | MACTH | MACTHA | NHTENAL | TENALLLY | ENHTENALL | ERIQRLGRVG |
| cWV | VVIF | YMACT | NPVYAT | HRVGKRF | HTENALLL | WYNVIDDVD | VKSVYILGKI |
| CVW | YAIY | ALLLY | CTHASN | THRVGKR | VGKRFCVK | CGHLDLSPK | NHVVYNHQEA |
| WWI | LICL | KYENH | PVYATL | RVGKRFC | NALLLYMA | VATNIIENG | FDRINVRRLF |
| WIY | VAYY | GPHNY | NHTENA | WMDENIK | GIPTIFLC | VKSVYILGK | LSTAKHSVDI |
| WFW | WYVD | RFFDL | HTENAL | LGPHNYL | KGGIPTIF | WLVRDRRPY | KLKNHTNSVM |
| CYC | YLYF | PHNYL | THRVGK | YATLKIR | LKHFKEFM | VQIKGGIPT | TKYGKPIQIK |
| WMC | YFTF | IYFYD | HRVGKR | VYATLKI | VTGGQYAS | KVTGGQYAS | NLNSNLDRIF |
| VWI | IAWL | KIRIY | SDVTRG | IRIYFYD | CGHLDLSP | IWMDENIKT | ISDVTRGNGI |
| FWC | YIAI | TMWAR | GPHNYL | KIRIYFY | RSLGPHNY | GVISINNVI | FRCMLAIKYL |
| VWC | YYVY | RIYFY | RVGKRF | LKIRIYF | WYNVIDDV | VTRGNGITH | QIKGGIPSIV |
| ICW | LLWF | LLYMA | LGPHNY | TLKIRIY | NVIDDVDP | NTKYGKPVQ | SETIHSRSYT |
| CFI | LHYY | PLYFK | VGKRFC | ATLKIRI | GKPVQIKG | TVTGGQYAS | ALEAIRFYVS |
| WYF | CIAL | LYFKI | WMDENI | GKRFCVK | YGKPVQIK | HVVYNHQEA | IRDLISVIRA |
| YIC | CLAI | RFCVK | MDENIK | KRFCVKS | PVQIKGGI | SCMKIDHCV | NLPTSAGKSL |
| IWW | LTWL | KIWMD | YATLKI | VTGGQYA | YNVIDDVD | YGTPMDFGQ | RVNNYVVYNQ |
| FIW | DIIC | GKIWM | ATLKIR | YENHTEN | YLKHFKEF | IQIKGGIPT | LKRLRFKGTV |
| IYW | YIPI | LLLYM | RIYFYD | ENALLLY | DVDPHYLK | ERIQRLGRV | FRCMLAVKYL |
| ICC | FIYF | IWMDE | IRIYFY | HTENALL | HYLKHFKE | SVYVLGKIW | ERIVSILEWD |
| WHC | YLFV | NNVIR | KIRIYF | NALLLYM | QEAGKYEN | KLSTAKHSV | TYSPDTLGYD |
| VWY | YYEI | NYLCG | LKIRIY | TENALLL | VVYNHQEA | DRINVRRLF | KQLSFFWRPE |
| YLW | VWVV | HNYLC | NPLYFK | GKIWMDE | MWARSLGP | KGKLKLSTA | LGKTTVVAIF |
| YLC | IGYF | IFLCN | RFCVKS | VGKRFCV | GKYENHTE | FKGKLKLST | NLSRQLGKTT |
| LWW | CFAL | LCGHL | GKRFCV | LLYMACT | LVRDRRPY | PETVHGFRC | FLVRDRRPVD |
| WIF | IWEI | TIFLC | KRFCVK | ALLLYMA | NHTNSVMF | QELRVLAAL | TSLYPSIIRQ |
| WFV | YLCD | YLCGH | KYENHT | KIWMDEN | VHGFRCML | IPFRAPTVK | KSCSQGGIRG |
| WVY | CLGI | AQRDW | VTGGQY | LLLYMAC | HGFRCMLA | NHGFTHRGT | MDFGQVFNMF |
| WVW | VVYC | KNYFL | GKIWMD | IWMDENI | CISDVTRG | YFLTYPQCS | PHLHVLIQFE |
| WCL | LKCF | YLKHF | NALLLY | HNYLCGH | CVSDVTRG | VQIKGGIPS | DFGQVFNMFD |
| WWH | VMFF | HFKEF | ALLLYM | NYLCGHL | FGQVFNMF | KNDLRDRFQ | KQAIELLPDF |
| LWY | CFLT | NYFLT | ENALLL | NPLYFKI | IKTKNHTN | NVIRAVRFA | EGDSRTGKTM |
| CCY | CLYL | FMGAQ | YENHTE | YLCGHLD | ISINNVIR | APTVKILSK | NYIESHRDEY |
| QWW | GLCF | FLCNP | ENHTEN | LCGHLDL | DENIKTKN | YVVYNHQEA | AVGSGKSTGL |
| FLW | CVVC | FFDLV | TENALL | GGIPTIF | MDENIKTK | FASLYPSII | VGSGKSTGLP |
| CQW | TIWN | CRELH | KIWMDE | GIPTIFL | TRGNGITH | TIHSRSYTH |  |
| LWF | VCVC | YFLTY | LLYMAC | IPTIFLC | KSVYILGK | APKDFVLQF |  |
| GCW | IFYV | NHNLR | LLLYMA | LKHFKEF | GNGITHRV | CMLAVKYLQ |  |
| YWW | CYLN | FGQVF | IWMDEN | PTIFLCN | IKLKNHTN | HFIVATNII |  |
| WFM | DCII | LGKIW | HNYLCG | ARSLGPH | ATNIIENG | VLCNPGEGA |  |
| CCV | IYNM | LLLLV | NYLCGH | KEFMGAQ | NIIENGVT | VKNDLRDRF |  |
| VWW | YLFQ | WYNVI | LKHFKE | KGGIPTI | VATNIIEN | GEMTVAGKK |  |
| IWL | FFIF | VVYNH | LCGHLD | KHFKEFM | VLQFHNLN | IRAVRFATD |  |
| LCY | FWLV | FNHNL | PLYFKI | WARSLGP | LGVISINN | LPTSAGKSL |  |
| WWV | WAVL | VISIN | YLCGHL | HFKEFMG | PKVYSNDA | EGRGQDYHA |  |
| IWI | CYNL | AWYNV | CGHLDL | IKGGIPT | TNIIENGV | VNNYVVYNQ |  |
| CCF | VIYF | IQFEG | IFLCNP | CGHLDLS | VISINNVI | YFLTYPKCS |  |
| YCY | WLYN | LILLL | GAQRDW | FKEFMGA | GKVMCISD | ELRVLAALS |  |
| YYF | ILWL | QVFNM | KHFKEF | TGGQYAS | NIKLKNHT | ERIVSILEW |  |
| YYC | YLPY | VYNHQ | GGIPTI | IDDVDPH | VYNHQEAG | RKALGIHKC |  |
| WYV | AWGY | DPHYL | EFMGAQ | PHNYLCG | GQYASKEQ | YSIELAQDL |  |
| YCC | CFTV | FLRVF | GIPTIF | GHLDLSP | QYASKEQA | GDFLTSLIN |  |
| CFL | CYGL | HVLIQ | KGGIPT | GPHNYLC | KLKNHTNS | LNFQVWTTS |  |
| IFC | CLYA | LHVLI | PTIFLC | LGKIWMD | VTRGNGIT | NGLMVWCIE |  |
| VCF | GFFY | VLIQF | TIFLCN | RSLGPHN | IIENGVTL | INSLYGALG |  |
| FCL | VIMV | ICREL | GGQYAS | WYNVIDD | QRLGRVGR | VAFDMRGQQ |  |
| HLW | WALF | AGKYE | HFKEFM | KYGKPVQ | INNVIRAV | FLGLPFNIA |  |
| QWC | VYIW | VVVVV | KEFMGA | KPVQIKG | YVLGKIWM | ALGPHNYLS |  |
| CLI | FALC | RCMLA | RSLGPH | NVIDDVD | MCISDVTR | WLAIQPVIS |  |
| IVW | FLIC | GFRCM | FKEFMG | NYFLTYP | YDLIRDLI | TSAGKSLIQ |  |
| VWH | FRYY | HTNSV | GHLDLS | PVQIKGG | ENIKLKNH | KICRELHEN |  |
| WCT | LVCF | FRCML | DDVDPH | VIDDVDP | RRPYGTPM | SKEQALVKK |  |


