



Protein function prediction with semi-supervised classification based on evolutionary multi-objective optimization

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Predicción de funciones de proteínas usando clasificación semi-supervisada basada en técnicas de optimización multi-objetivo

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Abstract

Proteins are the key elements on the path from genetic information to the development of life. The roles played by the different proteins are difficult to uncover experimentally as this process involves complex procedures such as genetic modifications, injection of fluorescent proteins, gene knock-out methods and others. The knowledge learned from each protein is usually annotated in databases through different methods such as the proposed by The Gene Ontology (GO) consortium. Different methods have been proposed in order to predict GO terms from primary structure information, but very few are available for large-scale functional annotation of plants, and reported success rates are much less than the reported by other non-plant predictors.

The most common approach to perform this task is by using strategies based on annotation transfer from homologues. The annotation process centers on the search for similar sequences in databases of previously annotated proteins, by using sequence alignment tools such as BLASTp. However, high similarity does not necessarily implies homology, and there could be homologues with very low similarity. As an alternative to alignment-based tools, more recent methods have used machine learning techniques trained over feature spaces of physical-chemical, statistical or locally-based attributes, in order to design tools that can be able of achieving high prediction performance when classical tools would certainly fail.

The present work lies on the framework of machine learning applied to protein function prediction, through the use of a modern paradigm called semi-supervised learning. This paradigm is motivated on the fact that in many real-world problems, acquiring a large amount of labeled training data is expensive and time-consuming. Because obtaining unlabeled data requires less human effort, it is of great interest to include it in the learning process both in theory and in practice.

A high number of semi-supervised methods have been recently proposed and have demonstrated to improve the accuracy of classical supervised approaches in a vast number of real-world applications.

Nevertheless, the successfulness of semi-supervised approaches greatly depends on prior assumptions they have to make about the data. When such assumptions does not hold, the inclusion of unlabeled data can be harmful to the predictor. Here, the main approaches to perform semi-supervised learning were analyzed on the problem of protein function prediction, and their underlying assumptions were identified and combined in a multi-objective optimization framework, in order to obtain a novel learning model that is less dependent on the nature of the data.

All the experiments and analyses were focused on land plants (*Embryophyta*), which constitutes an important part of the national biodiversity of Colombia, including most agricultural products.

Keywords: Bioinformatics, Gene Ontology, Semi-supervised Learning, Multi-objective optimization, Cuckoo search.

Resumen

Las proteínas son los elementos clave en el camino desde la información genética hasta el desarrollo de la vida. Las funciones desempeñadas por las diferentes proteínas son difíciles de detectar experimentalmente ya que este proceso implica procedimientos complejos, como las modificaciones genéticas, la inyección de proteínas fluorescentes, métodos de *knock-out* de genes y otros. El conocimiento aprendido de cada proteína es generalmente anotado en bases de datos a través de diferentes métodos como el propuesto por la Ontología Genética (GO). Se han propuesto diferentes métodos para predecir términos GO a partir de la información contenida en la estructura primaria, pero muy pocos están disponibles para la anotación funcional a gran escala de plantas, y las tasas de acierto reportadas son mucho menores que los reportados por otros predictores sobre especies no vegetales.

El enfoque más común para llevar a cabo esta tarea es mediante el uso de estrategias basadas en la anotación basada en transferencia de homólogos. El proceso de anotación se centra en la búsqueda de secuencias similares en bases de datos de proteínas anotadas anteriormente, mediante el uso de herramientas de alineación de secuencias como BLASTp. Sin embargo, una alta similitud no implica necesariamente una homología, y podría haber homólogos con una escasa similitud. Como alternativa a las herramientas de anotación basadas en alineamientos, los métodos más recientes han utilizado técnicas de aprendizaje de máquina entrenados sobre espacios de características físico-químicas o estadísticas, a fin de diseñar herramientas que pueden ser capaces de lograr un alto rendimiento de predicción cuando las herramientas clásicas sin duda fracasarían.

El presente trabajo se encuentra en el marco del aprendizaje de máquina aplicado a la predicción de funciones de proteínas, a través del uso de un paradigma

moderno llamado aprendizaje sem-supervisado. Este paradigma está motivada en el hecho de que en muchos problemas del mundo real, la adquisición de una gran cantidad de muestras de entrenamiento etiquetadas es cara y consume mucho tiempo. Debido a que la obtención de datos sin etiqueta requiere menos esfuerzo humano, es de gran interés para incluirlo en el proceso de aprendizaje, tanto en la teoría como en la práctica. Un gran número de métodos semi-supervisados se han propuesto recientemente y han demostrado mejorar la precisión de los enfoques clásicos supervisadas en un gran número de aplicaciones del mundo real.

Sin embargo, el éxito de los enfoques semi-supervisados depende en gran medida de las suposiciones previas que se tienen que hacer sobre los datos. Cuando estas suposiciones no se cumplen, la inclusión de datos sin etiqueta puede ser perjudicial para el predictor. En este trabajo, se analizan los principales enfoques para llevar a cabo el aprendizaje semi-supervisado sobre el problema de la predicción de funcionesde proteínas, y sus suposiciones subyacentes se identifican y se combinan en un marco de optimización multi-objetivo, con el fin de obtener un nuevo modelo de aprendizaje que sea menos dependiente de las la naturaleza de los datos .

Todos los experimentos y los análisis se centran en las plantas terrestres (*Embryophyta*), que constituyen una parte importante de la biodiversidad nacional de Colombia, incluyendo la mayoría de los productos agrícolas.

Palabras clave: Bioinformática, Ontología Genética, Aprendizaje Semi-supervisado, Otimización multi-objetivo, Búsqueda Cucú.

Introduction

This chapter presents the background, problem statement, hypothesis and objectives of this work. Many of the concepts introduced in this chapter will be discussed in more detail in subsequent chapters, but they are presented here in order to establish the motivation and the context for the rest of the document.

Background

Proteins are versatile macromolecules with a huge diversity of biological functions. They are responsible for most of the biochemical functions of the organelles and, consequently, they are directly involved in all chemical reactions occurring in cells. However, in spite of the wide variety of functions they perform, all proteins share a common basic configuration: a linear polypeptide chain composed by different combinations and repetitions of the twenty amino acids encoded by genes. Although there are almost 8 million sequences in non-redundant databases, for most, we know just that amino acid sequence deduced from the DNA chain ([Levitt, 2009](#)), and thus, the development of methods for determining protein functions from its primary structure becomes an important priority for current science. Since the experimental assessment of protein functions requires, in most cases, to be focused on specific proteins or functions, besides requiring either cloned DNA or protein samples from the genes of interest, some authors have concluded that the only effective route towards the elucidation of the function of some proteins may be by computational analysis ([Baldi and Brunak, 2001](#)).

Many computational resources have been developed in order to predict protein functions (full surveys are presented in ([Friedberg, 2006](#); [Pandey et al., 2006](#); [Zhao et al., 2008c](#))). Nevertheless, very few resources are available for large-scale

functional annotation of non-model species (Conesa and Götz, 2008) and, in particular, just a handful of methods have been recently proposed for predicting protein functions on vegetative species. This is a crucial issue for our country, since the potential development of Colombia through the exploitation of its biodiversity is one of the highest in the world (Cerón et al., 2009). The National Policy for the Promotion of Research and Innovation points out the importance of several research groups and centers that work on the improvement of key agricultural products, highlighting that those are also part of biodiversity in its interaction with human communities (COLCIENCIAS, 1995). Most of that agricultural products are land plants (*Embryophyta*), and thus, new methodologies and algorithms able to accurately predict protein functions over such organisms are strongly needed.

Problem statement

Machine learning methods are widely applied to the extraction of biological knowledge from proteins, in order to obtain models to both represent biological knowledge and to predict their functionality. This field has traditionally been divided into two sub-fields: *unsupervised learning*, in which the system observes an unlabeled set of items represented by their features and has to discover its underlying distribution in order to group them into clusters; and *supervised learning*, where the system observes a labeled training set and the objective is to predict the label y for any new input object. Among the unsupervised methods, the most common algorithm for protein function prediction is the Markov clustering algorithm, which has been mainly used for detecting remote protein families (Chen et al., 2007, 2006). However, as it is only a clustering tool, it is not actually predicting the functions of proteins but merely grouping them into sets whose potential usefulness for protein function prediction has to be elucidated in a posterior stage. Regarding supervised methods, the most prominent example are support vector machines, which have achieved high success in computational biology in general (Vert, 2005) and protein function prediction in particular (Bi et al., 2007; Cai, 2003; Jung et al., 2010). Nevertheless, it is a known fact that only a small number of proteins have actually been annotated for certain functions. Therefore, it is

difficult to obtain sufficient training data for the supervised learning algorithms and, consequently, the tools for protein function prediction have very limited scopes (Zhao et al., 2008c).

Under such circumstances, semi-supervised learning methods provide an alternative approach to protein annotation (Zhao et al., 2008c). Semi-supervised learning (SSL) is halfway between supervised and unsupervised learning: in addition to labeled data, the algorithm is provided with an amount of unlabeled data that can be used to improve the estimations about the data. While labeled instances (annotated proteins) are often difficult, expensive and time consuming to obtain (as they require the efforts of experienced human annotators), unlabeled data is relatively easy to collect in most protein databases.

In order to deal with labeled and unlabeled data, current semi-supervised algorithms have to make strong assumptions about the underlying joint probability of the data. There are several different semi-supervised learning methods, and each one makes different assumptions, but they can be summarized in the following two:

Cluster assumption: If points are in the same cluster, they are likely to be of the same class. Or equivalently: the decision boundary should lie in a low-density region.

Manifold assumption: The (high-dimensional) data lie (roughly) on a low-dimensional manifold. If the data happen to lie on a low-dimensional manifold, however, then the learning algorithm can essentially operate in a space of corresponding dimension, thus avoiding the curse of dimensionality.

There is, however, a strong drawback in semi-supervised algorithms. Since semi-supervised learning is possible only due to the special form of the data distribution that correlates the label of a data point with its location within the distribution, a bad matching of problem structure with model assumption can lead to degradation in classifier performance and, as a result, the inclusion of unlabeled data will degrade prediction accuracy (Chapelle and Schölkopf, 2006; Zhu, 2007).

Hypothesis

Most semi-supervised strategies implement the assumptions about data by introducing regularization terms in the solution of the optimization problem. It is quite straightforward to notice that regularization can be viewed as a special case of a multi-objective optimization problem, where several objective functions are being linearly combined by the introduction of linear weights (regularization constants).

Solving the regularized optimization problem for a unique combination of weights yields a solution that focuses on the objective functions with the highest weights. A more flexible solution can be obtained by directly applying a multi-objective optimization algorithm that deals with all the objective functions at the same time. In this setting, the optimization algorithm does not search for a unique solution, but for the set of all Pareto-optimal solutions with non-convex trade-off surfaces. The use of Pareto optimization provides the means to avoid the need for hard constraints and for a fixed weighting between unsupervised and supervised objectives. Consequently, one would expect a multi-objective approach to semi-supervised classification to perform more consistently across different data sets, and to be less affected by model assumptions.

We propose the hypothesis that tackling the semi-supervised classification problem within the framework of multi-objective optimization, will provide a more flexible framework for the integration of both unsupervised and supervised components and, consequently, it will provide an efficient method for automatic protein function prediction, outperforming supervised methods by exploiting the labeled and unlabeled data that is currently present in protein databases.

Objectives

General objective

Develop a semi-supervised classification strategy based on multi-objective optimization techniques oriented towards the protein function prediction problem.

Specific objectives

1. Select two or more objective functions that adequately reflect the underlying structure of the data for accomplishing semi-supervised assumptions.
2. Develop a multi-objective optimization methodology that finds a set of Pareto-optimal solutions according to the defined objective functions.
3. Develop a strategy for selecting the most biologically feasible solutions among the initial set of Pareto-optimal solutions.
4. Validate the proposed method on the particular case of protein function prediction.

All simulations were implemented on the *R* environment for statistical computing ([R Core Team, 2012](#)). Additional tools were mainly provided by Bioconductor ([Gentleman et al., 2004](#)), and the *seqinR* package ([Charif and Lobry, 2007](#)), all of them freely distributed under the *GNU* General Public License.

Chapter 1

Preliminary concepts

This chapter provides the fundamental concepts behind this work. First, an introduction to protein functionality and structures is given in order to provide

1.1 Functionality of proteins and their structure levels

Proteins are essential macromolecules in life. Their importance for living organisms is straightforward, not only for representing the second largest component in cells after water, but also, and more importantly, for the diversity of biochemical functions they are responsible for. For instance, binding proteins are capable of conforming a wide variety of structurally and chemically different surfaces, allowing themselves for “recognizing” other highly specific molecules in order to perform transport, reception and regulation functions; enzymes use binding plus specific chemical reactivity for speeding up molecular reactions; structural proteins constitute some of the main morphological components of living organisms, being the building blocks of many resistant structures and sources of biomaterials. And such examples just depicts the basis of protein functions universe, because they are only considering their functionality at molecular level. Going further, the scope of protein functions comprises not only the biochemical functions of isolated molecules, but also cellular functions they perform in conjunction with other molecules and even the phenotype they produce in the cell or organism

([Petsko and Ringe, 2004](#)). At cellular level, proteins perform most functions of organelles. Among other tasks, structural proteins in the cytoskeleton are responsible for maintaining the shape of the cell and keeping organelles in place; in the endoplasmatic reticulum, binding proteins transport other molecules between and within cells; in the lysosome, catalytic proteins break large molecules into small ones for carrying out digestion (for a deeper description of subcellular locations of proteins, see ([Chou and Shen, 2007](#))). Phenotypical roles of proteins are harder to determine, since phenotype is the result of many cellular function assemblies and its integration with environmental stimuli. However, by the comparison of genes descended from the same ancestor across many different organisms, or by studying the effects of modifying individual genes in model organisms, several thousands of gene products have been associated with phenotypes ([Benfey and Mitchell-Olds, 2008](#)), and specifically, with affected processes like cell growth or regulation of immune system processes, where proteins have fundamental roles.

