

Sequence analysis

iFeature: a Python package and web server for features extraction and selection from protein and peptide sequences

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Abstract

Summary: Structural and physiochemical descriptors extracted from sequence data have been widely used to represent sequences and predict structural, functional, expression and interaction profiles of proteins and peptides as well as DNAs/RNAs. Here, we present *iFeature*, a versatile Python-based toolkit for generating various numerical feature representation schemes for both protein and peptide sequences. *iFeature* is capable of calculating and extracting a comprehensive spectrum of 18 major sequence encoding schemes that encompass 53 different types of feature descriptors. It also allows users to extract specific amino acid properties from the AAindex database. Furthermore, *iFeature* integrates 12 different types of commonly used feature clustering, selection and dimensionality reduction algorithms, greatly facilitating training, analysis and benchmarking of machine-learning models. The functionality of *iFeature* is made freely available via an online web server and a stand-alone toolkit.

Availability and implementation: http://iFeature.erc.monash.edu/; https://github.com/Superzchen/iFeature/.

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1 Introduction

In recent years, machine learning techniques have been increasingly used as a powerful means to predict structural and functional properties of proteins and to assist in the annotation of genomic and proteomic data (Larranaga et al., 2006; Libbrecht and Noble, 2015). In this regard, it has proven crucial to transform protein and peptide sequences into effective mathematical expressions that describe their intrinsic correlation with the corresponding structural and functional attributes (Chou, 2011). Over the past decades, an increasing number of diverse feature encoding methods or descriptors extracted from protein and peptide sequence information have been proposed for improving various predictions. Applications include predicting protein structural and function classes (Chou and Fasman, 1978), protein-protein interactions, protein-ligand interactions (Cao et al., 2015; Shen et al., 2007), subcellular locations (Chou and Shen, 2008), enzyme substrates (Barkan et al., 2010; Rottig et al., 2010; Song et al., 2010), among others.

Several web servers and stand-alone software packages have been developed to calculate a variety of structural and physicochemical features, including PROFEAT (Li et al., 2006; Rao et al., 2011), PseAAC (Shen and Chou, 2008), PseAAC-Builder (Du et al., 2012), propy (Cao et al., 2013), PseAAC-General (Du et al., 2014), protr/ ProtrWeb (Xiao et al., 2015), Rcpi (Cao et al., 2015) and PseKRAAC (Zuo et al., 2017). However, in addition to feature extraction, feature selection and ranking analysis is an equally crucial step in machine learning of protein structures and functions. To the best of our knowledge, there is no universal toolkit or web server currently available that integrates both functions of feature extraction and feature selection analysis. It is in this spirit that we developed iFeature, a versatile open-source Python toolkit that bridges this gap. iFeature can be used not only to extract a great variety of numerical feature encoding schemes from protein or peptide sequences, but also for feature clustering, ranking, selection and dimensionality reduction, all of which will greatly facilitate users' subsequent efforts to identify relevant features and construct effective machine learning-based models. In order to facilitate users' interpretability of outcomes, the clustering and dimensionality reduction results can be visualized in form of scatter diagrams. iFeature also supports the integration of different feature types, making it more convenient to train models by combining different feature groups. Lastly, we developed a user-friendly web server for *iFeature*.

2 Implementation

An important advantage of *iFeature* is that it integrates the multi-faceted functionality of feature calculation, extraction, clustering, selection and dimensionality reduction analysis. A complete list of the 18 major encoding schemes is summarized in Table 1. We briefly discuss below.

The first group includes six feature sets, i.e. amino acid composition, composition of *k*-spaced amino acid pairs (Chen *et al.*, 2013; Liu *et al.*, 2017), enhanced amino acid composition, dipeptide composition, dipeptide deviation from expected mean (Saravanan and Gautham, 2015) and tripeptide composition (Bhasin and Raghava, 2004). The secondary group is labeled 'grouped amino acid composition', which also consists of five descriptors (Table 1). For this group, 20 amino acid types are first categorized according to their physicochemical properties, and then the composition of each category is calculated. The third group is the binary encoding scheme in which each amino acid is represented by a 20-dimensional binary vector. The fourth group