| VWV | VYIY | LIIGL | YNHQEA | YGKPVQI | DENIKLKN | KICRELHED |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| FYY | IFQY | CGCSY | VQIKGG | RFCVKSV | SFFSLKDP | RALDNLLDY |  |
| WCM | LFYC | ILSLI | KPVQIK | FCVKSVY | YLSGHLDF | LYQSCHILQ |  |
| LIC | VYVC | LLVVL | GKPVQI | AWYNVID | FFSLKDPI | AIELLPDFL |  |
| WCQ | CILV | GCGKT | PHNYLC | YLKHFKE | FSLKDPIP | SKRYLYQDN |  |
| YIF | LWFL | DAWYN | SLGPHN | KNYFLTY | IENGVTLD | NIFLAMLVN |  |
| VWL | CVKI | IIILL | HLDLSP | LIQFEGK | KVCVDDFN | LSGIKGQIG |  |
| CLV | IDCV | YVLGK | LGKIWM | FGQVFNM | MDENIKLK | QYDKYNDVN |  |
| IYY | RVWL | GITHR | WYNVID | HLDLSPK | NIDLHYFS | CGMYASALT |  |
| NCY | VTCK | NDAWY | KNYFLT | DDVDPHY | SLKDPIPW | VNNYVVYNH |  |
| FVF | WIVA | VHGFR | YGKPVQ | DVDPHYL | KSVYVLGK | VLQFHNLNA |  |
| IWT | WLVV | FLLLL | VIDDVD | EFMGAQR | LRKALGIH | TSLYPSIIR |  |
| HVW | LFRM | HGFRC | NLDRIF | VDPHYLK | PIPWKLYY | IDLHYFSSS |  |
| TWC | CHIL | ILLVL | NYFLTY | DPHYLKH | CVIEYRQQ | LGKTTVVAI |  |
| VCL | IGWI | QIRFN | PVQIKG | MGAQRDW | DRYQVLRK | TKNHTNTVM |  |
| IFI | RFCI | VLLLV | YFLTYP | VVYNHQE | NLRKALGI | SRQLGKTTV |  |
| ICL | WNLV | VVLAL | FNHNLR | TMWARSL | NRFFDLVS | RQLGKTTVV |  |
| FYF | CVYV | ALGIH | AWYNVI | PHYLKHF | PIQIKGGI | DCSSAVEDL |  |
| YLF | IGDW | GDLIY | CVKSVY | HYLKHFK | IEYRQQVP | HNLNANLDR |  |
| LYY | YVFI | ILILL | YLKHFK | LDLSPKV | IQRLGRVG | YKKAQAFDE |  |
| CIH | HHII | GNIIG | FGQVFN | DLSPKVY | KPIQIKGG | ICFAGDDMC |  |
| WRW | CNIT | NYVVY | GQVFNM | QEAGKYE | RIQRLGRV | GVSEGIHPI |  |
| FWM | CNLC | ALVIL | LIQFEG | VLIQFEG | SCMKIDHC | EIHAELNAI |  |
| ICA | LAFW | GKVMC | IQFEGK | AGKYENH | YGKPIQIK | ASLYPSIIQ |  |
| VVW | LLVW | IILLL | VYNHQE | EAGKYEN | AKYENHTE | VVAIFLAHF |  |
| YNW | CSIY | LLLLG | DVDPHY | HVLIQFE | FAKFKGKL | TDIAGYAGC |  |
| FLY | ICII | AVLLV | VDPHYL | LHVLIQF | IRCNIDLH | VHGMDADAE |  |
| FIY | LFMI | ISLLL | FMGAQR | KNHTNSV | STAKHSVD | NLGVISINN |  |
| CNW | VLWY | LGVVA | NNVIRA | VRDRRPY | YGTPMDFG | VDLPCGCSY |  |
| LCL | YDIC | AVIRF | PHYLKH | ICRELHE | LKLKHWKE | NGVGPLCKG |  |
| ACF | IIYM | ILGAV | DPHYLK | DVTRGNG | DGSQFDSS | SQFDSSLTP |  |
| LCV | CAFI | IINIL | MGAQRD | THRGTHH | WTFPIRCN | FLVRDRRPV |  |

## Table A19. Characteristic n-grams in bordered regions produced by combination of fractional difference, mole fractions and association rules

N -grams presented in the table characterize bordered regions by association rules, and have mole fractions>1E-7 and positive fractional difference in bordered regions. Table includes n-grams sorted according lift, confidence, and support, all in descending order.

| N -gram length |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 5 | 6 | 7 | 8 | 9 | 10 |
| VYKYE | FYDSVT | YFYDSVT | YFYDSVTN | MEGNRPTFV | ASMEGNRPTF |
| WDPLV | NRPTFV | FYDSVTN | EGNRPTFV | SMEGNRPTF | SMEGNRPTFV |
| NRPTF | VVYKYE | GNRPTFV | MEGNRPTF | LYDALEAPA | LYDALEAPAD |
| CHLKN | VYKYEE | EGNRPTF | QVVYKYEE | NYGHPRENF | IRIYFYDSIT |
| YKYEE | GHPREN | VVYKYEE | YFYDSITN | IYFYDSITN | RIYFYDSITN |
| FYDSQ | FYDSIT | QVVYKYE | LYDALEAP | RIYFYDSIT | KNYGHPRENF |
| HPREN | CHLKNP | YGHPREN | YGHPRENF | GQVVYKYEE | NKNYGHPREN |
| PRENF | HPRENF | YFYDSIT | GQVVYKYE | KNYGHPREN | KIRIYFYDSV |
| HLKNP | ICHLKN | FYDSITN | NYGHPREN | IRIYFYDSV | AYNGNDTEGL |
| WDPLL | YDSVTN | GHPRENF | IYFYDSIT | AYNGNDTEG | YNGNDTEGLL |
| FYDSV | IKFNLY | ICHLKNP | VICHLKNP | NDTEGLLKE | GNDTEGLLKE |
| DPLLN | YDSITN | VICHLKN | RIYFYDSV | NGNDTEGLL | NGNDTEGLLK |
| MKKII | GNRPTF | IYFYDSV | DTEGLLKE | YNGNDTEGL | SAYNGNDTEG |
| RPTFV | YFYDSV | GNDTEGL | YNGNDTEG | GNDTEGLLK | VSPTRSAHFH |
| LDYVG | NDTEGL | NDTEGLL | NGNDTEGL | MWDPLLNEF | MWDPLLNEFP |
| YDSIT | MWDPLV | DTEGLLK | NDTEGLLK | WDPLLNEFP | WDPLLNEFPE |
| FYDSI | SRGPAG | NGNDTEG | GNDTEGLL | DPLLNEFPE | KIRIYFYDSI |
| KFNLY | PLLNEF | GSKSEAL | DPLLNEFP | IRIYFYDSI | DPLLNEFPET |
| DLDYV | WDPLLN | MWDPLLN | MWDPLLNE | PLLNEFPET | PLLNEFPETV |
| IKFNL | TEGLLK | DPLLNEF | WDPLLNEF | YDALEAPAD | YDALEAPADT |
| CGGGR | YFYDSI | PLLNEFP | PLLNEFPE | WGEFQIDGR | WGEFQIDGRS |
| MKKLL | IKFNIY | WDPLLNE | RIYFYDSI | DALEAPADT | GEFQIDGRSA |
| DSVTN | DPLLNE | IYFYDSI | YDALEAPA | GEFQIDGRS | DALEAPADTP |
| PLYSG | DALEAP | TEGLLKE | DALEAPAD | YIDKDGDTL | LEWGEFQIDG |
| SRGPA | MWDPLL | YDALEAP | GEFQIDGR | EFQIDGRSA | SPTRSAHFHP |
| KFNIY | SKSEAL | DALEAPA | ALEAPADT | SPTRSAHFH | EFQIDGRSAR |
| LYIPE | DTEGLL | ALEAPAD | EFQIDGRS | ALEAPADTP | EWGEFQIDGR |
| YFYDS | ALEAPA | EFQIDGR | FQIDGRSA | CCPHCPR ${ }^{\text {c }}$ | CCCPHCPRHK |
| YDSVT | EFQIDG | FQIDGRS | WGEFQIDG | FQIDGRSAR | FQIDGRSARG |
| NDTEG | GELITA | GEFQIDG | RTGELITA | DKDGDTLEW | DKDGDTLEWG |
| KNPEK | FQIDGR | TGELITA | KDGDTLEW | LEWGEFQID | KSYIDKDGDT |
| EFQID | AFNYIE | DGDTLEW | PTRSAHFH | EWGEFQIDG | KDGDTLEWGE |