Interestingly, a key fact about proteins is that no matter this enormous variety of functions, they all share a common basic conformation: a linear polypeptide chain known as the “primary structure” of the protein. Such a chain is composed by different combinations and repetitions of the twenty amino acids encoded by genes, which in turn determines how the protein folds into higher-level structures. [Figure 1.1](#) depicts the different levels of protein structures.

The secondary structure of the protein can take the form either of alpha helices or of beta sheets, formed through regular hydrogen-bonding interactions between molecules in the main amine and carboxyl groups of amino acids, that is, in the invariant backbone of the chain. In the globular form of the protein, elements of either alpha helix, or beta sheet, or both, as well as loops and links that have no secondary structure, are folded into a tertiary structure. Many proteins are formed by association of the folded chains of more than one polypeptide; this constitutes the quaternary structure of a protein.

The huge variety of functions that can be performed by proteins comes from the large number of three dimensional folding patterns resulting from interactions among the side chains of these amino acids. However, it is important to point out that there are two constraints for a polypeptide to be a protein. First, it must be able to form a stable tertiary structure (or fold) under physiological conditions.

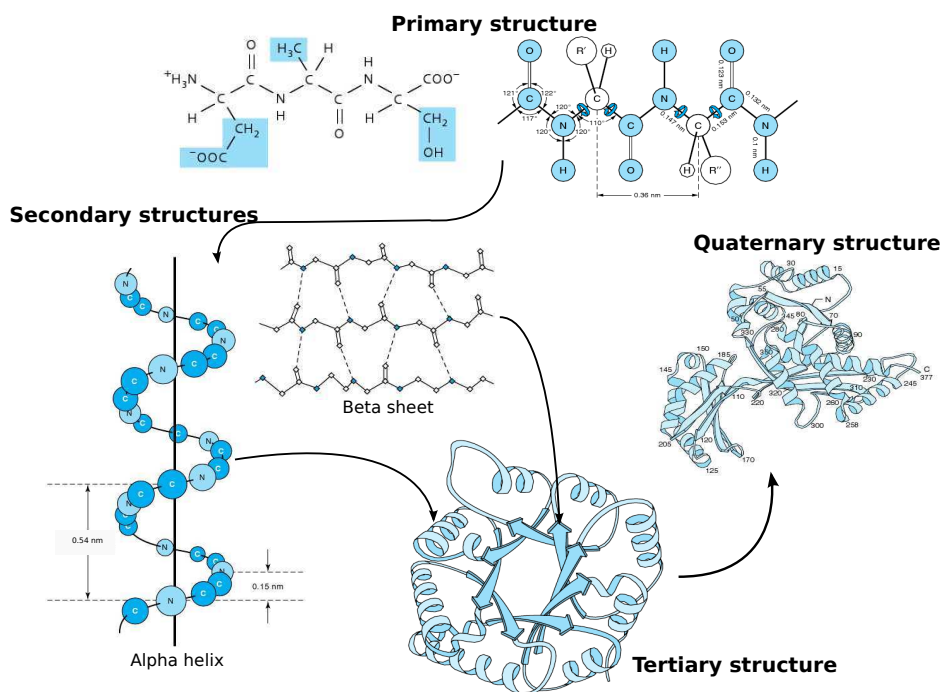


Figure 1.1: Levels of protein structure

Second, the folded pattern should have enough flexibility to interact with other molecules in order to allow protein functionality. Presumably because of these constraints, the number of folds adopted by proteins, though large, is limited (Petsko and Ringe, 2004).

1.2 Gene ontology

With all this functionality associated to proteins, the same definition of protein function turns into a fuzzy concept, leading different researchers to denote the functions of proteins differently. As an effort to provide consistent descriptors for key domains of molecular biology, the Gene Ontology (GO) project aims to construct controlled and structured vocabularies known as ontologies, and apply such vocabularies in the annotation of sequences, genes or gene products in biological databases (The Gene Ontology Consortium, 2004). An ontology is defined as a systematic arrangement of categories, together with the relations among them; in the case of ontologies defined at GO, each category corresponds to

a functional label or “GO term”, related with other terms through “is-a” or “part-of” relationships. Structurally, these ontologies could be modeled hierarchically as directed acyclic graphs, which means that every child node may have more than one parent node.

There are three ontologies in GO, defined to describe three non-overlapping domains of molecular biology: molecular function, cellular component and biological process. Molecular function (MF) refers to biochemical activities at molecular level, no matter what entities are in charge of accomplishing that function or the context where it takes place; examples of molecular functions are “enzyme regulator activity”, “binding activity” or “transport activity”. Cellular component (CC) refers to the specific sub-cellular location where a gene product is active, describing different parts of the eukaryotic cell; cellular components include “ribosome”, “cytoplasm” or “Golgi apparatus”. Biological process (BP) refers to a series of events or molecular functions, with a defined beginning and end, to which the gene or gene product contributes; examples are “reproduction”, “protein metabolic process” or “cell death”.

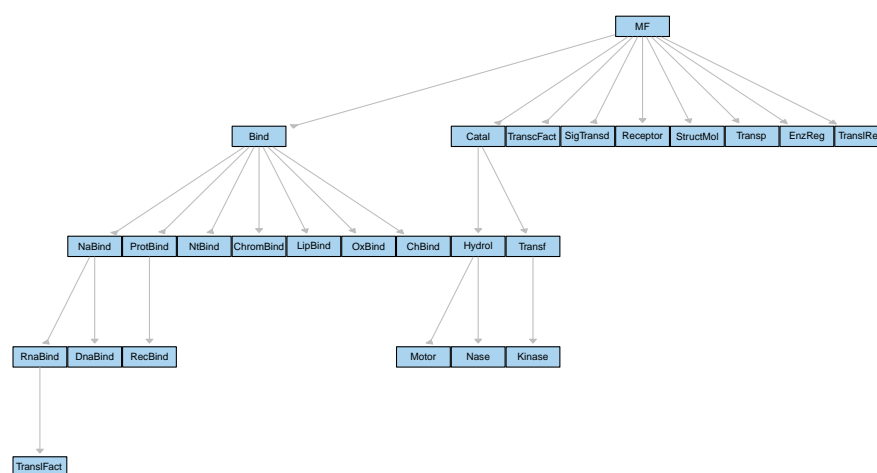


Figure 1.2: Molecular Function ontology from the plants GO-slim. The list of acronyms can be found in table 2.1

Currently, as of February 2013 there are 38137 defined GO terms, distributed over 9467 molecular functions, 3050 cellular components and 23928 biological

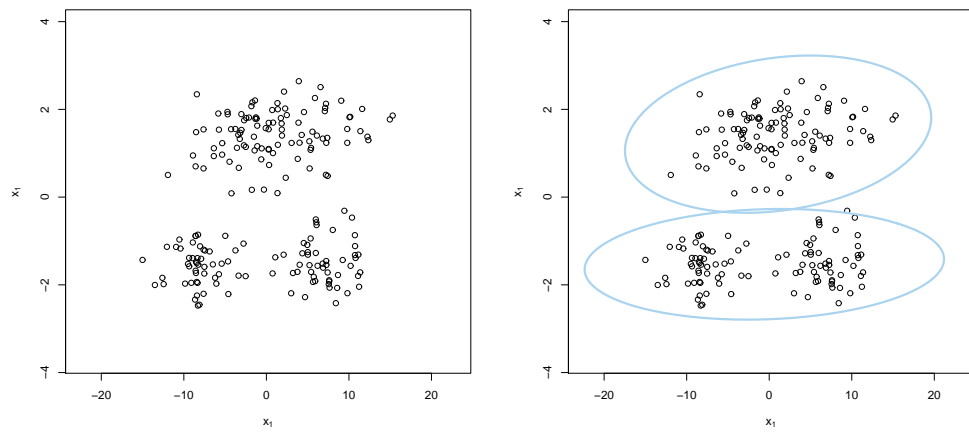
processes. However, it is often useful to have a less detailed set of categories to produce a high-level overview of functions distribution. For this reason, a number of custom datasets, named GO slims, are also maintained by The Gene Ontology Consortium. In those versions, more specific terms have been collapsed up into more general parent terms, and they are particularly useful for analyzing just a subsection of the GO in order to study a particular field or for a general genome-wide analysis. In particular, species-specific slims are maintained for plants (Berardini et al., 2004), *Candida Albicans* (Costanzo et al., 2006), *Schizosaccharomyces Pombe* (Aslett and Wood, 2006) and *Saccharomyces Cerevisiae* (Hirschman et al., 2006). As an example, figure 1.2 shows the GO-terms in the Molecular Function ontology for the plants GO slim.

1.3 Paradigms in machine learning

Machine learning provides the tools for constructing models that represent biological knowledge and use it to predict biological outcomes. In particular, this work is focused on semi-supervised learning methods for predicting protein functionality. However, in order to understand the nature of semi-supervised learning, it will be useful to first define classical supervised and unsupervised learning frameworks. Then, semi-supervised and semi-unsupervised learning will be properly defined.

1.3.1 Supervised and unsupervised learning

The field of machine learning has traditionally been divided into two sub-fields: *unsupervised learning* and *supervised learning*. In unsupervised learning, the system observes an unlabeled set of items represented by their D-dimensional feature vectors $\{\mathbf{x}_i\}_{i=1}^N$, $\mathbf{x}_i \in \mathbb{R}^D$, drawn from a feature space \mathcal{X} . The main objective is to discover its underlying distribution in order to group them into K clusters. In this setting, there is no “right” answer, since there is not a prior knowledge about the correct membership of the samples (which is why this paradigm is termed as unsupervised).



(a) Unclustered data

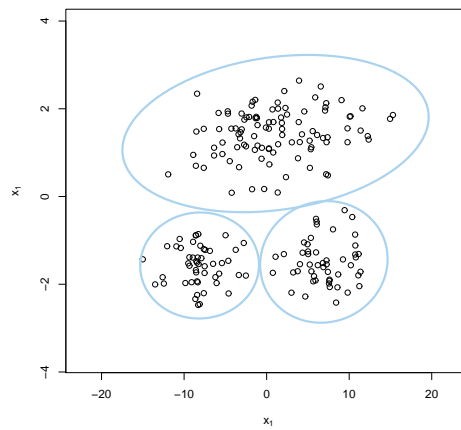
(b) Clustered data with $K=2$ (c) Clustered data with $K=3$ **Figure 1.3:** Example of unsupervised learning

Figure 1.3 depicts an example of the outcome of an unsupervised learner for two (Figure 1.3(b)) and three clusters (Figure 1.3(c)). Since there is no supervision, both clustering schemes could be valid solutions for different tasks.

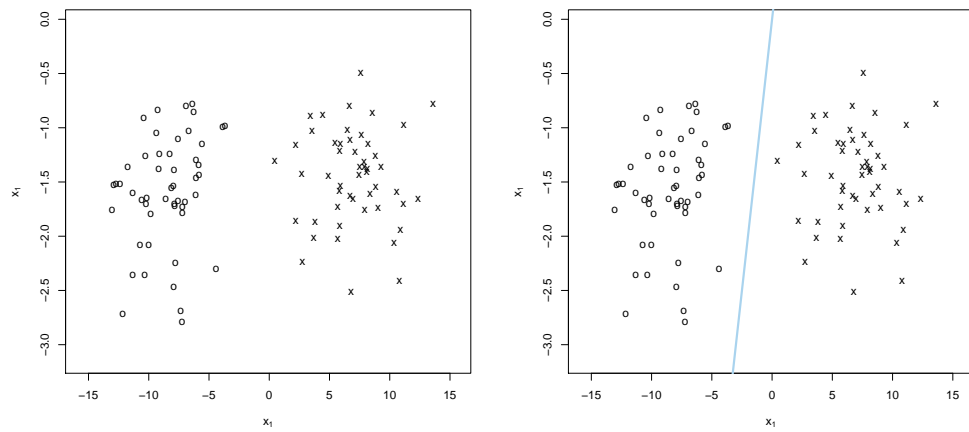
On the other hand, supervised learning algorithms have access to a labeled training set consisting of (feature, label) pairs, denoted by $\{(\mathbf{x}_i, y_i)\}_{i=1}^N$. In this setting, feature vectors are again represented by D -dimensional vectors $\mathbf{x}_i \in \mathbb{R}^D$, while labels are the desired predictions for each instance. When labels are continuous variables, that is, $y \in \mathbb{R}$, the constructed model is a regressor. In turn, when the predictions are constrained to a finite set of discrete labels, $y \in y_{j=1}^C$, the trained model is a classifier. Then, such classifier is a mathematical function $f(\mathbf{x})$, that associates each feature vector with its corresponding true label:

$$f : \mathcal{X} \mapsto \mathcal{Y} \quad (1.1)$$

Figure 1.4 depicts an example of a two-class supervised classification problem. In this setting, two approaches can be used for deriving the decision function $f(\mathbf{x})$: discriminative and generative. Discriminative classifiers focus on computing the decision frontier between the classes as in Figure 1.4(b), where a linear discriminant classifier is used. Generative classifiers, on the other hand, focus on modeling the data in order to obtain one model per class and provide membership probabilities for new instances. Figure 1.4(c) depicts the contour levels of two Gaussian probability distributions adjusted to the data.