includes three types of autocorrelation feature sets: normalized Moreau-Broto autocorrelation, Moran autocorrelation and Geary autocorrelation (Sokal and Thomson, 2006). This feature group allows users to select properties from the AAindex database (Kawashima et al., 2008). The fifth group consists of three feature sets: composition, transition and distribution (Dubchak et al., 1995, 1999). The sixth group is the conjoint triad (Shen et al., 2007). The seventh group contains two sequence-order feature sets, sequence-order-coupling number and quasi-sequence-order (Chou, 2000; Chou and Cai, 2004; Schneider and Wrede, 1994). The eighth group includes the pseudo-amino acid composition and the amphiphilic pseudo-amino acid composition (Chou, 2001, 2005). The ninth group includes two K-nearest neighbor features: KNNprotein and KNNpeptide (Chen et al., 2013). The tenth group is the PSSM encoding scheme, which extracts features from the position-specific scoring matrix (PSSM; Altschul, 1997) generated by PSI-BLAST. The eleventh group is the AAindex encoding scheme where each amino acid is represented by a 531-dimensional vector (Tung and Ho, 2008). The twelfth group is the BLOSUM matrix-derived descriptor (Lee et al., 2011). The thirteenth group is the Z-scale encoding where each amino acid is represented by five physicochemical descriptor variables. Feature groups 14 to 17 are derived from information about the predicted protein secondary structure, disorder, accessible surface area and torsional angles, respectively. The last group includes 16 types of pseudo K-tuple reduced amino acid compositions (Zuo et al.,

Moreover, as high-dimensional features can potentially cause over fitting or high-dimensional disaster (Bellman and Bellman, 1961) and increase of redundant information, machine learning models trained using such high-dimensional initial features often perform poorly in practice. To solve this problem, iFeature further integrates several commonly used feature clustering, selection and dimensionality reduction algorithms to filter out redundant features and retain the useful and relevant ones. All implemented feature analysis algorithms are listed in Table 2. All clustering methods support sample and feature clustering procedures. In cases where users are not familiar with computer programming using Python, we also implemented an online web server of iFeature. It is configured on the extensible cloud computing facility supported by the e-Research Centre at Monash University, equipped with 16 cores, 64 GB memory and a 2 TB hard disk. This configuration can be easily upgraded in line with increasing user demands in the future.

3 Results

In this work, we have developed *iFeature*, a comprehensive, flexible and open-source Python toolkit for generating various sequences, structural and physiochemical features derived from protein/peptide sequences. *iFeature* also allows users to integrate various feature clustering, selection and dimensionality reduction algorithms that facilitate feature importance analysis, model training and benchmarking of machine learning-based models. *iFeature* has been extensively tested to guarantee correctness of computations, and was purposely designed to ensure workflow efficiency. To the best of our knowledge, this is the first universal toolkit for integrated feature calculation, clustering and selection analysis. In the future, we will integrate more analysis and clustering algorithms to enable interactive analysis and machine learning-based modeling. *iFeature* is expected to be widely used as a powerful tool in bioinformatics, computational biology and proteome research.

iFeature 2501

Table 1. List of various descriptors calculated by iFeature

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Pseudo K-tuple reduced amino acids composition PseKRAAC (type1 to type16) —	Predicted main-chain torsional angles	Torsional angles (TS)	_
	Pseudo K-tuple reduced amino acids composition	PseKRAAC (type1 to type16)	_

Table 2. A list of various feature clustering, selection and dimensionality reduction algorithms available in *iFeature*

Type of functionality	Algorithm
Feature clustering	K-means (kmeans)
	Hierarchical clustering (hcluster)
	Mean shift (meanshift)
	DBSCAN (dbscan)
	Affinity propagation (apc)
Feature selection	Chi-square test (CHI2)
	Information gain (IG)
	Mutual information (MIC)
	Pearson's correlation coefficient (pearsonr)
Dimensionality	Principal component analysis (PCA)
reduction	Latent Dirichlet allocation (LDA)
	t-Distributed Stochastic Neighbor
	Embedding (t-SNE)

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