| QQRLI | TGELIT | DKDDGDTL | DGDTLEWG | DGDTLEWGE | PTRSAHFHPN |
| :---: | :---: | :---: | :---: | :---: | :---: |
| DTEGL | VSPTRS | KDDGTLE | SRTGELIT | PTRSAHFHP | DGDTLEWGEF |
| LKNPE | KDGDTL | RTGELIT | CCPHCPRH | KDGDTLEWG | DSRTGELITA |
| MKKLI | GEFQID | AFNYIES | EWGEFQID | SYIDKDGDT | MDSRTGELIT |
| CPRHK | GNDTEG | LVSPTRS | LVSPTRSA | SRTGELITA | TRSAHFHPNI |
| DSITN | RTGELI | VSPTRSA | VSPTRSAH | DSRTGELIT | DLVSPTRSAH |
| DPLVN | EGNRPT | WGEFQID | DLVSPTRS | CCCPHCPRH | FFDLVSPTRS |
| MWDPL | DGDTLE | CCPHCPR | TRSAHFHP | LVSPTRSAH | LVSPTRSAHF |
| AFNYI | QIDGRS | MEGNRPT | CCCPHCPR | TRSAHFHPN | FDLVSPTRSA |
| ALEAP | LGGAGG | TRSAHFH | LEAPADTP | DLVSPTRSA | PCCCPHCPRH |
| QIDGR | LEAPAD | SRTGELI | CPHCPRHK | VSPTRSAHF | QIDGRSARGG |
| FQIDG | SPTRSA | LEAPADT | DSRTGELI | FDLVSPTRS | LKIRIYFYDS |
| GPLYS | CCPHCP | QIDGRSA | QIDGRSAR | PCCCPHCPR | IDGRSARGGQ |
| PHCPR | CPHCPR | SPTRSAH | SPTRSAHF | MDSRTGELI | GNNSGQPSTV |
| HCPRH | DKDGDT | IDKDGDT | YIDKDGDT | QIDGRSARG | ASKVRRRLNF |
| PLLNE | IYFYDS | PHCPRHK | PCCCPHCP | KIRIYFYDS | SGQPSTVVDN |
| SPTRS | HCPRHK | CPHCPRH | IRIYFYDS | IDGRSARGG | NNSGQPSTVV |
| IEAAT | PHCPRH | CCCPHCP | IDGRSARG | GNNSGQPST | NSGQPSTVVD |
| FDSQT | IDGRSA | RIYFYDS | SGQPSTVV | SGQPSTVVD | PASKVRRRLN |
| SYIEK | SGQPST | PTRSAHF | NSGQPSTV | NSGQPSTVV | TPASKVRRRL |
| CPHCP | PTRSAH | IDGRSAR | NNSGQPST | NNSGQPSTV | GQPSTVVDNT |
| AVSNS | GQPSTV | GQPSTVV | GQPSTVVD | GQPSTVVDN |  |
|  | TRSAHF | SGQPSTV | ASKVRRRL | ASKVRRRLN |  |
|  | LVSPTR | NSGQPST | FDLVSPTR | PASKVRRRL |  |
|  | QPSTVV | DLVSPTR | QPSTVVDN |  |  |
|  |  |  |  |  |  |

## Table A20. Left components of characteristic inverse noncomplementary repeats (material downloaded from NCBI) related to disordered regions

Table includes, for each repeat length, (at most) first 100 n -grams sorted according confidence, lift, and support, all in descending order.

| Repeat length |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| PPP | QQQE | GGGGG | GGGGGG | PPPSPPP | PPPSPPPP | PPPPSPPPP | SSSSSSSSSS |
| PPS | HHHH | PPPPP | PPPPPP | GGGGGGG | GGGGGGGG | sSSSSSSSS | PPPPPPPPPP |
| QQQ | PEPE | PSPPP | PSPPPP | PPPPPPP | SSSSSSSS | PPPPPPPPP | GGGGGGGGGG |
| SPP | EQEQ | PPPSP | TTTTTT | PKPAPKP | PPPPSPPP | GGGGGGGGG | EEEEEEEEEE |
| PGP | GEGP | EEGEG | PPPPSP | DDDDDDD | PPPPPPPP | PPPPLPPPP | DDDDDDDDD |
| PSP | KPAP | PPPPS | PPSPPP | TTTTTTT | EEEEEEEE | QQQQQQQQQ | TTTTTTTTTT |
| PAP | QPQQ | PKPAP | SPPPPS | EEEEEEE | DDDDDDD | EPEPEPEPE | KPAPKPAPKP |
| PKP | DGKP | SPPPP | PPPSPP | PPSPPPP | TTTTTTTT | DDDDDDDD | QQQQQQQQQQ |
| PQP | QPPP | APAPA | PPPPLP | PAPAPAP | QQQQQQQQ | PSPPPSPPP | GAGGGAGAGG |
| GPR | APRP | SSTSS | TTAATT | PSPPPPS | PPPPLPPP | EEEEEEEEE | PPPSPPPPSP |
| QPQ | QQPQ | NNNNN | nNnNNN | PTPSPTP | SPPPSPPP | TTTTTTTTT | TTAATTAATT |
| PEP | QQAQ | PPAPP | PLPPPP | EPEPEPE | ATTAATTA | PPPPTPPPP | SPPPSPPPSP |
| RPP | EQEK | KKKKK | EDEEDE | QQQQQQQ | PSPPPPSP | PEPEPEPEP | PPSPPPPSPP |
| RPG | PPQP | PSPGP | AGGGGG | PPPPSPP | APKPAPKP | RSRSRSRSR | NNNNNNNNNN |
| GPP | PQPQ | PAPKP | GGGGGA | RSRSRSR | NNNNNNNN | SRSRSRSRS | PPPSPPPPLP |
| PTP | PQQQ | PSPSP | PRPPRP | PSPGPSP | cSSSSSSS | PPPSPPPSP | PEPEPEPEPE |
| PRP | QQVP | KKSKK | PKPAPK | SPPPPSP | PPPTPPPP | PPPPAPPPP | PPVVPPVVPP |
| SSS | GPGP | TETTN | DEEEED | PEPEPEP | EEAEEAEE | GAGGAGAGG | PSPPPPSPPP |
| EEE | QEQE | PEPGP | PEPEPE | SRSRSRS | EEDEEDEE | DSDSDSDSD | PKPAPKPAPK |
| PPA | PSKP | KKEKK | QAQQAQ | PTPPPTP | DEEYYEED | NnNnNNNNN | DGGDGGDGGD |
| PPG | ISPQ | ннннн | KРАРКР | NnNnNnN | TTAATTTA | SPPPSPPPP | GGAGGAGAGG |
| SPS | RPPP | PPEPP | KKKKKK | GEDEGED | TTAATTAA | GAGGAGGAG | GGGGGGGGGA |
| PPR | SPSR | PSKSP | HHHHHH | DEDEDED | PAPAAPAP | PAPAPAPAP | PSPPPSPPPP |
| EQE | KPEE | PTPSP | SPSPPP | PAPKPAP | SPSPPPPS | GGAGAGGAG | EEAEEEEAEE |
| EDD | PPSS | PSPTP | EDEGED | TTAATTA | PEPEPEPE | GEDEGEDEG | PAPTTPAPTT |
| EPE | AKRR | VPEPA | SGSGSG | APAPAPA | GGSGGSGG | APKPAPKPA | EPEPEPEPEP |
| DEG | SDSE | TTTTP | EEEAEE | DSDSDSD | PPPLPPPP | DEGEDEGED | GGAGAGGGAG |
| DDD | APSP | PGPPG | PPSSPP | GPPGPPG | GPSPGPSP | PPSPPPPSP | SPPPPSPPPP |
| APP | AQTQ | PQQQP | EEAEEE | PSPPPSP | PPSPPPPS | AEKAKAKEA | EDEDEEDEDE |
| DEE | GPQG | PPPLP | PAPKPA | PPPTPPP | SPPPPSPS | KPAPKPAPK | GPPGPPGPPG |
| QEQ | KPPP | GPPGP | csssss | SDSDSDS | GGGGGGAG | CSSSSSSSS | PLPPPPSPPP |
| SSP | TTET | PPPAP | PTPSPT | DEGEDEG | PPPPAPPP | DEDDEDDED | PPPTPPPTPP |
| RRR | DGED | PTPKP | GGDGGD | KPAPKPA | ATTTAATT | PTPSPTPTP | APAPAAPAPA |
| EED | PAQQ | LPPPP | PTTTTT | DDEDEDD | GAGAGGAG | PTPTPSPTP | GGRGGGGRGG |
| DDE | PQPP | PKPTP | PPPSPS | GGAGGAG | APTGGTPA | PPPPNPPPP | PAPKPAPKPA |