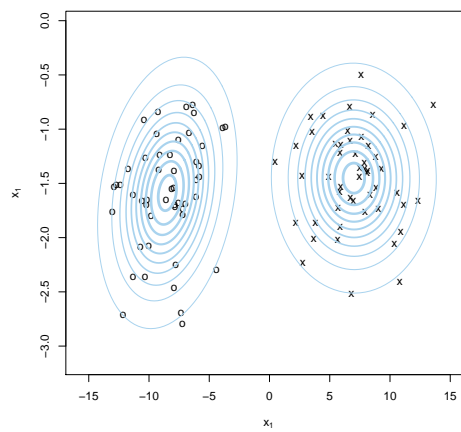
1.3.2 Transductive, semi-unsupervised and semi-supervised learning

The idea of using both labeled and unlabeled data for designing robust predictors has been on the machine learning community since around the middle sixties with several proposals on self-training (see, for example, Scudder III (1965) and Fralick (1967)) and transductive inference (Vapnik and Chervonenkis, 1974). The main motivation behind this kind of learning comes from the fact that in many real-world problems, acquiring a large amount of labeled training data is expensive and time-consuming. Because obtaining unlabeled data requires less human effort, it



(a) Data

(b) Discriminant classifier



(c) Generative classifier

Figure 1.4: Example of supervised learning

is of great interest to include it in the learning process both in theory and in practice.

There are several ways of combining those two sources of data, given rise to different paradigms in machine learning. Consider a system that can observe two sources of data: the points $\mathcal{X}_L = \{\mathbf{x}_i\}_{i=1}^L$ for which labels $\{y_i\}_{i=1}^L$ are provided, and the points $\mathcal{X}_U = \{\mathbf{x}_i\}_{i=L+1}^{L+U}$ the labels of which are not known. “Transductive” learning is only interested on predicting the labels of the unlabeled data in the training dataset, that is, in learning a function of the form:

$$f : \mathcal{X}_U \mapsto \mathcal{Y}_U \quad (1.2)$$

here, f is expected to be a good predictor on the unlabeled data and is defined only on the given training sample, and is not required to make predictions outside. On the other hand, “inductive semi-supervised” learning is interested on designing a function able to predict the labels on future test data. That is:

$$f : \mathcal{X} \mapsto \mathcal{Y} \quad (1.3)$$

here, f is expected to be a good predictor over the whole feature space and, implicitly, it is expected to be better than the supervised classifier trained on the labeled data alone. Like in supervised learning, a common estimation of the performance of the system with future data can be obtained by using a separate test sample, which is not available during training. This setting is sometimes simply called semi-supervised classification and constitutes the main subject of the present work.

An interesting analogy presented in (Zhu and Goldberg, 2009), proposes that semi-supervised learning is like an in-class exam, where the questions are not known in advance, and a student needs to prepare for all possible questions; in contrast, transductive learning is like a take-home exam, where the student knows the exam questions and needs not prepare beyond those.

Finally, other forms of partial supervision are also possible. Constrained clustering is an extension to conventional unsupervised clustering which, in addition to the unlabeled data, is fed with some supervised information about the clusters. Such information is commonly provided in the form of “must-link” and

“cannot link” constraints, imposing restrictions over pairs of instances that must be or cannot be in the same cluster. [Seeger \(2006\)](#) defines this category of algorithms as “semi-supervised learning”, since their main objective is to estimate the probability distribution of the data as in unsupervised learning methods.

1.3.3 Remarks on the application of machine learning methods for protein function prediction

Following the notation of ([King and Guda, 2008](#)), let \mathcal{P} be the protein space (the set of all possible protein sequences). Labeled data will be noted as \mathcal{P}_L while unlabeled data will be noted by \mathcal{P}_U . Then, we have:

$$\mathcal{P} = \mathcal{P}_L \cup \mathcal{P}_U \quad (1.4)$$

First, let \mathcal{X} be the feature space generated from a characterization function $\zeta : \mathcal{P} \mapsto \mathcal{X}$. This function accepts as input a protein sequence and returns a feature vector $\mathbf{x} \in \mathbb{R}^D$, with D physical-chemical and/or statistical attributes of the protein sequence (this will be explained in more detail in the next chapter). In general terms, the function ζ is neither injective nor surjective, that means that several elements in \mathcal{P} can be mapped into the same element of \mathcal{X} . Besides, there may be some feature vectors in \mathcal{X} for which no instance in \mathcal{P} could possibly exist in nature ([King and Guda, 2008](#)).

Let \mathcal{Y} be the label set, with all the discrete labels that can be assigned to a given protein. This set corresponds to all the functional annotations that can be associated to proteins (the set of GO terms). It is necessary to have into account that those labels are not mutually exclusive as in traditional supervised classification, that is, one protein can be associated to more than one GO term at the same time. Then, each labeled instance will be associated not only to a single label y but to a set of Q labels that is a subset of the whole set of possible labels $\mathbf{y} \subseteq \mathcal{Y}$.

This gives rise to a multi-label learning problem, a branch of machine learning where multiple target labels must be assigned to each instance. Multi-label learning methods can be transformed into one or more single-label classification problems by employing several topologies [Tsoumakas and Katakis \(2007\)](#). In this

regard, a recent paper by [Giraldo-Forero et al. \(2013\)](#), empirically demonstrated that the performance of the “binary relevance” topology, together with a technique of class balance, remains above several recently proposed techniques for the problem of predicting protein functions. Binary relevance decomposes the multi-label decision function $f : \mathcal{X} \mapsto \mathcal{Y}^Q$, into Q binary classifiers $f^q : \mathcal{X} \mapsto \{+1, -1\}$, where each binary classifier decides whether or not a given instance should be associated to the q -th class. This approach is also known as “one against all”, and will be used for all the experiments in this work. Therefore, the labels will be assumed to be binary throughout the rest of the document.

Having defined \mathcal{X} and \mathcal{Y} , it is now possible to apply any machine learning algorithm (supervised or unsupervised) to the prediction of protein functions.

1.4 Single-objective and multi-objective optimization for machine learning methods

The training of most machine learning method comprise of two steps: selecting a candidate model (that in general terms is a parametric function) and then, estimating the parameters of the model using an optimization algorithm and available data. Therefore, all learning problems can be considered as optimization problems ([Jin and Sendhoff, 2008](#)).

For the particular case of protein function prediction, let $\mathbf{x}_i \in \mathcal{X}$ be the feature vector representing the protein $p_i \in \mathcal{P}$, and let $y_i \in \mathcal{Y}$ be the label assigned to that element. The predictor function f should satisfy $f(\mathbf{x}_i) = y_i$, $i = 1, 2, \dots, N$. In general terms, this function can belong to a family of parametric functions with parameters $\boldsymbol{\theta}$:

$$f \in \{f_{\boldsymbol{\theta}}\}_{\boldsymbol{\theta} \in \mathcal{T}} \tag{1.5}$$

where \mathcal{T} is the space of all the vectors of parameters. Learning the predictor is achieved by the correct selection of the vector of parameters $\boldsymbol{\theta}$, and such adjustment can be understood as an optimization process depending on the labeled data (for the supervised case) or both the labeled and unlabeled data (for the semi-supervised case).

Table 1.1: Examples of loss functions and the optimization methods used in several supervised machine learning algorithms

Method	Loss function	Optimization algorithm
Least squares classifier	$(y_i - f_{\theta}(\mathbf{x}_i))^2$	Analytical solution
Multilayer perceptron	$-(f_{\theta}(\mathbf{x}_i)y_i) H(f_{\theta}(\mathbf{x}_i)y_i)^{\dagger}$	Backpropagation (gradient descent)
Support vector machine	$\max(0, 1 - y_i f(\mathbf{x}_i))$	Quadratic programming

[†] $H()$ stands for the Heaviside step function.

In order to perform the estimation of parameters, it is first necessary to define one or multiple optimization criteria. The most common criterion for supervised and (inductive) semi-supervised learning is to define an objective function that reflects the quality of the adjusted model, by minimizing the prediction error over the training set:

$$o_{err}(\boldsymbol{\theta}) = \sum_{i=1}^N \ell(f_{\boldsymbol{\theta}}(\mathbf{x}_i), y_i) \quad (1.6)$$

where ℓ is some loss function that depends on the desired labels y_i and the predicted labels $f_{\boldsymbol{\theta}}(\mathbf{x}_i)$ of the training set. Table 1.1 shows several examples of the loss functions used in common machine learning algorithms. The optimization process must find the optimal vector of parameters $\boldsymbol{\theta}^*$ such that:

$$\boldsymbol{\theta}^* = \arg \min_{\boldsymbol{\theta} \in \mathcal{T}} o_{err}(\boldsymbol{\theta}) \quad (1.7)$$

However, minimizing the training error is not the only objective to be considered, since this may result in over-fitting the training data and consequently obtaining poor performance on unseen data. To avoid this problem, the complexity of the model must be also controlled. This is usually expressed by defining

another objective function in terms of the norm of f_{θ} :

$$o_{comp}(\theta) = \|f_{\theta}\| \quad (1.8)$$

The most common approach for including this new objective in the training process, is by aggregating the two objectives into a scalar objective function. This process is known as “regularization”:

$$\theta^* = \arg \min_{\theta \in \mathcal{T}} \{o_{err}(\theta) + \lambda o_{comp}(\theta)\} \quad (1.9)$$

where λ is a positive scalar constant defined by the user, and the correct determination of this parameter is not a trivial task. The work by [Marler and Arora \(2009\)](#) provides an analysis that reveals several fundamental deficiencies of the scalarization of multi-objective optimization problems, showing that although the weighted sum method is easy to use, it provides only a linear approximation of the preference function.

In fact, almost every real-world problem involves simultaneous optimization of several incommensurable and often competing objectives ([Zitzler, 1999](#)). A more flexible approach to deal with this kind of problems comes from the Pareto-based optimization strategies. In this setting, the objective function does not provide a scalar output, but a vector with the evaluation of all the objectives considered:

$$\mathbf{o}(\theta) = [o_1(\theta), o_2(\theta), \dots, o_M(\theta)] \quad (1.10)$$

where M is the number of objectives. It is rarely the case that there is a single solution that optimizes all the objectives at the same time. Therefore, instead of a single solution, a set of trade-off solutions must be returned by the optimization algorithm. Here, it is necessary to modify the notion of optimality. The most commonly adopted notion is the generalized by [Pareto \(1896\)](#), called “Pareto optimality”. in a minimization multi-objective problem, a solution θ^* is said to be Pareto-optimal if there is no other feasible solution $\theta \in \mathcal{T}$ which would decrease one of the objective functions without causing a simultaneous increase in at least one other objective function. A formal definition can be established by first

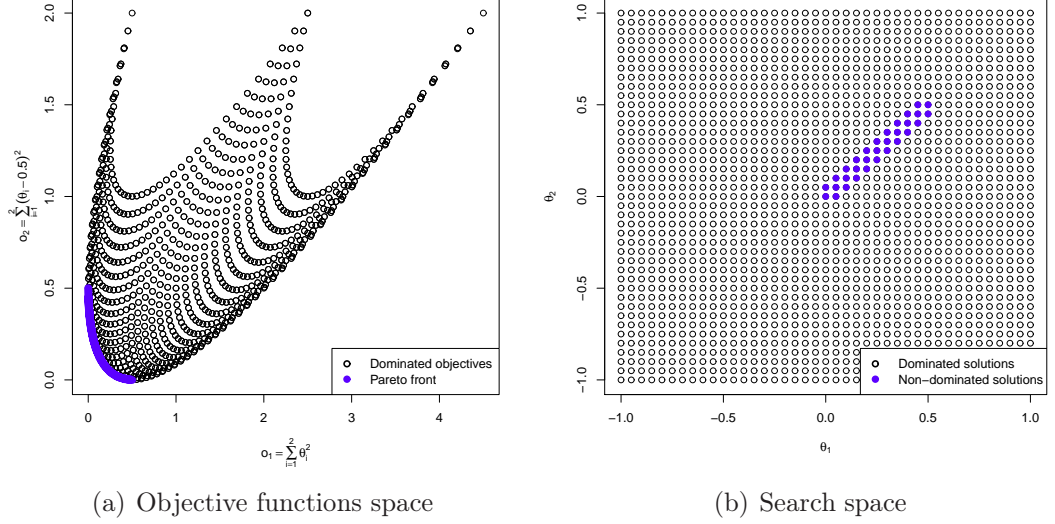


Figure 1.5: Example of a Pareto front

defining the notion of Pareto-dominance (Zitzler, 1999) between two solutions θ and ϕ :

$$\theta \preceq \phi \quad (\theta \text{ weakly dominates } \phi) \iff \forall i \in \{1, 2, \dots, M\} : o_i(\theta) \leq o_i(\phi) \quad (1.11)$$

$$\theta \prec \phi \quad (\theta \text{ dominates } \phi) \iff \theta \preceq \phi \wedge o(\theta) \neq o(\phi) \quad (1.12)$$

then, a solution θ^* is said to be Pareto-optimal if and only if:

$$\nexists \theta \in \mathcal{T} : \theta \prec \theta^* \quad (1.13)$$

The whole set of Pareto-optimal solutions is called the “non-dominated set”, and the set of objective vectors corresponding to the evaluations of the non-dominated set is called the “Pareto front”. As an example, Figure 1.5 depicts the Pareto front obtained when minimizing two competing objective functions $o_1(\theta) = \sum_{i=1}^2 \theta_i^2$ and $o_2(\theta) = \sum_{i=1}^2 (\theta_i - 0.5)^2$, defined over a two-dimensional search space $\theta = [\theta_1, \theta_2]$ restricted to the intervals $-1 \leq \theta_1, \theta_2 \leq 1$.