| PSS | TPSP | PTPPP | EPEPEP | TTAATTT | DDEDDEDD | APAPAPAPA | PPPPSPPPSP |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| KPK | VEED | PKRKP | PPAAPP | PTPTPTP | HHHHHHHH | DEDEDEDED | PPPSPPPPPP |
| PPT | RSPS | PPDPP | PSPPSP | GGGGGAG | PAPTTPAP | GDGGDGGDG | QAQQAQAQQA |
| APA | TDGK | QPAPQ | EEQEQE | AGGGGGA | PPTPPTPP | PKPAPKPAP | APKPAPKPAP |
| TPP | VPQQ | REQER | RSRSRS | SPSPSPS | SALSSLAS | GGGGGGGAG | EEEAEEAEEE |
| RPR | GNMN | KRPRK | AAPPAA | TTTPTTT | TSALSSLA | PEEVVVEEP | PPPPPSPPPP |
| GEG | QQPP | SHTHS | GEEGEG | AGGGGGG | DGEDEGED | SSSSGSSSS | PPPPSPPPPS |
| RRS | RMAE | PPPTP | SSPPSS | GRGRGRG | SSASSASS | PPPSPPPPS | QAQAQQAQAQ |
| EDE | RPGE | CSSSS | PAPAPA | PPPAPPP | SSSSSSSC | TTTTPTTTT | Scssssssss |
| SRS | SPTS | QAQQA | TPTTTT | CSSSSSS | AGGGGGGA | PSPTPSPTP | SSSASSASSS |
| RSR | AEGP | RSPSK | PPPTPP | GEGEGEG | EPEPEPEP | EDEDEDEDE | GGDGGDGGDG |
| DED | ERKR | PGPEP | SSGGSS | DDDGDDD | GDGGDGGD | EEAEEEAEE | GGGGYYGGGG |
| PDP | NGPQ | QGNGQ | AASAAS | PAPEPAP | RKSKKSKR | GEGEGEGEG | NPPPPSPPPP |
| RKR | NNGP | KEAER | QPQQPQ | PSPTPSP | SDSDDSDS | PAPKPAPKP | PAAPAAPAAP |
| RQR | PGMG | QQTQQ | SRSRSR | APASAPA | PPAPPAPP | QAQQAQQAQ | PSPTPPPTPS |
| KKK | PSPT | PPSPS | AEKKEA | APKPAPK | PPTPSPTP | AGSTATSGA | RSRSRSRSRS |
| QRQ | QAID | EAEEE | PPRRPP | ATTAATT | GRGRGRGR | EKQASAQKE | SRSRSRSRSR |
| PMP | SHEA | EEEDE | PSSSSP | DGGDGGD | PAPKPAPK | PGPPGPPGP | AAPAAAAPAA |
| EGE | SQGG | PVPKP | RGGGGG | EDDEDDE | PKPAPKPA | PPAPAPAPP | AQQAQAQQAQ |
| GGG | TGPD | QEREQ | SKKKKS | TETKTET | PPGAAGPP | AGGGGGGGA | DEDDEDDEDD |
| KRK | DEAP | GDGGD | SKRRKS | ннннннн | PPPAPPPP | AKKAPAKKA | DKEDKKDEKD |
| SES | GDCG | PQSQP | TTTTPT | KKKKKKK | GGRGGRGG | EEEAEEAEE | EDDEDDEDDE |
| PHP | KGGD | PTPIP | AAAAQG | PAPPPAP | KKKKKKKK | EETSESTEE | EEDEEDEEEE |
| QAQ | RAQA | EEEED | GGGGGS | LPPAPPL | PPPPPSPP | PTPKPTPKP | EPKPEEPKPE |
| QKQ | RSPP | QQHQQ | DAKKAD | APATAPA | SPSPGPSP | PTPTPTPTP | GAGGAGGAGG |
| SDS | QQQR | AEKAK | PQQQQP | KEEAEEK | AGGGGGGG | APAAPAAPA | GGAGAGGAGG |
| EEK | SPKP | AQQAQ | PTDDTP | PVPKPAP | KKSKKSKK | GGGGSGGGG | ннннннннннн |
| PVP | APPR | PEEPK | SDDDDS | EEDEDEE | PPSPPPSP | GRGRGRGRG | KKKAEEAKKK |
| EEA | AQPQ | DDDED | SSRRSS | EEDEEDE | RRRSSRRR | HHHHHHHHH | PTPTPPTPTP |
| EAE | EAER | DEDDE | AKAKEA | SSSTSSS | RSRSRSRS | PPPPVPPPP | PVRRRRRRVP |
| GPG | EQTP | EDEEE | EEPKPE | EDEEEDE | APAKKAPA | RRSPSPSRR | RRSPSPSRRS |
| ESE | KKQP | EPQPE | EESSEE | NKKSKKN | DGGDGGDG | SDSDSDSDS | SSSSAASSSS |
| ERE | QRQQ | GCQCG | GSGGGG | PKPTPKP | DSSSSSSD | SESESESES | TPSPTPSPTP |
| DSD | KDEK | PDPGP | KPAPAP | PSPSPSP | EDEDEDEE | ERERERERE | AGGGGGGGGA |
| SQS | KEDK | QQQVP | PPPPAP | RGRGRGR | EDEDEEDE | GDGDGDGDG | Eedeeeedee |
| PNP | PGSP | RRRSS | PPQQPP | SSSASSS | EDEGEDEG | GGGGAGGGG | EEEEDDEEEE |
| QSQ | PPDP | TPAAP | STSSTS | APPAPPA | EEDEDEDE | PKPAPKPKP | EEEEIIEEEE |
| PGA | PTSP | APPAP | APKPAP | APPPPPA | GEDEGEDE | PPPSPPPPP | ELDALLADLE |
| APS | QEAK | APVPV | EEDEDE | Eeedeee | GGGGGGGA | PPPTPSPTP | GEEEEEEEEG |
| QTQ | QSSS | ELTGP | PTTPAP | ERERERE | GRSSSSRG | SPPPSPPPS | GRGNGNGRGN |
| AAP | TSKS | GGKEA | SGGGGG | PPSSPPS | QAQAQQAQ | EEEEDEEEE | GSSGSSGSSG |
| RER | DSES | PAAPP | ANPPNA | SSSPSSS | QRKTTKRQ | ERREKERRE | KLKKYYKKLK |
| EKE | EKAY | PAPTP | DGGDGG | PDPLPDP | SSGSSGSS | ESESESESE | PPPAPPAPPP |
|  | NEAK | PPPQP | EPKPEE | PEPTPEP | TTTAATTT | KGKGKGKGK | PPPPPPSPPP |
|  | PPVP | PTGGT | ESDDSE | RLEEELR | DDEEEEDD | PEPSPEPSP | PPPPSPPPPP |
|  | QQPA | QAARE | KPKKPK | TTAPATT | DDSDDSDD | PPPSPSPPP | PPPSPPSPPP |
|  | QTQA | QEEQE | SDEEDS | eemaeee | GGEGGEGG | QAAQAQAAQ | PPSSPPSSPP |
|  | SFDD | SAGGG | SGTTGS | GGNSNGG | GGGAGAGG | SGSGSGSGS | PTTTTTTTTP |
|  | SPTR | TTTPT | SRRRRS | GRGGGRG | GGTGGTGG | SPTPTPSPT | RRRRRRRRRRR |
|  | SQRS | AASSG | TSTTST | GSGSGSG | KKDKKDKK | cSSSSSSSS | SESESESESE |
|  | TEPE | DPTTS | TTEETT | KKAPAKK | scssssss | DDDDGDDDD | SGGNGGNGGS |
|  | EDSD | EDDED | DEEDEE | PEPAPEP | SDSSSSDS | EKEKEKEKE | AAAAPPAAAA |
|  | EESE | EETSE | DEGGED | SGSGSGS | SRSRSRSR | KEKEKEKEK | AAAASAAAAK |
|  | GQQA | GGGDG | DSSSSD | AAPAPAA | TTTPTTTT | PKPKPEPEP | ARAARAADAA |
|  | KPKR | MEREM | EPEEPE | AKPQPKA | TTTTPTTT | PSPPPLPPP | ASSASSASSA |
|  | NQNA | PAPAA | GGGGRG | APAYAPA | EDEEDEDE | PTPKPKPTP | DDDDDEDDD |
|  | PHPH | PAPEP | KKGGKK | DDDNDDD | EDEEEEDE | PTPSPTPSP | DDEDDDDEDD |
|  | PSEP | PQRQP | REAAER | EDEGEDE | EEEAEEAE | RERERERER | GGNGGGGNGG |
|  | PSRS | QRSRQ | SAASAA | GDDGDDG | EPEGDDDG | SAGAGAGAS | NNKKNNKKNN |
|  | QQEQ | SATSS | AEEEEA | GGGQGGG | GDDDDDDG | SPSPSPSPS | PAPAPPAPAP |
|  | RASQ | STTTT | DSDSDS | KEKEKEK | KEDKKDEK | ADADADADA | PPSPPPPPSPS |
|  | RPAR | DDEEE | EEKPEE | KGDPDGK | QPPQQPPQ | GSGSGSGSG | PTPPPTPSPT |
|  | GKEE | KPEEP | EREERE | SSRRRSS | RGGGGGGR | PPSPPPSPP | QEEQEEQEEQ |
|  | PPEE | PGPSP | QPPPPQ | SSSDSSS | RGRGRGRG | PTPPTPPTP | SAASAAASAA |
|  | PTDD | SPSSS | RRQQRR | TTTATTT | SSPSSPSS | SDAKRKADS | SAASAASAAS |

# Table A21. Left components of characteristic inverse noncomplementary repeats (material downloaded from NCBI) related to ordered regions 

Table includes, for each repeat length, (at most) first 100 n-grams sorted according confidence, lift, and support, all in descending order.

| Repeat length |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| YLL | YLLY | VLLLV | NYVVYN | GLGAGLG | INLKKLNI | GLSVPVSLG | VVLALLALVV |
| FYF | CVVC | VVVVV | NLKKLN | EVIRIVE | GAGLLGAG | ELGNKNGLE | TGVtttivg |
| YLY | YVVY | GVNVG | KLYYLK | ALAAALA | GKRHHRKG | vcvevcvev | LSILLLLISL |
| LWL | LCCL | ILSLI | VSVVSV | ASVDVSA | AGAGGAGA | GAAAGAAAG | AVGLLLLGVA |
| YFY | FYYF | IILII | ERYYRE | IERVREI | GAGAAGAG | GVGFGVGFG | CGFGCGFGCG |
| YVY | IINN | LGVGL | NKYYKN | GFGAGFG | KNLGGLNK | GLGAGLGAG | GCCGCCGCCG |
| IFI | YFFY | VLGLV | LLRRLL | NGDWDGN | YNLDDLNY | NELLSLLEN | IIIILLIIII |
| IIL | CIIC | ILFLI | GLAALG | RALDLAR | YQLLLLQY | IVKDRDKVI | LALLLLLLAL |
| YIY | VPVL | LIKIL | VVVVVV | VAGSGAV | DVKTTKVD | AAWAAAWAA | STGGFFGGTS |
| FIF | MFFM | LIPIL | IILLII | LILKLIL | GGGLLGGL | cVCVCVCVC | VVVVVVVVVV |
| VWV | LLSL | LIDIL | IINNII | LLVRVLL | ILLIILLI | GVGVGVGVG | YNNYNNYNNY |
| FVF | CCCC | GLGAG | DELLED | vcvevcv | LSELLESL | IGILLLIGI | AgAGAAGAGA |
| IYI | WAAW | IIDII | GTLLTG | VNRLRNV | SLFDDFLS | DGDLRLDGD | LLLLLLLLLL |
| CVC | LNIS | LLYLL | LLGGLL | AALALAA | AAYAAYAA | GYLSFSLYG | GAAGAAGAAG |
| LII | IMMI | IIFII | DVIIVD | AATQTAA | EILSSLIE | LLVLFLVLL | GAGAGGAGAG |
| ILI | CYYC | IIGII | FVLLVF | AVDEDVA | EVLEELVE | FIENFNEIF |  |
| CLC | LVAL | VLTLV | IIKKII | HMSDSMH | SAFGGFAS | GSAFGFASG |  |
| LLF | WEEW | TLQLT | IYKKYI | PIQVQIP | VNVGGVNV | HAILTLIAH |  |
| III | AIGG | IINII | ALVVLA | AAGKGAA | IIIIIIII | LSDVGVDSL |  |
| FLL | ALVA | DINID | ILLLLI | GGALAGG | NATAATAN | AAGLVLGAA |  |
| LIL | LWWL | ILALI | ISFFSI | IKNKNKI | YDKAAKDY | ALDDADDLA |  |
| GIV | SVRV | LIIIL | LVLLVL | AIEYEIA | GKCAACKG | ALNTFTNLA |  |
| LFL | VAGL | LVIVL | TIDDIT | KATVTAK | AgAttaga | EALLELLAE |  |
| VYV | VCCV | NITIN | VLAALV | LDKIKDL | GGFGGFGG | EGSRIRSGE |  |
| FLF | VGSV | IIAII | IKEEKI | LFSSSFL | GSLIILSG | FPKTVTKPF |  |
| ICI | QWWQ | VVIVv | LFIIFL | RYLVLYR | ILILKLIL | GAGFGAGFG |  |
| LCL | ICCI | LINIL | LFLLFL | WGCSCGW | KFGAAGFK | GVIPDPIVG |  |
| VCV | AVGL | LVPVL | VLDDLV | CEVRVEC | LVKEEKVL | IEKFKFKEI |  |
| LLV | FMMF | YPDPY | KKLLKK | ELLELLE | NNYNNYNN | LRLRLRLRL |  |
| IIN | PLDI | VLFLV | LIIIIL | MMDYDMM | RIEGGEIR | LSNVGVNSL |  |
| LYL | CHHC | FLLLF | LLVVLL | GVGFGVG | TGIAAIGT | LTASSSATL |  |
| IIG | VVDG | IISII | GAGFGA | IVLLLVI | TIAIIAIT | NYNNYNNYN |  |
| VVL | YTGL | IAFAI | IKNNKI | LKSASKL | WIEKKEIW | PALLNLLAP |  |
| LIT | ALLL | LIAIL | LGAAGL | NATITAN | YNAIIANY | SIESASEIS |  |
| ILL | ILGI | FLALF | NINNIN | VAGVGAV | AETTTTEA | STSTETSTS |  |
| YAY | ASFV | VSFSV | RIGGIR | ADEIEDA | AIYKKYIA | TTAGTGATT |  |
| VLL | TITI | LVTVL | VEDDEV | ELFNFLE | ALAGGALA | TVGSYSGVT |  |
| YTY | TLTV | NVLVN | VLLLLV | IIIIIII | ATAVVATA | YISISISIY |  |
| YQY | TYLR | FILIF | VVAAVV | IVFTFVI | DDIDDIDD | AAGGIGGAA |  |
| VFV | IGNG | ILVLI | AKNNKA | KLAVALK | FEKVVKEF | CGCCMCCGC |  |
| VIV | LVVV | LYTYL | DIGGID | KTIDITK | GGSIISGG | CGFGCGFGC |  |
| YRY | EPDY | ILTLI | ILGGLI | RAKLKAR | GILSSLIG | DSALHLASD |  |
| YKY | FSGI | IRGRI | KAVVAK | YGGAGGY | IAAVVAAI | GAAGAGAAG |  |
| VIG | IIKN | IVGVI | LLIILL | AAYEYAA | IYKNNKYI | GAGLGLGAG |  |
| LVL | VDGT | LLCLL | YYGGYY | AFAGAFA | KIKIIKIK | GTSVWVSTG |  |
| YGY | GCGL | vgYgV | GGIIGG | cVCVCVc | KNLTTLNK | GVGFGFGVG |  |
| IAI | LALP | IVAVI | LAVVAL | GGYPYGG | LIIIIIIL | IIIIIIIII |  |
| IGI | YRLF | LKKLN | LFFFFL | GKMLMKG | LLPLLPLL | IMKFLFKMI |  |
| LLI | IILI | AIIIA | LGDDGL | GTGSGTG | LPGLLGPL | KNRNSNRNK |  |
| IVV | IINK | IVDVI | LKVVKL | ILITILI | LSALLASL | LAALSLAAL |  |
| IVI | LLLK | LFAFL | AVRRVA | LLLFLLL | NEYNNYEN | NIIKEKIIN |  |
| FFF | GLLL | IVNVI | FFFFFF | LQFIFQL | NKRYYRKN | NNIINIINN |  |
| FAF | GNIA | LFSFL | FILLIF | NIFEFIN | NNSVVSNN | SLWGNGWLS |  |
| IDV | GVVS | LITIL | ITGGTI | QTEVETQ | RRWRRWRR | TIAGFGAIT |  |
| VLV | KLNI | VIIIV | LLLGVA | YIESEIY | SQNIINQS | VQGLGLGQV |  |
| LIN | LLLV | AIRIA | tLAALT | AGAFAGA | TEVKKVET | VTDTVTDTV |  |
| LLL | LTGN | GFGVG | VALLAV | AGFGAGF | TLLTtLLT | AAAAHAAAA |  |
| NII | AILS | LIYIL | VVGGVV | DGVEVGD | VVMVVmVV | AAARYRAAA |  |
| IHI | AITG | LVYVL | DFIIFD | DVDYDVD | vVVVVVVV | AADAAAYAA |  |
| YNY | INSN | LIFIL | DLTTLD | LQAAAQL | AALLLLAA | AAGLFLGAA |  |
| AIV | LGLL | FLGLF | KIAAIK | VARHRAV | AATIITAA | AINLVLNIA |  |
| IIK | LTTG | ITATI | KKSLIR | AAGNGAA | AIRAARIA | CGCCTCCGC |  |
| ILT | NGTL | LFLFL | LFKKFL | AALYLAA | ALAAAALA | DELVLVLED |  |
| IID | VLKT | LIGIL | NAIIAN | AAVVVAA | ELFGGFLE | DIDDIDDID |  |
| FRF | GTVN | VIDIV | NIKKIN | ARVKVRA | EnYVVYne | DLKGTGKLD |  |
| IMI | LVPV | VISIV | NLEELN | AVAGAVA | FTFAAFTF | EVPFEFPVE |  |
| ILA | SQTV | VCVCV | RLHHLR | DDGMGDD | GTLMMLTG | FGCGFGCGF |  |
| VMV | VGVG | FTYTF | TKDDKT | GGLTLGG | IIAIIAII | FLSLCLSLF |  |
| FTF | VLAL | GLILG | VGLGVG | GIGAGIG | IILLLLII | GAGFGFGAG |  |
| INL | DIYS | IDLDI | VLGGLV | GITETIG | ILILLILI | GAGLGAGLG |  |
| FPF | GTLN | IGVGI | AGFGAG | HFANAFH | ILLLLLLI | GFDLTLDFG |  |