While the minimum of o_1 is obtained for $\theta = [0, 0]$, this solution causes o_2

to produce an output of 0.5. In turn, the minimum value of o_2 is reached for $\boldsymbol{\theta} = [0.5, 0.5]$, causing o_1 to produce an output of 0.5. These two points correspond to the ends of the Pareto front in Figure 1.5(a), while all the intermediate points lying on the Pareto front are also good trade-off solutions for the multi-objective problem: in the absence of preference information, none of the corresponding trade-offs can be said to be better than the others. Figure 1.5(b), depicts the location of the non-dominated solutions in the original search space.

Chapter 2

Supervised Gene Ontology prediction for *Embryophyta* organisms

Currently, there are almost 8 million sequences in non-redundant databases, including the complete genomes of $\approx 1,800$ different species. However, for most of them, we know just their primary structure: the linear amino acid sequence deduced from the DNA chain ([Levitt, 2009](#)). Assessment of protein functions needs in most cases of experimental approaches carried out in the lab. Such approaches require either cloned DNA or protein samples from the genes of interest. Experimental procedures for probing protein function includes DNA micro-arrays for providing expression patterns of genes; two dimensional gel electrophoresis which can separate complex protein mixtures into their components for being identified by mass spectrometry; gene knockout experiments for studying phenotypical effects of inactivating determined gene products and experiments with green fluorescent protein for determining gene products locations.

Unfortunately, this procedures must be usually focused on specific proteins or functions, and the current dimensions of data bases makes of manual annotation a difficult and almost intractable problem. Additionally, experimental determination of the function of many proteins is very likely to be hard, because the function may be related specifically to the native environment in which a par-

ticular organism lives. Such perspective has lead some authors to conclude that the only effective route toward the elucidation of the function of some proteins may be computational analysis and prediction from amino acid sequences, obtaining a first hint toward functionality that later can be subjected to experimental verification ([Baldi and Brunak, 2001](#), chapter 1).

2.1 Gene Ontology predictors

Many approaches have been developed in this matter (for full surveys, see ([Friedberg, 2006](#); [Pandey et al., 2006](#); [Zhao et al., 2008c](#))). One of the earliest applications, yet still one of the more popular bioinformatics tools is the Basic Local Alignment Search Tool for proteins (BLASTP) ([Altschul et al., 1997](#)) which has been applied for obtaining annotation transfers based on sequence alignments. Also, a high number of methods (GOblet ([Groth et al., 2004](#)), OntoBlast ([Zehetner, 2003](#)), GOFigure ([Khan, 2003](#)) and GOtcha ([Martin et al., 2004](#))) are based on the idea of refining and improving initial results from classic alignment tools such as BLASTp, by performing mappings and weightings of GO terms associated to BLASTP predictions. However, in such methods, the failure of conventional alignment tools to adequately identify homologous proteins at significant E-values is not considered ([Hawkins et al., 2009](#)). The same applies for some more recent methods that have improved specific points of this methodology such as speeding up the procedure through decision rules (([Jones et al., 2008](#))), including additional functionality for visualization and data mining (([Conesa and Götz, 2008](#))) or also including some statistics of GO terms to refine selection (([Vinayagam et al., 2006](#))). In order to avoid the dependency to BLAST alignments in the cases where the alignment-based annotation transfer approach is not so effective, more recent methods have used machine learning techniques trained over feature spaces of physical-chemical, statistical or locally-based attributes. Those methods employ techniques such as neural networks (ProtFun ([Jensen et al., 2003](#))), Bayesian multi-label classifiers (([Jung and Thon, 2008](#))) and support vector machines (SVM-Prot ([Cai, 2003](#)), GOKey ([Bi et al., 2007](#)), PoGO ([Jung et al., 2010](#))), obtaining high performance results in their own re-

spective databases, mostly composed by model organisms such as bacteria and a few high order species.

There are, however, several aspects that must be discussed about current performance in prediction of GO terms, when applied to non-model organisms such as land plants (*Embryophyta*). First, from the previously described methods, only Blast2GO (Conesa and Götz, 2008) was specialized for predicting GO terms in plant proteins. In fact, as it is pointed out by the authors of Blast2GO, very few resources are available for large-scale functional annotation of non-model species. Some methods specialized on vegetative species have been proposed recently, but they are only intended for performing cellular component predictions (Predotar (Small et al., 2004), TargetP (Emanuelsson et al., 2000), Plant-mPloc (Chou and Shen, 2010)). Moreover, Predotar and TargetP can discriminate among only three or four cellular location sites. Plant-mPloc, in turn, covers twelve different location sites and it was rigorously tested over a set of proteins with less than 25% of identity among them, where homologue-based tools like BLASTP would certainly fail. For such dataset, they obtained an overall success rate of 63.7%, much less than reported by other cellular location predictors tested over non-plant datasets. Second, none of the existing methods can be used to deal with plant proteins that can simultaneously exist or move between two or more different location sites (Chou and Shen, 2010), or belong to multiple functional classes at the same time (Briesemeister et al., 2010).

In order to improve the performance of current GO term predictors for land plants, it would be useful to have a better understanding of the underlying relationships between primary structure information and protein functionality. However, the structure of the machine learning models behind high-accuracy predictors often makes difficult to understand why a particular prediction was made (Briesemeister et al., 2010). In this sense, a recent method called Yloc (Briesemeister et al., 2010) was proposed for analyzing what specific features are responsible for given predictions. This method, nevertheless, is not intended to predict GO terms, but instead, it uses annotation information from PROSITE (Sigrist et al., 2010) and GO as inputs to the predictor. Additionally, their study is only focused on predicting protein locations in the cell.

Since most of the current GO prediction methods are limited to a few arbitrary functional classes and single ontologies, they cannot provide any information about relationships on the predictability at the various levels of protein functionality (molecular, cellular, biological), which could be another key element for determining how the information of the primary structure is related to protein function.

2.2 Proposed methodology: prediction of GO terms in *Embryophyta* organisms with pattern recognition techniques

This section presents an analysis on the predictability of GO terms over the *Embryophyta* group of organisms, which is composed by the most familiar group of plants including trees, flowers, ferns, mosses, and various other green land plants. The analysis provides the following key elements: predictions are made by using features extracted solely from primary structure information; analysis comprises most of the available organisms belonging to the *Embryophyta* group; biases due to protein families are avoided by considering only proteins with low similarity among them and a strong evidence of existence; predictions and analysis are made over a set of categories belonging to the three ontologies; proteins are allowed to be associated to several GO terms simultaneously.

Results from this chapter answer whether it is possible to predict most GO-slim terms from primary structure information, what categories are more susceptible to be predicted, which ontology is most related to the information contained in the primary structure and what relationships among ontologies could be influencing the predictability at different levels of protein functionality in land plants.

The implemented methodology for assessing predictability of GO terms in *Embryophyta* proteins comprises the following parts: (i) selection of the protein sequences conforming the database in order to cover the highest number of available plant proteins, while ensuring high confidence annotations and avoiding possible biases; (ii) categories describing positive and negative samples associated to each GO term are determined in order to minimize the impact of hierarchical

relationships; (ii) protein sequences are mapped into feature vectors that encode a number of attributes of varied nature; (iii) computed features are clustered into groups of similar information content; (iv) one binary classifier is learned for each GO term and each feature cluster in order to evaluate the prediction power of individual clusters, and finally (v) one binary classifier is learned for each GO term using the whole set of features in conjunction with an automatic feature selection strategy in order to determine the global predictability of each GO term. The following subsections describe the methods employed for each part of the methodology.

2.2.1 Database

The database comprises all the available *Embryophyta* proteins at UniProtKB/Swiss-Prot database ([Jain et al., 2009](#), file version: 10/01/2013), with at least one annotation in the Gene Ontology Annotation (GOA) project ([Barrell et al., 2008](#), file version: 7/01/2013). The resulting set comprises proteins from 189 different land plants.

In order to avoid the presence of protein families that could bias the results, the dataset was filtered at several levels of sequence identity using the Cd-Hit software ([Li and Godzik, 2006](#)). The main results are reported for the lowest identity cutoff (30%). However, additional analyses at 40%, 50%, 60%, 70% and 80% were also performed in order to provide further information on the robustness of the method. It is important to mention that, according to ([Petsko and Ringe, 2004](#))[chapter 4], function can be inferred from homology based transfer methods in cases where a protein has more than about 40% sequence identity to another protein whose biochemical function is known, and if the functionally important residues are conserved between the two sequences. Identities under 40% do not allow to make function prediction with high confidence when using the classical methods.

The main set comprises a total of 3368 protein sequences, from which 1973 sequences are annotated with molecular functions, 2210 with cellular components and 2798 with biological processes. Finally, it is also important to clarify that, although computational and indirectly derived annotations increase

coverage significantly, they probably contain a higher portion of false positives (Rhee et al., 2008). Consequently, annotations associated to evidence codes from automatically-assigned annotations were not included in the analyses.

2.2.2 Definition of classes

Although, in principle, the method can be trained to predict any GO term for which there are enough training sequences, all tests were performed over the set of categories defined by the plants GO slim developed by The Arabidopsis Information Resource - TAIR (Berardini et al., 2004, file version: 14/03/2012). This choice was made because GO includes a large number of categories that do not occur in plants, due to its broad size. In turn, slims are smaller, more-manageable sub-sets of GO, that focus on terms relevant to a specific problem or data set (Davis et al., 2010), thus allowing to generate higher-level annotation more robust to tests of statistical significance (Rhee et al., 2008).

Positive and negative samples associated to each GO term are selected by considering the propagation principle of GO in order to avoid hierarchical relationships. Otherwise, as members of child categories are also included on parent categories, classes would be totally overlapped and no standard classification tool could be successful. If a protein is predicted to be associated to any given GO term, it must be automatically associated to all the ancestors of that category and thus, it is enough to predict only the lowest level entries. Consequently, for each GO term, positive samples are all those proteins that have been annotated with this term or any of its descendants, excepting those descendants that are also included as categories. All the remaining samples in the database are selected as negative samples for that GO term. It is very important to keep in mind the interpretability implications of this procedure, in order to correctly understand posterior results. As said before, GO slims are reduced versions of GO, where more specific terms have been collapsed up into parent terms. For instance, the term “nucleic acid binding” has five children in the original GO molecular function ontology, while it only has three children in the plants GO slim, what means that the remaining two children were merged up into the parent. After removing

redundant entries, sequences remaining in “nucleic acid binding” category are precisely the ones associated to that two merged categories, as well as to every other function not explicitly listed in GO. For that reason, a correct interpretation of this category should be now “other members of nucleic acid binding”, clarifying the disruption with the three explicit children. For avoiding possible confusions, redefined classes are marked with an asterisk throughout the document.

After defining the membership of the sequences, categories with less than 30 proteins were discarded because they did not have enough samples to train a statistically reliable classifier. The final set is thus comprised by 14 GO terms in the molecular function ontology, 20 GO terms in the cellular component ontology and 41 GO terms in the biological process ontology. Table 2.1 shows the final list of categories, as well as the acronyms used to cite them throughout this paper and the number of samples in each one for the 30% identity cutoff.

2.2.3 Characterization of protein sequences

Protein sequences were mapped into feature vectors by extracting three types of attributes: physical-chemical features, primary structure composition statistics and secondary structure composition statistics (see Table 2.2).

The first group is intended to provide information directly related with biochemistry of the molecule and is constituted by six properties related to basic physical-chemical attributes of amino acids. Weight of the sequence is influenced by the size of the constituent amino acids; polarity of amino acid side chains determines the percentage of positively and negatively charged residues in the sequence; acidic or basic nature of amino acids determines the isoelectric point of the sequence and grand average of hydropaticity index (GRAVY) measures whether the protein is hydrophobic (positive GRAVY) or hydrophilic (negative GRAVY).

The second group of features is based on the increasing use of document classification techniques for protein sequence classification (Cheng et al., 2005; Ganapathiraju et al., 2005), where characterization is done by counting the occurrences of all possible subsequences of a fixed length n , called n -grams or n -mers, over the primary structure of the protein. In this work, only features corresponding

Table 2.1: Definition and size of the classes. The list of GO terms covered by this analysis is intended to provide a complete landscape of GO predictability at the three levels of protein functionality in *Embryophyta* plants. For classification purposes, classes marked with an asterisk (*) were redefined. The number of samples in those categories corresponds to the sequences associated to that class and none of its also listed descendants.