| TLI | IIDK | INYNI | AIFFIA | INLKKKLN | ILVVVVLI | GGSAGASGG |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| FNF | ISSV | LTMTL | GKIIKG | LTTITTL | INNIINNI | HYHYHYHYH |  |
| IDI | ITIT | ISISI | GVAAVG | SIYRYIS | ITRFFRTI | IDEIEIEDI |  |
| LVV | LLGL | IVKVI | ILIILI | TGVHVGT | IWNNNNWI | IGAATAAGI |  |
| LNI | NNID | NIVIN | ISIISI | VGAKAGV | LAALLAAL | IILIIILII |  |
| LFG | NNIL | IVLVI | IVLLVI | VLLSLLV | LDRYYRDL | ISDVYVDSI |  |
| CGC | VGLP | VIGIV | KDVVDK | VTEEETV | LKQYYQKL | IYIYIYIYI |  |
| IIT | YFGN | VIVIV | LKLLKL | AAAWAAA | LLRLLRLL | KEVFEFVEK |  |
| INI | FAVG | VVNVV | NVFFVN | AGGWGGA | LNTLLTNL | KINNYNNIK |  |
| VVV | IDLV | DLALD | NYPPYN | ANFTFNA | LRRRRRRL | LAVGAGVAL |  |
| ITI | IIGN | FLSLF | VIVVIV | AVFGIVA | LTNIINTL | LCPCLCPCL |  |
| VIA | LINN | GVGFG | AIGGIA | AVQYQVA | NLEIIELN | LFDEMEDFL |  |
| GIL | LLLD | HGFGH | AWAAWA | DGVLVGD | NQLLLLQN | LKTKNKTKL |  |
| INN | NGTT | IFEFI | DNVVND | DVSGSVD | NVNAANVN | LLLLTLLLL |  |
| IVA | NINI | IGIGI | ETVVTE | ELLPLLE | QGELLEGQ | LLPLLLPLL |  |
| ILN | NNIK | ILQLI | EVNNVE | GADVDAG | SDAIIADS | MLLLSLLLM |  |
| YDY | YTTT | ILYLI | IISSII | GNFAFNG | SLVGGVLS | NDLMSMLDN |  |
| IVN | ALVL | ISYSI | LDTTDL | IAGGGAI | TEAIIAET | NLKKLKKLN |  |
| LVT | AVVL | IVIVI | LGAGAG | LIIIIIL | TGVAAVGT | PCLCPCLCP |  |
| YSY | GLVA | IVVVI | LLTTLL | MFISIFM | TGYTTYGT | RLCCFCCLR |  |
| IVG | LGQI | LLWLL | LTLLTL | PFVNVFP | TLSAASLT | SIDDEDDIS |  |
| VTI | LLKG | VGNGV | LVGGVL | RGIEIGR | TVSGGSVT | SLALMLALS |  |
| ITV | LPGK | VILIV | NCNNCN | RGSFSGR | VGLAALGV | SRLYRYLRS |  |
| YEY | LVLA | VLKLV | NLTTLN | RTGVGTR | VLLLLLLV | VDDVVVDDV |  |
| FKF | TDLY | YAVAY | RNHHNR | RVACAVR | VNTAATNV | VIVIVIVIV |  |
| LML | TGDG | FGAGF | RVPVTE | TELFLET | VQLEELQV | VLPLCLPLV |  |
| TIV | VGVV | KYLYK | VLSSLV | TTGHGTT | YARLLRAY | VVVVVVVVV |  |
| NIL | VSGT | LFIFL | VTNNTV | VVVAVVV | YGAVVAGY | YAYDADYAY |  |
| ILG | WALN | VLYLV | VVSSVV | YLLNLLY | YYNYYNYY | YRPDADPRY |  |

## Table A22. Left components of characteristic inverse noncomplementary repeats (material downloaded from NCBI) related to borderline regions

Table includes, for each repeat length, (at most) first 100 n -grams sorted according confidence, lift, and support, all in descending order.

| Repeat length |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| PVRV | GMWMG | RMSSMR | TPDFDPT | SPIGGIPS | LTPPTPPTL | RLRGLLGRLR |
| GDIA | MKWKM | QKIIKQ | TSAVAST | DNLEELND | AGDKIKDGA | DEVVEEVVED |
| AENR | MAWAM | GPWWPG | NAWGWAN | FSLEELSF | KSFKEKFSK | PVVPVVPVVP |
| MWWM | QMAMQ | GDKKDG | RVAQAVR | IELPPLEI | VKTSPSTKV |  |
| RDES | MYYYM | MLEELM | DIAEAID | SAGGGGAS | AAAAEAAAA |  |
|  | ALAGS | MVTTVM | EITETIE | HFHHHHFH | AAAAGAAAA |  |
|  | HEFEH | RGEEGR | KFLDLFK | IEDEEDEI | LKLSDSLKL |  |
|  | MFQFM | EFFFFE | PRVFVRP | TDDRRDDT | LQMKLKMQL |  |
|  | MYMYM | ERGGRE | KLTITLK | AKLTTLKA | ММКМТМКММ |  |
|  | PFWFP | ETIITE | REEFEER | DKGAAGKD | PAAAAAAAP |  |
|  | QDFDQ | PRDDRP | RFQVQFR | EFKKKKKFE | RTQVKVQTR |  |
|  | QVFVQ | SIQQIS | RSQPQSR | NKEFFEKN | VAEAEAEAV |  |
|  | SNLKK | AAAAPA | ADLMLDA | SLVEEVLS | VLEELEELV |  |
|  | WDWDW | ADMMDA | AEREREA | YEVRRVEY | VPVVPVVPV |  |
|  | WEEEW | ARTTRA | EANHNAE | SSAAAASS | Yadamaday |  |
|  | צммму | DHDDHD | LSGSGSL | KELEELEK | DIDIDIDID |  |
|  | EYFYE | GNEENG | AEATAEA | AAASSAAA | RRRRWRRRR |  |
|  | ELRAL | ITEETI | AVARAVA | MKMEEMKM | ADAADAADA |  |
|  | ECECE | NHDDHN | EEIWIEE | AALAALAA |  |  |
|  | QRFRQ | RIEEIR | ELIAILE | AIKLLKIA |  |  |
|  | EYHYE | RRIIRR | GAMLMAG | LDLDDLDL |  |  |
|  | DWNWD | RDIIDR | GKIDIKG | LTQGGQTL |  |  |
|  | MNYNM | VVPPVV | HEEREEH |  |  |  |
|  | MVTVM | KKYYKK | INGNGNI |  |  |  |
|  | RYDYR | LNKKNL | LGNFNGL |  |  |  |



## Table A23. Left components of characteristic inverse noncomplementary repeats (material downloaded from DisProt) related to disordered regions

Table includes, for each repeat length, (at most) first 100 n-grams sorted according confidence, lift, and support, all in descending order.

| Repeat length |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| YTP | EQQE | PSYSP | SPSYSP | PAPAPAP | PQQPQQPF | PAPAPAPAP | APAPAPAPAP |
| PSY | DSDS | EKSEV | VPKKPV | PQQPQQP | QQPFPQQP | GGGGQGGGG | PQQPQQPFPQ |
| EQQ | NDDK | KKPVP | EEEEKE | QPFPQQP | QPQQPFPQ | PFPQQPPQP | QPFPQQPQQP |
| YSP | VPVP | PKKPV | EVQQVE | QPQQPFP | KPKAAKPK | QQPQQPFPQ | DDDDDDDDDD |
| QQE | PPPG | PVPKK | GVVVVG | GGGWGGG | AGAAAAGA | DEDEDEDED | GGGGGGGGGG |
| FSF | QQPF | QPQQP | QPQQPF | KHKDKHK | AAAAAAAA | QPQLPFPQQ |  |
| GWG | QQVE | EVEVE | PPPPPPG | SPSYSPS | DDDDDDDD | TPTPTPTPT |  |
| EQK | GPPP | QQPFP | APAAPA | DEDEDED | EEEEEEEE | APAPAPAPA |  |
| KLK | PEVP | PEEEE | EDEEDE | EAEAEAE |  | DDDDDDDDD |  |
| ADA | GFSF | VPKKP | LVEEEE | GGAPAGG |  | SDSDSDSDS |  |
| EEP | KKEP | PQQPQ | EDDDDE | GLFDFLG |  |  |  |
| DSD | KPVP | QPFPQ | EEYEEY | ITSNSTI |  |  |  |
| PVP | GHHG | EEPEE | EPKKPV | KKAPAKK |  |  |  |
| SPS | EVQQ | PFPQQ | GLGGLG | KKPVPVP |  |  |  |
| DDK | PQQS | KAPPA | KAEEAK | LPTGTPL |  |  |  |
| DKD | FPQQ | PPPPPG | KGKKGK | PKPEPKP |  |  |  |
| VPV | FSFS | PVKVP | KSLLSK | RDRDRDR |  |  |  |
| SEV | PFPQ | KGKGK | NNNNNN | SDSHSDS |  |  |  |
| VES | QPFP | KPPPP | PFPQQP | TPTPTPT |  |  |  |
| QSQ | QQPQ | VAVAV | DDDDDD | SDSDSDS |  |  |  |
| APA | VKKP | EEGEE | AAAAAA | DDDDDDD |  |  |  |
| GTG | AAAE | EEKKP | RRRRRR | PPPPPPP |  |  |  |
| GYG | EKKP | EKGKE | PPPPPP | AAAAAAA |  |  |  |
| KEK | EVEE | GHHGP | AAPPAA | APAPAPA |  |  |  |
| NNN | PKKV | KAPPP | EEKKEE | DEEDEED |  |  |  |
| QQP | EAPP | KKPEV | TAAAAT | EPEPEPE |  |  |  |
| PKK | EPVP | KKPPP | EEEEEE | PKKAKKP |  |  |  |
| PPP | VQQE | LVEEE | SSSSSS | EEEEEEE |  |  |  |
| DED | FEEE | PVAKK |  |  |  |  |  |
| QEL | KKAV | PVPVP |  |  |  |  |  |
| PEE | PKVP | QQGYS |  |  |  |  |  |
|  | TNTG | RRQRR |  |  |  |  |  |
|  | YTPS | VEAPP |  |  |  |  |  |
|  | EDKD | AAEAA |  |  |  |  |  |
|  | EEKV | AQAQA |  |  |  |  |  |
|  | EGAA | DKHKD |  |  |  |  |  |
|  | FSFG | EDWDE |  |  |  |  |  |
|  | GGQG | EEWEE |  |  |  |  |  |
|  | KKIV | GGQGG |  |  |  |  |  |
|  | KNDK | KKAKK |  |  |  |  |  |
|  | PVKP | KKVKK |  |  |  |  |  |
|  | QGYS | PEEPV |  |  |  |  |  |
|  | SAEK | PEVPP |  |  |  |  |  |
|  | SEEN | PRPRP |  |  |  |  |  |
|  | TDDT | PVPEE |  |  |  |  |  |
|  | TFSF | QNNNQ |  |  |  |  |  |
|  | YSQQ | RRNRR |  |  |  |  |  |
|  | AAES | RRSRR |  |  |  |  |  |
|  | APPV | SFGSG |  |  |  |  |  |
|  | DKDE | VEPPP |  |  |  |  |  |
|  | EDED | VPPPK |  |  |  |  |  |
|  | EPPP | VPVPK |  |  |  |  |  |
|  | GCCG | VVEEK |  |  |  |  |  |
|  | KKPK | AAAPA |  |  |  |  |  |
|  | PKKE | AAKAA |  |  |  |  |  |
|  | PPKE | AAPAE |  |  |  |  |  |
|  | PPPE | AGAGA |  |  |  |  |  |
|  | QGGG | AGPGA |  |  |  |  |  |
|  | QNNQ | AVPVP |  |  |  |  |  |
|  | SAAP | DDSDD |  |  |  |  |  |
|  | SIIS | EDKDE |  |  |  |  |  |
|  | VEAE | EEEPP |  |  |  |  |  |
|  | VIKK | EEPVP |  |  |  |  |  |
|  | VPKE | EEYEE |  |  |  |  |  |
|  | AAEA | EGKGE |  |  |  |  |  |
|  | AQTT | EKKPV |  |  |  |  |  |
|  | DSKE | EPEEV |  |  |  |  |  |
|  | DTTD | EPNPE |  |  |  |  |  |
|  | DYEE | EPQPE |  |  |  |  |  |
|  | EAAP | EQKQE |  |  |  |  |  |
|  | EAVV | ESESE |  |  |  |  |  |


|  | EEAG EHHE EREE GGMG KDSA KDVE KGKG KPAA KSFG NEEN NNNQ NPPN NRTP PAAA PPAE PTFS PVEL PVPT QNNN QSQQ QSYG RKEE RRSR SDED SDKD SEED SKSD SNQG SNSN | GGTGG GPPPP GQQSQ GYGYG KAKKP KKAPP KKDKK KPPPA NASAN NDDKK NNNNN NYQQY PGAGP PPPPK PVVVP QAQAQ QPLPQ QQPYP RDRDR RRKRR SDEDS SDVDS SEVSK SPAPS SQQPF VEPEV VPAPA VPEEP VPPPPV |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

## Table A24. Left components of characteristic inverse noncomplementary repeats (material downloaded from DisProt) related to ordered regions

Table includes, for each repeat length, (at most) first 100 n-grams sorted according confidence, lift, and support, all in descending order.

| Repeat length |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| DSG | PPGP | GPPGP | PGPPGP | GPPGPPG | PGPPGPPG | SSSSSSSSS | SDSDSDSDSD |
| NEA | PGPP | PGPPG | PTPPTP | DSDSDSD | KKKIIKKK | DSDSDSDSD | SSSSSSSSSS |
| NVA | GAPG | GGRGG | GRGRGR | GPPGPAG | AEVEAAKK | PGPPGPPGP | EEEEEEEEEE |
| AEL | EKQK | GAPGP | KKSAAE | AEATAEA | GPPGPPGP | GGGGSGGGG | QQQQQQQQQQ |
| VRF | KQKE | PGPAG | ASKKAA | AAKKSAA | нннннннн | QQQQQQQQQ |  |
| VAT | GTPP | PGAPG | EATAEA | AASKKAA | SDSDSDSD | EEEEEEEEE |  |
| FRV | EKRE | ASKKA | LLLLLL | GGGGSGG | SSSSSSSS | SSSSSSSSS |  |
| TLK | ERKE | AGAPG | PPDIPD | GPSSSPG | GGGGGGGG | DSDSDSDSD |  |
| VAS | AAKK | KKAAE | PPGPPS | GQPGPAG | PGPPGPPG | PGPPGPPGP |  |
| TVR | PPSF | EAAKK | SPPGPP | GVPFPVG | KKKIIKKK | GGGGSGGGG |  |
| AAN | VPGP | GQPGP | VEAAKK | PEPSPEP | AEVEAAKK | QQQQQQQQQ |  |
| AVN | ENEA | VEKRE | AGPPGA | QQQAQQQ | GPPGPPGP | EEEEEEEEE |  |
| LEI | GSPG | AANVA | EAASKK | QVEGEVQ | нннннннн |  |  |
| TTK | PGPV | ADAVK | GGRRGG | SLSSLSS | SDSDSDSD |  |  |
| GDY | GRGG | AKKSA | KKVVKK | VVASAVV | SSSSSSSS |  |  |
| KTT | PTGP | ASSSA | PGPAGA | GGGSGGG | GGGGGGGG |  |  |
| ATN | ARVR | AYRYA | SLSSLS | GGGGGGG |  |  |  |
| KNV | KLTV | GPAGA | AAKKSA | GGRGRGG |  |  |  |
| ITA | KAAE | LDADL | AGAPGP | PEPEPEP |  |  |  |
| TAV | KEVI | AGPPG | APPGPP | ннннннн |  |  |  |
| NKA | KGSD | GSPGP | EEELKL | QQQQQQQ |  |  |  |
| SVR | GGRG | KVADA | FLAALF |  |  |  |  |
| YKG | NVAS | RGPPG | GAGGAG |  |  |  |  |
| LTA | VTLT | LLALL | GPPGPV |  |  |  |  |
| IKS | YDGG | SSISQ | GRGGRG |  |  |  |  |
| IGE | ASKK | AGKPG | ISTTSI |  |  |  |  |
| YGI | KKKV | ATSTA | KKNNKK |  |  |  |  |
| TLV | PKKK | ELEKQ | LFEEFL |  |  |  |  |
| PRG | EDSG | GPPGA | LVGGVL |  |  |  |  |
| SDA | ESEA | IENEA | LVVVVL |  |  |  |  |
| RSV | KDGK | IRSGG | PGGPGG |  |  |  |  |



Table A25. Order levels and lengths of homorepeats found in association rules

| Order level | Amino acid | Homorepeat length | Rule lift | Rule confidence |
| :---: | :---: | :---: | :---: | :---: |
| DD | A | 6 | 0.863 | 53.18 |
|  |  | 7 | 0.863 | 55.39 |
|  |  | 8 | 0.703 | 59.04 |
|  |  | 9 | 0.863 | 70.73 |
|  |  | 10 | 0.877 | 80.00 |
|  | D | 3 | 3.063 | 68.65 |
|  |  | 4 | 2.574 | 85.52 |
|  |  | 5 | 2.157 | 93.82 |
|  |  | 6 | 1.568 | 96.65 |
|  |  | 7 | 1.559 | 100.00 |
|  |  | 8 | 1.191 | 100.00 |
|  |  | 9 | 1.220 | 100.00 |
|  |  | 10 | 1.096 | 100.00 |
|  | E | 3 | 3.303 | 74.02 |
|  |  | 4 | 2.758 | 91.63 |
|  |  | 5 | 2.268 | 98.64 |
|  |  | 6 | 1.619 | 99.79 |
|  |  | 7 | 1.559 | 100.00 |
|  |  | 8 | 1.191 | 100.00 |
|  |  | 9 | 1.220 | 100.00 |
|  |  | 10 | 1.096 | 100.00 |
|  | G | 3 | 2.558 | 57.33 |
|  |  | 4 | 2.493 | 82.84 |
|  |  | 5 | 2.299 | 100.00 |
|  |  | 6 | 1.623 | 100.00 |
|  |  | 7 | 1.559 | 100.00 |
|  |  | 8 | 1.191 | 100.00 |
|  |  | 9 | 1.220 | 100.00 |
|  |  | 10 | 1.096 | 100.00 |
|  | H | 4 | 3.010 | 100.00 |
|  |  | 5 | 2.299 | 100.00 |
|  |  | 6 | 1.623 | 100.00 |
|  |  | 7 | 1.559 | 100.00 |
|  |  | 8 | 1.191 | 100.00 |
|  |  | 9 | 1.220 | 100.00 |
|  |  | 10 | 1.096 | 100.00 |
|  | K | 3 | 2.587 | 57.98 |
|  |  | 4 | 2.368 | 78.68 |
|  |  | 5 | 2.299 | 100.00 |
|  |  | 6 | 1.623 | 100.00 |
|  |  | 7 | 1.559 | 100.00 |
|  |  | 8 | 1.191 | 100.00 |
|  |  | 9 | 1.220 | 100.00 |
|  | N | 4 | 1.901 | 63.17 |
|  |  | 5 | 2.299 | 100.00 |
|  |  | 6 | 1.623 | 100.00 |
|  |  | 7 | 1.559 | 100.00 |
|  |  | 8 | 1.191 | 100.00 |
|  |  | 9 | 1.220 | 100.00 |
|  |  | 10 | 1.096 | 100.00 |
|  | P | 3 | 4.313 | 96.65 |
|  |  | 4 | 2.993 | 99.45 |
|  |  | 5 | 2.299 | 100.00 |
|  |  | 6 | 1.623 | 100.00 |
|  |  | 7 | 1.559 | 100.00 |
|  |  | 8 | 1.191 | 100.00 |
|  |  | 9 | 1.220 | 100.00 |
|  |  | 10 | 1.096 | 100.00 |
|  | Q | 3 | 4.026 | 90.23 |
|  |  | 4 | 2.919 | 97.00 |
|  |  | 5 | 2.276 | 98.97 |
|  |  | 6 | 1.619 | 99.74 |
|  |  | 7 | 1.559 | 100.00 |
|  |  | 8 | 1.191 | 100.00 |
|  |  | 9 | 1.220 | 100.00 |
|  |  | 10 | 1.096 | 100.00 |
|  | R | 3 | 2.886 | 64.68 |
|  |  | 4 | 2.208 | 73.37 |
|  |  | 5 | 1.863 | 81.02 |
|  |  | 6 | 1.298 | 80.00 |
|  |  | 7 | 1.357 | 87.03 |
|  |  | 8 | 1.128 | 94.73 |
|  |  | 9 | 1.220 | 100.00 |