Class	Acronym	Size	Class	Acronym	Size
Molecular Function			Biological Process		
Nucleotide binding	Ntbind	47	Reproduction*	Reprod*	337
Molecular function*	MF*	268	Carbohydrate metabolic process	ChMet	315
DNA binding	DnaBind	107	Generation of precursor metabolites and energy	MetEn	150
Transcription factor activity	TranscFact	307	Nucleobase, nucleoside, nucleotide, nucleic acid metabolic process*	NaMet*	712
RNA binding	RnaBind	43	DNA metabolic process	DnaMet	191
Catalytic activity*	Catal*	334	Translation	Transl	82
Receptor binding	RecBind	38	Protein modification process	ProtMod	391
Transporter activity	Transp	125	Lipid metabolic process	LipMet	324
Binding*	Bind*	173	Transport	Transport	531
Protein binding*	ProtBind*	630	Response to stress	StressResp	790
Kinase activity	Kinase	68	Cell cycle	CellCycle	234
Transferase activity*	Transf*	173	Cell communication*	CellComm*	66
Hydrolase activity	Hydrol	190	Signal transduction	SigTransd	305
Enzyme regulator activity	EnzReg	41	Cell-cell signaling	Cell-cell	53
Cellular Component			Multicellular organismal development*	MultDev*	490
Cellular component*	CC*	234	Biological process*	BP*	879
Extracellular region	ExtcellReg	109	Metabolic process*	Met*	279
Cell wall	CellWall	77	Cell death	CellDeath	95
Intracellular*	Intracell*	167	Catabolic process	Catabolic	479
Nucleus*	Nucleus*	421	Biosynthetic process*	Biosint*	1125
Nucleoplasm	NuclPlasm	51	Response to external stimulus*	ExtResp*	65
Nucleolus	Nucleolus	84	Tropism	Tropism	36
Cytoplasm*	CitPlasm*	168	Response to biotic stimulus	BioResp	275
Mitochondrion	Mitochond	244	Response to abiotic stimulus	AbioResp	642
Endosome	Endosome	58	Anatomical structure morphogenesis	StrMorph	366
Vacuole	Vacuole	171	Response to endogenous stimulus	EndoResp	332
Peroxisome	Peroxisome	32	Embryonic development	EmbDev	139
Endoplasmatic reticulum	EndRet	109	Post-embryonic development*	PostDev*	375
Golgi apparatus	GolgiApp	100	Pollination	Poll	43
Cytosol	Cytosol	389	Flower development	FlowerDev	228
Ribosome	Ribosome	98	Cellular process*	CP*	1486
Plasma membrane	PlasmMb	353	Response to extracellular stimulus	ExtcellResp	59
Plastid	Plastid	696	Photosynthesis	Photosyn	102
Thylakoid	Thylk	147	Cellular component organization	CellOrg	757
Membrane*	Mb*	472	Cell growth	CellGrowth	133
			Protein metabolic process*	ProtMet*	187
			Cellular homeostasis	CellHom	53
			Secondary metabolic process	SecMet	164
			Cell differentiation	CellDiff	267
			Growth*	Growth*	64
			Regulation of gene expression, epigenetic	RGE	103

Table 2.2: Initial set of features extracted from amino acid sequences. Features are divided into three broad categories: physical-chemical features, primary structure composition statistics and secondary structure composition statistics.

Nature	Description	Number
Physical-chemical	Sequence length	1
	Molecular weight	1
	Positively charged residues (%)	1
	Negatively charged residues (%)	1
	Isoelectric point	1
	GRAVY	1
Primary structure statistics	Amino acid frequencies	20
	Amino acid dimer frequencies	400
Secondary structure statistics	Structure frequencies	3
	Structural dimer frequencies	9
Total		438

to $n = \{1, 2\}$ are employed, provided that the size of the feature space grows exponentially with n . Simple countings were converted to relative frequencies (summing to one) in order to obtain information relative to the sequence composition more than simple presence statistics.

The third group is analog to the second one, but applying the n-gram characterization to the predicted secondary structure of the protein. Predictions were calculated by employing the Predator 2.1 software (Frishman and Argos, 1997), whose output is a linear sequence with three structural domains: alpha helices, beta sheets and coils. It is worthy to note that even though this group of features were computed over secondary structure information, they must be still considered as derived from primary structure because there is no full certainty about correctness of the secondary structure prediction.

It is important to point out that in the case of ambiguity characters in the sequence, that is, characters used to denote lack of information at certain positions (B for asparagine or aspartic acid, J for isoleucine or leucine, Z for glutamine or glutamic acid, X for any amino acid), each feature was computed as its statistical expected value with natural abundance percentages of amino acids as their prior probabilities (Buxbaum, 2007, chapter 1). Additionally, since different groups of features are very heterogeneously scaled (for instance, sequence length is ranged

Table 2.3: Description of the clusters of features with similar information content

Group	Main feature	Size	Group	Main feature	Size
1	Protein length	34	9	Proline	14
2	Negative charge / Acidic	8	10	Glutamine	35
3	Positive charge / Basic	30	11	Arginine	26
4	Alanine	10	12	Tryptophan	38
5	Cysteine	38	13	Tyrosine	35
6	Hidrophobic	46	14	Alpha helices	6
7	Histidine	29	15	Beta sheets	4
8	Asparagine / Methionine	85			

from 7 to 5138, while n-grams are just in the interval from 0 to 1), z-score normalization was performed so that each feature has a zero mean and unitary standard deviation.

2.2.4 Feature clusters

As a first step to perform an analysis of discriminant features for each GO term, features were hierarchically clustered into groups of similar information content. For this purpose, the Ward clustering algorithm was used, with absolute Pearson correlation distance as metric. This procedure yielded 15 clusters that are summarized in Table 2.3 and are used for assessing the influence of specific feature groups on the predictability of each category.

2.2.5 Feature selection strategy

The feature selection procedure was carried out before trying to induce any decision rule (classifier), because, having a limited number of training examples, excessive features (irrelevant or redundant) would possibly overfit the training data. For this purpose, an analysis of relevance and redundancy was applied.

Let ϕ_i , $i = 1, 2, \dots, N$, be the initial set of features, \mathbf{y} be the vector of labels, $c_{ij} = \text{cor}(\phi_i, \phi_j)$ be the linear correlation computed between any pair ϕ_i and ϕ_j and $c_{iy} = \text{cor}(\phi_i, \mathbf{y})$ be the linear correlation between ϕ_i and \mathbf{y} . Defining this, relevance of features can be quantified by computing c_{iy} for all features and then, redundant ones can be identified by analyzing the $N \times N$ feature

correlation matrix. In order to speed up the calculus, an algorithm based on the *Fast Correlation-Based Filter* of (Yu and Liu, 2004) was used.

First, an initial feature subset is selected based on a predefined threshold on c_{iy} . Then, the concept of *approximate Markov blanket* is used to select predominant features from this relevant set. As defined by (Yu and Liu, 2004), the feature ϕ_i forms an approximate Markov blanket for ϕ_j , ($i \neq j$) if and only if $c_{iy} \geq c_{jy}$ and $c_{ij} \geq c_{jy}$. This principle states that if the information shared between any pair of features is less or equal than information shared between one of them and the class labels, the one with lesser information could be discarded. Since the feature with highest correlation with the class labels does not have any approximate Markov blanket, it must be one of the predominant features and is taken as reference. The algorithm proceeds to compute correlation between the reference and all the remaining features by decreasing order of relevance; if the reference happens to form an approximate Markov blanket for some feature ϕ_i , the last is removed from the relevant features set. After one round of filtering features based on the first relevant one, the algorithm will take the remaining feature right next to it as the new reference to repeat the filtering process. The algorithm stops when no more predominant features can be selected.

2.2.6 Decision making

Support vector machines (SVM) were chosen as base classifiers for running all the supervised tests. SVMs are powerful tools for solving classification problems, designed over a strong theoretical background based on the idea of minimizing the structural risk (Vapnik, 1998). For a linear SVM, the objective is to find a classification function of the form:

$$f_{(\mathbf{w},b)}(\mathbf{x}) = \langle \mathbf{w}, \mathbf{x} \rangle + b \quad (2.1)$$

where $\langle \cdot, \cdot \rangle$ represents the dot product. Following the notation in section 1.4, a vector of parameters can be defined as $\boldsymbol{\theta} = [\mathbf{w}, b]$, and the optimization problem

can be stated as follows:

$$\boldsymbol{\theta}^* = \arg \min_{\boldsymbol{\theta} \in \mathcal{T}} \left\{ \frac{1}{2} \|\boldsymbol{\theta}\|^2 + C \sum_{i=1}^L \ell(f_{\boldsymbol{\theta}}(\mathbf{x}_i) y_i) \right\} \quad (2.2)$$

where $\ell(t) = \max(0, 1-t)$ is the hinge loss function and C is a trade-off parameter regulating the complexity of the model. For the non-linear case, the data are first mapped in a high dimensional Hilbert space \mathcal{H} through a mapping $\Phi : \mathcal{X} \mapsto \mathcal{H}$, and then a linear decision boundary is constructed in that space. The mapping Φ can be explicitly computed or only implicitly through the use of a kernel function K such that $K(\mathbf{x}_1, \mathbf{x}_2) = \langle \Phi(\mathbf{x}_1), \Phi(\mathbf{x}_2) \rangle$. The Representer Theorem can be used to show that the solution function has the form:

$$f_{\boldsymbol{\theta}^*}(\mathbf{x}) = \sum_{i=1}^L \alpha_i K(\mathbf{x}, \mathbf{x}_i) \quad (2.3)$$

where the coefficients α_i can be found with a conventional quadratic optimization algorithm. The Gaussian kernel is the most commonly used because of its attractive features such as structure preservation (Liu et al., 2012). This kernel is computed by:

$$K(\mathbf{x}_1, \mathbf{x}_2) = e^{-\frac{\|\mathbf{x}_1 - \mathbf{x}_2\|^2}{2\sigma}} \quad (2.4)$$

where σ is the dispersion parameter that must be properly chosen by the user. In this work, the SVM is trained with the 'kernlab' package, available in R-CRAN (Karatzoglou et al., 2004). Dispersion of the kernel and trade-off penalization parameter of the SVM are tuned for each test with a particle swarm optimization meta-heuristic, a bio-inspired optimization method that has been used in multiple applications in the past years (Kennedy and Eberhart, 1995).

In order to allow samples to be associated to multiple categories, decision making was implemented following the one-against-all strategy. The method produced a strong class imbalance since it trains a number of binary classifiers, each one intended to recognize samples from one class out of the whole training set. In other words, since all the proteins outside of the target GO term are seen as negative samples and there are only a relatively small number of proteins

annotated with the function, the number of negative samples may be hundreds and even thousands times the one of positive samples. Therefore, the classifier may be degraded by the false negative samples or imbalanced data (Zhao et al., 2008d). To overcome the problems that imbalanced classes commonly produce in pattern recognition techniques, the Synthetic Minority Over-sampling Technique (SMOTE) was employed (Chawla et al., 2002).

In order to estimate the performance of the predictive model, a 5-fold cross-validation strategy is implemented. In such strategy, the test procedure is repeated five times, and each time an 80% of the data is used for adjusting the SVM parameters and training the model, while the remaining 20% is used as testing samples. This strategy also allows providing an estimation of the reliability of the model by computing the variability of the results through the five repetitions.

2.3 Results and Discussion

2.3.1 Analysis of predictability with individual feature clusters

Classification results with individual feature clusters, for an identity cutoff of 30%, are condensed in Figure 2.1. The square root of the product between sensitivity and specificity (geometric mean), is depicted as global performance measure and the color scale has been adjusted to highlight the highest (green) and the lowest (blue) performance. Note that the rows and columns have been ordered to explicitly locate best predicted GO terms on top and most discriminant groups to the left.

Figure 2.1(a) shows the analysis for the molecular function ontology. For all feature groups, *Receptor binding* achieved the highest classification scores. This category is intended to comprise proteins that interact selectively and non-covalently with one or more specific sites on a receptor molecule. About 63% of the proteins associated to this category in the database are proteins involved with binding of serine/threonine kinase receptors, which turned out to be easily predicted from most of the defined features.

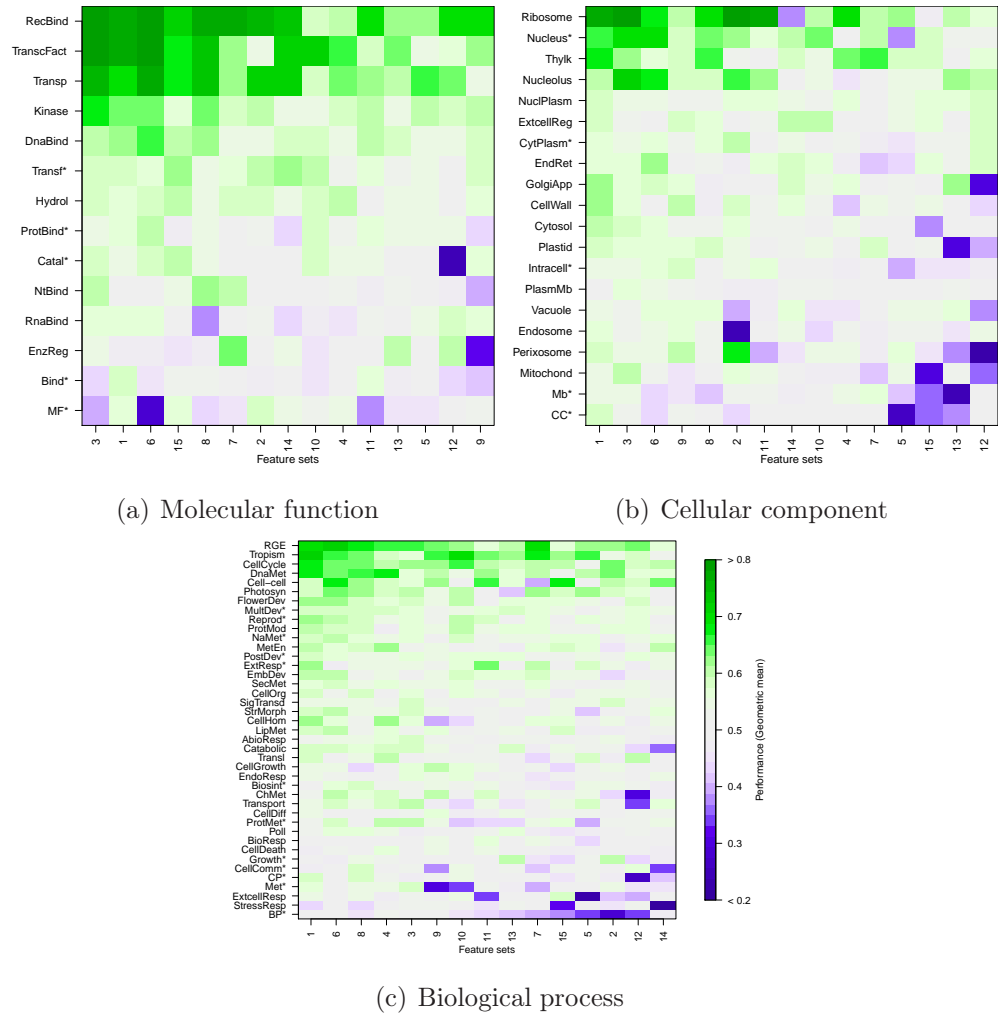


Figure 2.1: Prediction performance with different feature clusters. Rows represent classes in Table 2.1 while columns represent feature groups in Table 2.3. For each ontology, best predicted categories are ordered from top to bottom while most discriminant feature groups are ordered from left to right.