|  |  | 10 | 1.096 | 100.00 |
| :---: | :---: | :---: | :---: | :---: |
|  | S | 3 | 3.353 | 75.14 |
|  |  | 4 | 2.830 | 94.04 |
|  |  | 5 | 2.264 | 98.48 |
|  |  | 6 | 1.620 | 99.82 |
|  |  | 7 | 1.558 | 99.90 |
|  |  | 8 | 1.191 | 100.00 |
|  |  | 9 | 1.220 | 100.00 |
|  |  | 10 | 1.096 | 100.00 |
|  | T | 4 | 2.183 | 72.52 |
|  |  | 5 | 2.141 | 93.13 |
|  |  | 6 | 1.623 | 100.00 |
|  |  | 7 | 1.559 | 100.00 |
|  |  | 8 | 1.191 | 100.00 |
|  |  | 9 | 1.220 | 100.00 |
|  |  | 10 | 1.096 | 100.00 |
| OO | C | 4 | 1.953 | 100.00 |
|  |  | 5 | 2.024 | 100.00 |
|  |  | 6 | 3.028 | 100.00 |
|  |  | 7 | 3.209 | 100.00 |
|  | F | 3 | 1.615 | 96.00 |
|  |  | 4 | 1.889 | 96.70 |
|  |  | 5 | 1.984 | 98.03 |
|  |  | 6 | 3.028 | 100.00 |
|  |  | 7 | 3.209 | 100.00 |
|  | I | 3 | 1.644 | 97.70 |
|  |  | 4 | 1.930 | 98.82 |
|  |  | 5 | 1.968 | 97.22 |
|  |  | 6 | 2.954 | 97.56 |
|  |  | 7 | 3.209 | 100.00 |
|  |  | 8 | 7.755 | 100.00 |
|  |  | 9 | 6.629 | 100.00 |
|  | L | 3 | 1.609 | 95.61 |
|  |  | 4 | 1.901 | 97.31 |
|  |  | 5 | 1.981 | 97.86 |
|  |  | 6 | 2.987 | 98.64 |
|  |  | 7 | 3.109 | 96.87 |
|  |  | 8 | 7.238 | 93.33 |
|  |  | 9 | 5.800 | 87.50 |
|  |  | 10 | 11.998 | 87.50 |
|  | M | 5 | 2.024 | 100.00 |
|  |  | - | 3.028 | 100.00 |
|  | V | 3 | 1.577 | 93.72 |
|  |  | 4 | 1.843 | 94.33 |
|  |  | 5 | 2.024 | 100.00 |
|  |  | 6 | 3.028 | 100.00 |
|  |  | 7 | 3.209 | 100.00 |
|  |  | 8 | 7.755 | 100.00 |
|  |  |  | 6.629 | 100.00 |
|  |  | 10 | 13.712 | 100.00 |
|  |  | 4 | 1.953 | 100.00 |
|  | W | 5 | 2.024 | 100.00 |
|  |  | 6 | 3.028 | 100.00 |

## Biography

Samira Almokhtar Alshafah was born on 29th December 1978 in Zawia, Libya. She finished primary school in Almajed School (Harsha-Libya) 1993, high school in Almajed School, (Harsha-Libya) 1996, and Bachelor studies in computer science at Zawia University, Faculty of Engineering, Department of Electronic Engineering (Zawia-Libya) 2001. She was on master studies in Computer science at Libya Academy of Graduate Studies (Tripoli) from 2004 to 2007. and graduate at 2007. Samira enrolled PhD studies in 2010 at Faulty of Mathematics, University of Belgrade, Serbia, as a candidate from Zawia University for PhD studies.

She worked as a teacher in computer science in high school (Harsha- Libya) from 2002 to 2004, and as a lecturer at the Faculty of Engineering, Department of Electronic Engineering (Zawia-Libya) from 2004. Member of the examination committee in the Faculty of Engineering (Zawia-Libya) became 2004 and 2005, and a faculty member (professor) at Faculty of Engineering, Department of Electronic Engineering (Zawia-Libya) after graduate of the Master studies 2007. Samira also taught courses in Java language at the Institute of Higher education of Computer Technologies (Enjela-Libya) 2007-2008. She was the supervisor of the project Graduated bachelor degree at the Higher Institute of Computer Technologies (EnjelaLibya) 2009.

Her research interest was in developing DC motor drive using computer, Handwritten Arabic Characters Recognition, and after enrolled in PhD studies bioinformatics and data mining.

Samira Published two research papers in International Journals and two papers in conferences.

She is married and has three children.

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Име и презиме аутора
Број индекса $\qquad$

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Дозвољавам да се објаве моји лични подаци везани за добијање академског назива доктора наука, као што су име и презиме, година и место рођења и датум одбране рада.
Ови лични подаци могу се објавити на мрежним страницама дигиталне библиотеке, у електронском каталогу и у публикацијама Универзитета у Београду.

Потпис аутора
У Београду,

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Овлашћујем Универзитетску библиотеку „Светозар Марковић" да у Дигитални репозиторијум Универзитета у Београду унесе моју докторску дисертацију под насловом:

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[^0]:    ${ }^{1}$ This can be important for ' N ' regions.
    ${ }^{2}$ Probability of occurrence some AA in region is equal to mole fraction of this AA (taken as a monogram) in this region.

[^1]:    ${ }^{3}$ https://www.ncbi.nlm.nih.gov/Taxonomy/Utils/wprintgc.cgi?

[^2]:    ${ }^{4}$ For example, in protein DP01070 (P42568) only positions 490-567 are annotated as disorder. On the other hands, all predictors listed in MobiDb (http://mobidb.bio.unipd.it/) recognized region 137-475 as disorder. This is in accordance with content of the region which includes mainly disorder promoting AAs, among others sequence of 42 consecutive Serine AAs. In previous version of DisProt database (up to version 6.0.2) some proteins include information about experimentally verified ordered regions, but in the new version (from 7.03) such explicit information are removed.
    ${ }^{5}$ Maybe some region of protein that is predicted (by predictor) as disorder is not annotated as disorder in current version of DisProt, and is count as order in verification process.

[^3]:    ${ }^{6}$ Table not include some n-grams lengths necessary to demonstrate the trend of decreasing percents of retained material. Maximum n-gram length is equal 30 whih corresponds to AA n-grams with length 10.

[^4]:    ${ }^{7}$ Support level and previously used threshold guarantee that no n-gram with small (e.g. statistically nonsignificant) number of occurrences will appear in results. For example, if number of n-grams with specific length is 1.500 .000 than n-gram of such length which occurs less than 150 times will not be taken into account.

[^5]:    8 "Double order level" DO is different from OD because DO determines that left component of repeat is related to disordered while right component is related to ordered region, and vice-versa for OD.

[^6]:    ${ }^{9}$ Support 0.0005 additionally filters input dataset, so palindromes with small number ( $<5$ ) of occurrences in used material were eliminated and not appear in association rules. Due to smaller number of data, for DisProt data threshold for elimination palindromes is less than 2 occurrences.

[^7]:    ${ }^{10}$ These numbers may look like an error because set of statistically significant repeats is subset of set of all repeats. But, because of larger number of repeats in set of all repeats large number of rules have minimal support which didn't passed filter. This is evident if compare average support per rule for all repeats and statistically significant repeats: $0.0128 / 0.0150$ for and $0.0035 / 0.0040$ for repeats with length 3 and 4 repsectively.

[^8]:    Len. Repeats
    7 AEATAEA, DSDSDSD, GGGGGGG, GGGGSGG, GGGSGGG, GGRGRGG, GPPGPPG, HHHHHHH, PEPEPEP, PEPSPEP, QQQAQQQ, QQQQQQQ
    8 GGGGGGGG, GPPGPPGP, HHHHHHH, PGPPGPPG, SSSSSSSSS
    9 DSDSDSDSD, EEEEEEEEE, GGGGSGGGG, PGPPGPPGP, QQQQQQQQQ, SSSSSSSSS
    10 EEEEEEEEEE, QQQQQQQQQQ, SSSSSSSSSS

[^9]:    ${ }^{11}$ Tandem repeats are defined as pair of identical sequences with minimal sequence length 2 . According this definition minimal repeat length that can include tandem repeat is 4 , so percentage calculation is not applicable on repeats with length 3.

[^10]:    ${ }^{12}$ Low capability is consequence of using repeat sequences only in model construction. N-grams have categorical type with possible (depends on their length) very large number of values. As dataset used for model construction does not include all possible n-gram values, class for previously unseen value can not be predicted correctly.