Transcription factor activity achieved was easily predicted from the feature groups 1, 3, 6, 8 and 14. Not so surprising is the fact that *DNA binding* also presents a similar behavior since most transcription factors must interact with DNA molecules and consequently they are also included in this category. However it is worthy to note that several other proteins also perform DNA cleavage, such as polymerases, nucleases and histones, and they were also well predicted from the same feature groups. The conclusion from these results becomes more evident by observing the results of the *DNA metabolic process* in Figure 2.1(c), which confirm the high predictability of all proteins involved with transcription when using the mentioned features groups. A similar behavior is also observed for *nucleus** in Figure 2.1(b), supported by the fact that the transcription process is mostly carried out in that sub-cellular location.

Transporter activity refers to proteins that enable the directed movement of substances into, out of, within or between cells. Most of them are integral transmembrane proteins, that are distinguished by their high content of hydrophobic residues (Whitford, 2005). In fact, some of the highest performance of *transporter activity* were reached with the groups 3 and 6, which include GRAVY index as well as monomer and dimer frequencies of three out of the four most hydrophobic residues: leucine, isoleucine and phenylalanine. Additionally, predictability of this molecular function is reflected, while in a minor degree, on the *transport* biological process, which reaches its highest values for the same feature groups (see Figure 2.1(c)). The main difference between those GO terms lies in that *transport* is a broader category, including external agents such as oxygen carriers and lipoproteins that perform transport within multicellular organisms.

On the other hand, the root node of the molecular function ontology was GO terms with the lowest average prediction performance. Remember that the root node contains the proteins that do not belong to any of its descendant categories, so it keeps a small set of sequences of a sparse nature, which explains the impossibility to model and predict them as a group. It is interesting to note that the same behavior is observed for the other two ontologies (figures 2.1(b) and 2.1(c)).

Concerning the cellular component ontology, it can be observed in Figure 2.1(b) that *ribosome* category has reached the highest classification accuracies,

specially with groups 1, 2, 3 and 11. Such groups mainly consist of the four charged residues: lysine, arginine, glutamic acid and aspartic acid. This can be explained since ribosomal proteins must interact with the negatively charged phosphodiester bonds in the RNA backbone, so they are expected to have a high percentage of positively charged residues to neutralize such charge repulsion. In agreement with this, (Whitford, 2005) describes the composition of isolated ribosomal proteins as showing a high percentage of lysine and arginine residues and a low aromatic content. Hence, there is enough evidence to establish that ribosomal proteins are another highly predictable category from primary structure information.

As explained before, *nucleus** becomes easily predicted from the same feature groups that have shown high discriminant capabilities for transcription related proteins. A similar behavior is also observed for proteins belonging to the *nucleolus* component, which encompasses proteins including RNA polymerases, transcription factors, processing enzymes and ribosomal proteins among others, which must interact with nucleic acids and have shown low isoelectric points in comparison to the remaining proteins in the database.

Thylakoid proteins also presented high prediction performance with several feature groups. Further studies would be required to explain this results.

Broad categories such as *membrane** showed poor performance with most feature groups, presumably due to its high diversity. However, some rather well-defined categories such as *mitochondrion* and *perixosome* were also ranked in the lowest places in figure 2.1(b), simply proving to be poorly predictable from the extracted feature groups.

Concerning figure 2.1(c), the biological process that was better predicted for most group features is *regulation of gene expression, epigenetic*. This GO term encloses proteins involved in modulating the frequency, rate or extent of gene expression and is highly composed by histones. In fact, since histones are highly alkaline proteins, it is consistent to observe that this category became particularly well predicted from groups 3, 6 and 7, which are mainly conformed by frequencies of phenylalanine, leucine, isoleucine, lysine and histidine residues. Also, cysteine related frequencies were highly discriminant for *regulation of gene expression, epigenetic* (group 5 which can be explained by the fact that altering the redox

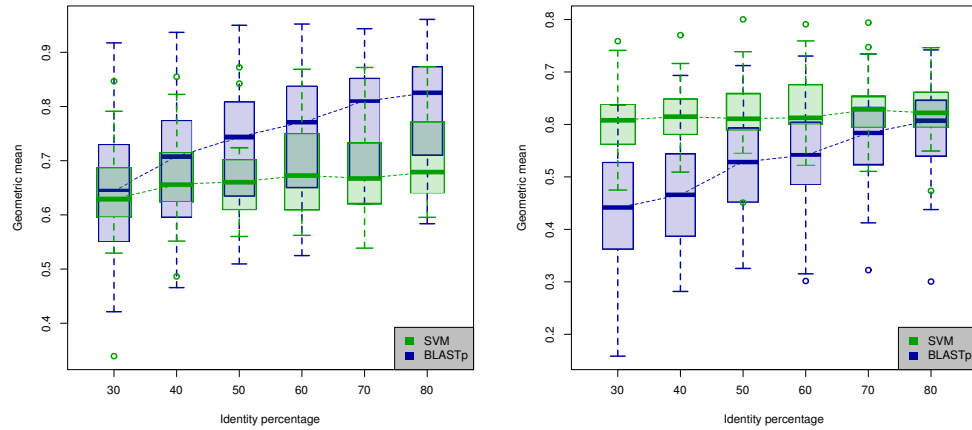
state of cysteines serves for modulating protein activity, and several transcription factors become activated by the oxidation of cysteines that form disulfide bonds (Arrigo, 1999).

Tropism and *Cell Cycle* also appeared near the top of figure 2.1(c), just before *DNA metabolic process* which was already discussed.

2.3.2 Analysis of predictability with the full set of features

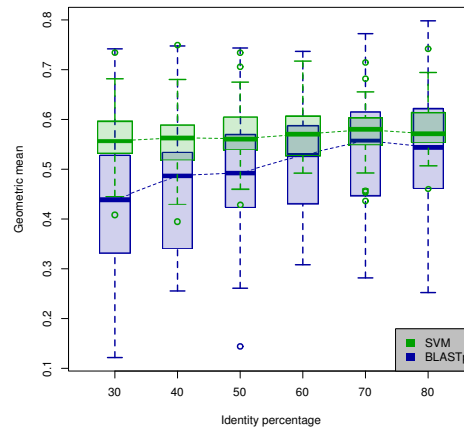
Analyses in the previous section were done after discarding sequences with identities superior to 30%. Otherwise, the predictability of certain terms could be enhanced from the fact that many proteins in training and testing sets are copies (or close relatives) from another, rather than from predictive value of certain sequence-derived features. However, in order to provide further information on the robustness of the proposed methodology when the identity cutoff changes, Figure 2.2 presents an analysis of predictability with the full feature set (although applying the feature selection procedure described in the methods section), while varying the identity cutoff. For comparison purposes, results achieved by BLASTP are depicted in blue, while results of the proposed methodology are depicted in green. The first thing that can be noted from Figure 2.2 is the fact that alignment-based predictions are more sensitive to the variation of the identity percentage than the proposed methodology. It can be clearly seen that BLASTP suffers a strong performance degradation as the identity filter is more stringent, while the performance of the proposed methodology remains more stable. Moreover, although in Figure 2.2(a) it can be seen that, when predicting molecular functions, BLASTP is superior than the proposed methodology for high identity cutoffs, the difference at 30% is not statistically significant. Conversely, the proposed methodology clearly outperforms BLASTP for low identity percentages when predicting cellular components and biological processes (figures 2.2(b) and 2.2(c)).

Figure 2.3 depicts detailed results of predicting each class with the full feature set for an identity cutoff of 30%. Left plots show sensitivity, specificity and geometric mean (green line) achieved with the five-fold cross-validation procedure, as well as the performance of the BLASTP algorithm for comparison purposes (blue



(a) Molecular function

(b) Cellular component



(c) Biological process

Figure 2.2: Performance variation in function of the identity cutoff. Green and blue plots show the variation of the general prediction performance for SVM and BLASTp, respectively, according to the identity percentage cutoff used in the dataset. Boxplots show the dispersion throughout the 75 GO terms.

line). Right plots depicts the corresponding p-values obtained from a paired t-test between the BLASTP method and the SVM. The dashed black line is located at 0.95 and the colored bars indicate which values are statistically significant at a 95% significance level. Green bars show the cases when the SVM method is significantly better than BLASTP, while blue bars show otherwise.

Note that GO terms were ordered again from top to bottom according to their predictability, but this order is not strictly the same as in Figure 2.1. Some interesting results in figure 2.3(b) are provided by categories such as *plastid*, which was not easily predicted with any feature set independently, but reached medium to high classification results when the complete set was used. Such behavior is a clear example of the multivariate associations that could be missed when analyzing only individual feature sets.

Other results were consistent with the insights provided by the previous analyses, showing that some of the best predicted GO terms were *transporter activity*, *transcription factor activity*, and *DNA binding* in molecular functions; *ribosome*, *nucleus**, *nucleolus* and *thylakoid* in cellular components; *regulation of gene expression*, *epigenetic*, *Cell cycle*, *Photosyntesis* and *DNA metabolic process* in biological processes.

A reduced number of categories had performance under 50%, most of them from the biological process ontology and a few from the molecular function ontology. It is important to note that the majority of those categories achieved very high specificities and low sensitivities, pointing out to a high dispersion of such categories over the feature space, which yields to a very high number of false negatives. Also, the high dispersions observed in the boxplots for some of the worst predicted classes demonstrate that there is a high variability among repetitions of the experiment which means that those low performance are not confident. Conversely, the categories with high performance show also low dispersions associated to them, hinting consistency in the predictors.

Although the main purpose of this chapter is not to design a highly accurate GO term predictor, but to provide a comprehensive analysis of the predictability of GO terms from primary structure information, it is important to mention how this method compares with currently used prediction tools. The blue and green lines in Figure 2.3 represent the prediction performance of BLASTP and the SVM

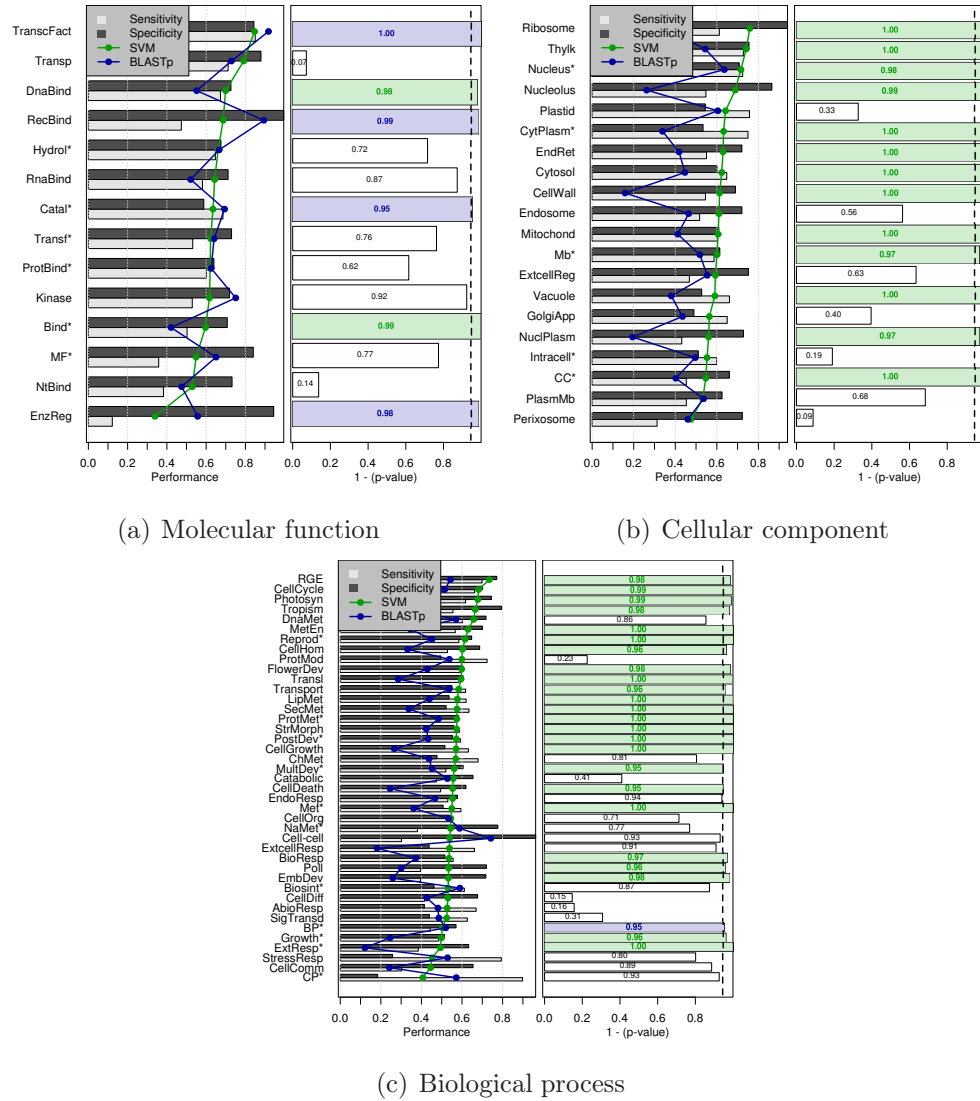


Figure 2.3: Prediction performance with the complete set of features. Bars in the left plots show sensitivity and specificity of SVMs. Lines depict geometric mean as a global performance measure for SVM (green) and BLASTP (blue). Right plots depict the p-values obtained by paired t-tests at a 95% significance level. For each ontology, the best predicted categories are ordered from top to bottom.

based predictor used in this work, respectively. Both methods were tested over the same database described in the methods section. From Figure 2.3(a) it is possible to conclude that the two methods provide similar prediction capabilities for the molecular function ontology at this identity cutoff, with the SVM significantly outperforming BLASTP only for two molecular functions, while the opposite case occurs for three cases. In eight out of fourteen molecular functions, there is no statistically significant difference between the methods. On the other hand, figures 2.3(b) and 2.3(c) show that the SVM out-performed BLASTP for most cellular components and biological processes, with only a few exceptions. It is also important to point out that the results achieved here are competitive with those reported by (Chou and Shen, 2010), which is one of the more recent and effective predictors dedicated to plant proteins.

Finally, Figure 2.4 depicts the accuracy obtained in each category, when predictions of inferior GO terms were propagated up to their parents. Observe that asterisks have been removed to point out that GO terms are now including all their descendants.

It is notable how categories with the major number of descendants have been negatively affected by their false positives. This is especially observed in Figure 2.4(b) for *cytoplasm*, and *intracellular*, and Figure 2.4(c) for *cellular process* and *metabolic process*. Conversely, a few classes that were lacking sensitivity were favored by the contributions of their descendants, as it is the case of the root nodes of the ontologies.

2.4 Concluding remarks

An analysis of GO terms predictability in land plants proteins was carried out in order to determine single categories or groups of related functions that are more related with primary structure information. For this purpose, pattern recognition techniques were employed over a feature set of physical-chemical and statistical attributes computed over the primary structure of the proteins. High predictability of several GO terms was observed in the three ontologies. Proteins associated to transport activities showed high correct prediction rates when using hydropathicity related features. Also, proteins involved with transcription (and



Figure 2.4: Propagated prediction performance. Prediction performance when propagating predictions of children nodes to their parents. Note that asterisks in the category names have been removed since categories include all their member now.

therefore associated to the nucleus) presented high discriminability from the extracted features. Ribosomal and other proteins involved with translation, proved to be highly predictable from features related to electric charges of the amino acid sequence. At the biological process level, proteins related to regulation of gene expression and nucleic acid metabolic process were easily predicted, while some other biological processes showed low predictability from the extracted primary structure features.

Chapter 3

Semi-supervised Gene Ontology prediction for *Embryophyta* organisms

Supervised machine learning uses a labeled set of instances to train the classifier. Labeled instances, however, are often difficult, expensive, or time consuming to obtain, as they require the efforts of experienced human annotators. Meanwhile unlabeled data may be relatively easy to collect, but there has been few ways to use them ([Zhu, 2007](#)).

In the particular case of protein function prediction, it is a known fact that only a small number of proteins have actually been annotated for certain functions. Therefore, it is difficult to obtain sufficient training data for the supervised learning algorithms and, consequently, the tools for protein function prediction have very limited scopes ([Zhao et al., 2008c](#)). Besides, it is particularly hard to find the representative negative samples because the available information in the annotation databases, such as Gene Ontology (GO) ([The Gene Ontology Consortium, 2004](#)), only provides information about which protein belongs to which functional class but there is no certainty about which protein does not belong to the class ([Zhao et al., 2008a](#)). To see this, consider the functional path of a GO term as the path on the ontology from the root to the node representing that GO term. Since the current functional annotation might be incomplete, it is hard to

justify whether or not the proteins annotated with the nodes on the functional paths of a GO node (the ancestors of the term) would belong to this GO node if a deeper study was performed. Therefore these proteins must only be considered as unlabeled samples regarding that GO term (Bi et al., 2007).

Under such circumstances, semi-supervised learning methods provide an alternative approach to protein annotation (Zhao et al., 2008c). Semi-supervised learning (SSL) is halfway between supervised and unsupervised learning: in addition to labeled data, the algorithm is provided with an amount of unlabeled data that can be used to improve the estimations about the data. This chapter presents an analysis of the predictability of GO terms with semi-supervised learning methods, also providing an analysis of the assumptions that the different methods do, in order to understand their successfulness.

3.1 State of the art in semi-supervised classification

The idea of using both labeled and unlabeled data for designing robust predictors has been on the machine learning community since around the middle sixties with several proposals on self-training, where a supervised learning method is iteratively re-trained with its own predictions over a set of unlabeled data for successively refining the final output (see, for example, Scudder III (1965) and Fralick (1967)). Later, the work in semi-supervised learning moved to the generative models, considering that each class has a Gaussian distribution. This is equivalent to assuming that the complete data comes from a mixture model and, with large amount of unlabeled data, the mixture components can be identified with the expectation-maximization (EM) algorithm (McLachlan and Krishnan, 2007; Miller and Uyar, 1996). The interest in semi-supervised learning increased in the nineties, mostly due to applications in natural language problems and text classification (Blum and Mitchell, 1998; Joachims, 1999). According to Chapelle and Schölkopf (2006), the work by Merz et al. (1992) was the first to use the term “semi-supervised” for classification with both labeled and unlabeled data.

A more complete historical perspective of semi-supervised learning can be found on (Chapelle and Schölkopf, 2006, chapter 1).

One significant difference between supervised and semi-supervised methods is that, unlike supervised learning, in which a good generic learning algorithm can perform well on a lot of real-world data sets without specific domain knowledge, in semi-supervised learning there is commonly accepted that there is no “black box” solution and a good understanding of the nature of the data is required to achieve successful performance (Chapelle and Schölkopf, 2006, Chapter 21). This is mainly due to the fact that, in order to deal with labeled and unlabeled data, current semi-supervised algorithms have to make strong assumptions about the underlying joint probability measure $P(\mathcal{X}, \mathcal{Y})$ e.g. a relation of the probability of the feature space $P(\mathcal{X})$ and the joint probability of the feature space and the label set $P(\mathcal{Y}, \mathcal{X})$. There are several different semi-supervised learning methods, and each one makes different assumptions about this link. These methods include generative models, graph-based models, semi-supervised support vector machines, and so on (Zhu and Goldberg, 2009).

The main assumption made by semi-supervised learning algorithms is the “semi-supervised smoothness assumption” (Chapelle and Schölkopf, 2006, chapter 1):

Semi-supervised smoothness assumption: If two points \mathbf{x}_1 , and \mathbf{x}_2 in a high-density region are close, then so should be their corresponding label sets \mathbf{y}_1 , \mathbf{y}_2 . Note that by transitivity, this assumption implies that if two points are linked by a path of high density (e.g., if they belong to the same cluster), then their outputs are likely to be close. If, on the other hand, they are separated by a low-density region, then their outputs need not be close.

Such assumption originates the two common assumptions used in semi-supervised learning:

Cluster assumption: If points are in the same cluster, they are likely to be of the same class. This assumption does not imply that each class forms a single, compact cluster, it only means that there are no instances of

two distinct classes in the same cluster. The cluster assumption can be formulated in an equivalent way:

Low density separation: The decision boundary should lie in a low-density region.

Manifold assumption: The (high-dimensional) data lie (roughly) on a low-dimensional manifold. Instances that are close according to the manifold geodesic distance are likely to be of the same class.

According to each assumption, there are three main families of semi-supervised methods: generative methods (cluster assumption), density-based methods (low density separation), and graph-based methods (manifold assumption). In the following sub-sections each family of methods will be reviewed, emphasizing on the assumptions made by each one. It should be pointed out that, since semi-supervised learning is a rapidly evolving field, the review is necessarily incomplete. A wider review in this matter can also be found on ([Zhu and Goldberg, 2009](#)).

3.1.1 Generative methods

Generative methods follow a common strategy of augmenting the set of labeled samples with a large set of unlabeled data and combining the two sets with the Expectation-Maximization algorithm, in order to improve the parameter estimates [Cozman et al. \(2003\)](#). They assume a probabilistic model $p(\mathbf{x}, y) = p(y)p(\mathbf{x}|y)$, where $p(\mathbf{x}|y)$ is an identifiable mixture distribution. The most commonly employed distributions are the Gaussian Mixture Models:

$$p(\mathbf{x}|y) = \sum_{k=1}^K \pi_k \mathcal{N}(\mathbf{x}|\boldsymbol{\theta}) \quad (3.1)$$

where $\mathcal{N}(\mathbf{x}|\boldsymbol{\theta})$ is the gaussian distribution with parameters $\boldsymbol{\theta} = [\boldsymbol{\mu}_k, \boldsymbol{\Sigma}_k]$, being $\boldsymbol{\mu}_k$ the mean vector and $\boldsymbol{\Sigma}_k$ the covariance matrix of the k -th Gaussian component, and π_k the mixing components such that $\sum_{k=1}^K \pi_k = 1$ for $k = 1, 2, \dots, K$.

Ideally only one labeled example per component is needed to fully determine the mixture distribution. In this setting, any additional information on $p(\mathbf{x})$

is useful and the EM algorithm can be used for estimating θ . A strength of the generative approach is that knowledge of the structure of the problem or the data can naturally be incorporated by modeling it (Chapelle and Schölkopf, 2006, chapter 1). However, generative techniques provide an estimate of $p(\mathbf{x})$ along the way, although this is not required for classification, and in general this proves wasteful given limited data. For example, maximizing the joint likelihood of a finite sample need not lead to a small classification error, because depending on the model it may be possible to increase the likelihood more by improving the fit of $p(\mathbf{x})$ than the fit of $p(y|\mathbf{x})$ (Chapelle and Schölkopf, 2006, chapter 2).

The aforementioned works of Miller and Uyar (1996) and McLachlan and Krishnan (2007), among others, shown to be strong methods for classifying text data. Also, Nigam et al. (2000) have applied the EM algorithm on mixture of multinomial for the task of text classification, showing better performance than those trained only from the supervised set. Fujino et al. (2005) extend generative mixture models by including a “bias correction” term and discriminative training using the maximum entropy principle. However, anecdotal evidence is that many more studies were not published because they obtained negative results, showing that learning a mixture model will often degrade the performance of a model fit using only the labeled data (Zhu and Lafferty, 2005); one published study with these conclusions is Cozman et al. (2003). This is due to the strong assumption done by generative methods: that the data actually comes from the mixture model, where the number of components, prior $p(y)$, and conditional $p(\mathbf{x}|y)$ are all correct (Zhu and Goldberg, 2009).

3.1.2 Density-based methods

With the rising popularity of support vector machines (SVMs), Semi-Supervised SVMs (S³VMs) emerged as an extension to standard SVMs for semi-supervised learning. S³VMs find a labeling for all the unlabeled data, and a separating hyperplane, such that maximum margin is achieved on both the labeled data and the (now labeled) unlabeled data. As a result, unlabeled data guides the decision boundary away from dense regions. The assumption of S³VMs is that the classes are well-separated, such that the decision boundary falls into a low density region

in the feature space, and does not cut through dense unlabeled data (Zhu and Goldberg, 2009, chapter 6).

In a similar way than the conventional SVMs described in section 4.1.2, the optimization problem for an S³VMs can be stated as follows:

$$\boldsymbol{\theta}^* = \arg \min_{\boldsymbol{\theta} \in \mathcal{T}} \left\{ \frac{1}{2} \|\boldsymbol{\theta}\|^2 + C \sum_{i=1}^L \ell(f_{\boldsymbol{\theta}}(\mathbf{x}_i)y_i) + \lambda \sum_{i=L+1}^{L+U} \ell(|f_{\boldsymbol{\theta}}(\mathbf{x}_i)|) \right\} \quad (3.2)$$

where $\ell(t) = \max(0, 1 - t)$ is the hinge loss function, C is the trade-off parameter and λ is a new regularization parameter. The first two terms in the above equation correspond to the traditional solution for the standard supervised SVM shown in equation (2.2), while the last term puts $f_{\boldsymbol{\theta}}(\mathbf{x}_i)$ of the unlabeled points \mathbf{x}_i away from 0 (thereby implementing the low density assumption) (Chapelle et al., 2006).

Again, as in the supervised case, the kernel trick can be used for constructing non-linear S³VMs. While the optimization in SVM is convex and can be solved with QP-hard complexity, optimization in S³VM is a non-convex combinatorial task with NP-Hard complexity. Most of the recent work in S³VM has been focused on the optimization procedure (a full survey in this matter can be found in (Chapelle et al., 2008)). Among the proposed methods for solving the non-convex optimization problem associated with S³VMs, one of the first implementations is the S³VM^{light} by (Joachims, 1999), which is based on local combinatorial search guided by a label switching procedure. Chapelle and Zien (2005) presented a method based on gradient descent on the primal, that performs significantly better than the optimization strategy pursued in S³VM^{light}; the work by Chapelle et al. (2006) proposes the use of a global optimization technique known as “continuation”, often leading to lower test errors than other optimization algorithms; Collobert et al. (2006) uses the Concave-Convex procedure, providing a highly scalable algorithm in the nonlinear case.

Other recent proposals include (Li et al., 2010) which focuses on the class-imbalance problem and proposes a cost-sensitive S³VM; Qi et al. (2012) which describes laplacian twin support vector machines; and several approaches to adaptive regularizations like (Xu et al., 2009) and (Wang et al., 2011).

3.1.3 Graph-based methods

Graph-based methods start with a graph where the nodes are the labeled and unlabeled data points, and (weighted) edges reflect the similarity of nodes. The assumption is that nodes connected by a large-weight edge tend to have the same label, and labels can propagate throughout the graph. In other words, graph-based methods do the assumption that labels are “smooth” with respect to the graph, such that they vary slowly on the graph. That is, if two instances are connected by a strong edge, their labels tend to be the same (Zhu and Goldberg, 2009, chapter 5).

This family of methods enjoy nice properties from spectral graph theory. They commonly use an energy function as objective in the optimization problem, ensuring that the labels will change slowly through the graph (consequently implementing the manifold assumption) (Hein et al., 2005). Some common graphs include the following (Zhu and Goldberg, 2009, chapter 5):

Fully connected graphs: The graph needs to be weighted so that similar nodes have large edge weight between them. The disadvantage is in computational cost as the graph is dense. A common choice in this case is the “exp-weighted graph” where the weights between instances \mathbf{x}_i and \mathbf{x}_j are defined as:

$$W_{ij} = \exp\left(-\frac{\|\mathbf{x}_i - \mathbf{x}_j\|^2}{2\alpha}\right) \quad (3.3)$$

where α is a bandwidth hyperparameter that controls the decay rate.

Sparse graphs: In this kind of graphs each node connects to only a few nodes, making them computationally fast. They also tend to enjoy good empirical performance. There are two common choices: k -NN graphs and ϵ -NN graphs. In the former, instances \mathbf{x}_i and \mathbf{x}_j are connected by an edge if any of them are included into the k nearest neighbors of the other. k is a hyperparameter that controls the density of the graph. Small k may result in disconnected graphs. For ϵ -NN graphs, \mathbf{x}_i and \mathbf{x}_j are connected by an edge, if the distance $\|\mathbf{x}_i - \mathbf{x}_j\| \leq \epsilon$. The hyperparameter ϵ controls neighborhood radius.

A graph is represented by the $(L + U) \times (L + U)$ weight matrix \mathbf{W} , $W_{ij} = 0$ if there is no edge between instances \mathbf{x}_i and \mathbf{x}_j . Once the graph has been defined, a real function over the nodes can be defined $f_{\boldsymbol{\theta}} : \mathcal{X} \rightarrow \mathbb{R}$. In order to achieve that unlabeled points that are similar (as determined by the edges of the graph) to have similar labels, the quadratic energy function can be used as objective:

$$\boldsymbol{\theta}^* = \arg \min_{\boldsymbol{\theta} \in \mathcal{T}} \left\{ \frac{1}{2} \sum_{ij} W_{ij} (f_{\boldsymbol{\theta}}(\mathbf{x}_i) - f_{\boldsymbol{\theta}}(\mathbf{x}_j))^2 \right\} \quad (3.4)$$

Since this objective function is minimized by constant functions, it is necessary to constrain $f_{\boldsymbol{\theta}}$ to take values $f_{\boldsymbol{\theta}}(\mathbf{x}_i) = y_i$, for all the labeled data $\mathbf{x}_i \in \mathcal{X}_L$. Finally, let \mathbf{D} be the diagonal degree matrix, where $D_{ii} = \sum_j \mathbf{W}_{ij}$ is the degree of node \mathbf{x}_i . The combinatorial Laplacian Δ is defined as:

$$\Delta \equiv \mathbf{D} - \mathbf{W} \quad (3.5)$$

and it is easy to verify that:

$$\boldsymbol{\theta}^* = \arg \min_{\boldsymbol{\theta} \in \mathcal{T}} \{ \mathbf{f}_{\boldsymbol{\theta}}^T \Delta \mathbf{f}_{\boldsymbol{\theta}} \} \quad (3.6)$$

There are many related methods that exploit the idea of obtaining a target function being smooth on the graph. Such methods include, among others, the Mincut ([Blum and Chawla](#)), which partitions the graph for minimizing the number of pairs of linked instances with different labels; graph random walks ([Azran, 2007](#)); harmonic functions ([Zhu and Lafferty, 2005](#)); local and global consistency ([Zhou et al., 2004](#)) and manifold regularization ([Belkin et al., 2006](#); [Sindhwani et al., 2006](#)). Some more recent proposals have focused on the problem of large graph construction, like [Liu et al. \(2010\)](#) and [Lin \(2012\)](#).

Most graph-based methods are inherently transductive, giving predictions for only those points in the unlabeled set, and not for an arbitrary test point. The simplest strategy for extending the method for unseen data is by dividing the input space into Voronoi cells centered on the labeled instances. From an algorithmic point of view, this strategy is equal to classify instances by its 1-nearest-neighbor. [Zhu and Lafferty \(2005\)](#) proposed an approach that combines generative mixture models and discriminative regularization using the graph Laplacian

in order to provide an inductive model. Laplacian SVMs, proposed by [Belkin et al. \(2006\)](#), provides a natural inductive algorithm since it uses a modified SVM for classification. The optimization problem in this case is regularized by the introduction of a term for controlling the complexity of the model according to equation (3.6):

$$\theta^* = \arg \min_{\theta \in \mathcal{T}} \left\{ \sum_i \ell(f_{\theta}(\mathbf{x}_i)y_i) + \lambda \sum_{ij} W_{ij} (f_{\theta}(\mathbf{x}_i) - f_{\theta}(\mathbf{x}_j))^2 \right\} \quad (3.7)$$

where W_{ij} is the weight between the i -th and j -th instances in the graph and λ is again a regularization parameter. A lot of experiments show that Laplacian SVM achieves state of the art performance in graph-based semi-supervised classification [Qi et al. \(2012\)](#).

3.1.4 Applications of semi-supervised learning for protein function prediction

A few semi-supervised methods have been applied for both gene function prediction (over the DNA sequence) and protein function prediction (over the amino acids sequence). [Kasabov and Pang \(2003\)](#) used a S³VMs for promoter recognition, improving predictive performance by 55% over the standard inductive SVM results. [Li et al. \(2003\)](#) used a “co-updating” schema of two SVMs, each one trained over a different source of data, for discriminating among five functional classes in the yeast genome. For the problem of predicting the functional properties of proteins, [Krogel and Scheffer \(2004\)](#) conducted an extensive study on the caveats of incorporating semi-supervised learning and transduction for predicting various functional properties of proteins corresponding to genes in the yeast genome, founding that S³VMs significantly decrease performance compared to inductive SVMs. [Shin and Tsuda \(2006\)](#) used graph-based semi-supervised learning for functional class prediction of yeast proteins, using protein interaction networks for obtaining the graphs.

More recently, [King and Guda \(2008\)](#) proposes a generative semi-supervised method for protein functional classification and provide experimental results of

classifying a set of eukaryotic proteins into seven subcellular locations from the Cellular Component ontology of GO. Zhao et al. (2008b) proposed a new algorithm to define the negative samples in protein function prediction. In detail, the one-class SVMs and two-class SVMs are used as the core learning algorithm in order to identify the representative negative samples so that the positive samples hidden in the unlabeled data can be recovered. Shin et al. (2009) propose a method for integrating multiple graphs within a framework of semi-supervised learning and apply the method to the task of protein functional class prediction in yeast. The proposed method performs significantly better than the same algorithm trained on any single graph.

3.2 Proposed methodology: semi-supervised learning for predicting gene ontology terms in *Embryophyta* plants

3.2.1 Selected semi-supervised algorithms

In order to test the efficiency of semi-supervised learning in the task of predicting protein functions, two state of the art methods were chosen, each one implementing a different semi-supervised assumption: S³VM following the concave-convex optimization procedure (CCP) (Collobert et al., 2006) (implementing the low-density separation assumption and consequently the cluster assumption) and Laplacian-SVM Belkin et al. (2006) (implementing the manifold assumption).

CCP S³VM: The S³VM proposed by (Collobert et al., 2006; Sinz et al., 2007) was chosen since it provides high scalability in the nonlinear case, making it the most suitable choice for the amounts of *Embryophyta* proteins in the databases used in this work. Consider the set of labeled points $\mathcal{X}_L = \{\mathbf{x}_i\}_{i=1}^L$ for which labels $\{y_i\}_{i=1}^L$ are provided, and the points $\mathcal{X}_U = \{\mathbf{x}_i\}_{i=L+1}^{L+U}$ the labels of which are not known. The objective function to be optimized in

this case, corresponds to:

$$J_{S^3VM}(\boldsymbol{\theta}) = \frac{1}{2} \|\boldsymbol{\theta}\|^2 + C \sum_{i=1}^L \ell(f_{\boldsymbol{\theta}}(\mathbf{x}_i)y_i) + \lambda \sum_{i=L+1}^{L+U} \ell(|f_{\boldsymbol{\theta}}(\mathbf{x}_i)|) \quad (3.8)$$

where the function $\ell(t) = \max(0, 1 - |t|)$ is the hinge loss function. The main problem with this objective function, in contrast to the classical SVM objective, is that the additional term is non-convex and gives rise to local minima. Additionally, it has been experimentally observed that the objective function tends to give unbalanced solutions, classifying all the unlabeled points in the same class. A constraint should be imposed on the data to avoid this problem ([Chapelle and Zien, 2005](#)):

$$\frac{1}{L} \sum_{i=1}^L y_i = \frac{1}{U} \sum_{i=L+1}^{L+U} f_{\boldsymbol{\theta}}(\mathbf{x}_i) \quad (3.9)$$

which ensures that the number of unlabeled samples assigned to each class will be the same fraction as in the labeled data. CCP decomposes a non-convex function J into a convex component J_{vex} and a concave component J_{cave} . At each iteration, the concave part is replaced by the tangential approximation at the current point and the sum of this linear function and the convex part is minimized to get the next iterate. The first two terms in equation (3.8) are convex, while the third term can be decomposed into the sum of a convex function plus a concave one:

$$J_{vex} = \max(0, 1 - |t|) + 2|t| \quad (3.10)$$

$$J_{cave} = -2|t| \quad (3.11)$$

If an unlabeled point is currently classified positive, then at the next itera-

Algorithm 1 CCP for S³VM**Require:** Initial θ from the supervised SVM**while** convergence of y_i is not met **do** $y_i \leftarrow f_{\theta}(\mathbf{x}_i), i = L + 1, L + 2, \dots, L + U$

$$\theta = \arg \min \left\{ \frac{1}{2} \|\theta\|^2 + C \sum_{i=1}^L \ell(f_{\theta}(\mathbf{x}_i)y_i) + \lambda \sum_{i=L+1}^{L+U} \tilde{\ell}(f_{\theta}(\mathbf{x}_i)y_i) \right\}$$

end while**return** θ

tion, the convex loss on this point will be:

$$\tilde{\ell}(t) = \begin{cases} 0 & \text{if } t \geq 1, \\ (1 - t) & \text{if } |t| < 1, \\ -4t & \text{if } t \leq -1 \end{cases} \quad (3.12)$$

The CCP algorithm for the semi-supervised support vector machines is presented in Algorithm 1.

Laplacian SVM: Regarding the graph-based algorithms, Laplacian support vector machines (Lap-SVM) were chosen since, according to Qi et al. (2012), many experiments show that Lap-SVM achieves state of the art performance among graph-based semi-supervised classification methods. This method, as proposed in Belkin et al. (2006), uses an objective function that is slightly different to equation (3.7), that is:

$$J_{LapSVM}(\theta) = \frac{1}{L} \sum_{i=1}^L \ell(f_{\theta}(\mathbf{x}_i)y_i) + \lambda_A \|\theta\|^2 + \frac{\lambda_I}{(L + U)^2} \mathbf{f}_{\theta}^T \Delta \mathbf{f}_{\theta} \quad (3.13)$$

where λ_A and λ_I are two regularizing constants that must be set by the user. Belkin et al. (2006), also demonstrated a modified version of the Representer Theorem that ensures that the solution function can be given again by linear combination of kernel functions as in equation (2.3), and the Lap-SVMs can be implemented by using a standard SVM quadratic solver.

3.2.2 Databases

Before proceeding with the prediction of protein functions, a set of three benchmark problems constructed by [Chapelle and Schölkopf \(2006\)](#) was used to test the algorithms in order to provide an analysis of the relation between the performance of the semi-supervised learning methods and the different dataset structures. The benchmark were artificially created in order to create situations that correspond to certain assumptions. The three datasets have 750 instances, from which 100 are labeled while the remaining are unlabeled.

g241n This data set was constructed to have potentially misleading cluster structure, and no manifold structure. It has 375 points drawn from each of two unit-variance isotropic Gaussians, the centers of which have a distance of 6 in a random direction; these points form the positive class. Then the centers of two further Gaussians for the negative class were fixed by moving from each of the former centers a distance of 2.5 in a random direction. The identity matrix was used as covariance matrix, and 375 points were sampled from each new Gaussian. A two-dimensional projection of the resulting data is shown on [Figure 3.1\(a\)](#).

g241c This data set accomplishes the cluster assumption, that is, the classes correspond to clusters but the manifold assumption does not hold. It has 750 points drawn from each of two unit-variance isotropic Gaussians, the centers of which had a distance of 2.5 in a random direction. The class label of a point represents the Gaussian it was drawn from. All dimensions are standardized (shifted and rescaled to zero-mean and unit variance). A two-dimensional projection of the resulting data is shown on [Figure 3.1\(b\)](#).

Digit1 This data set was designed to consist of points close to a low-dimensional manifold embedded into a high-dimensional space, but not to show a pronounced cluster structure. It comes from a system that generates artificial writings (images) of the digit “1” ([Hein and Audibert, 2005](#)). The numbers are generated from a function with five degrees of freedom and a sequence of transformations is later applied to introduce noise and different biases

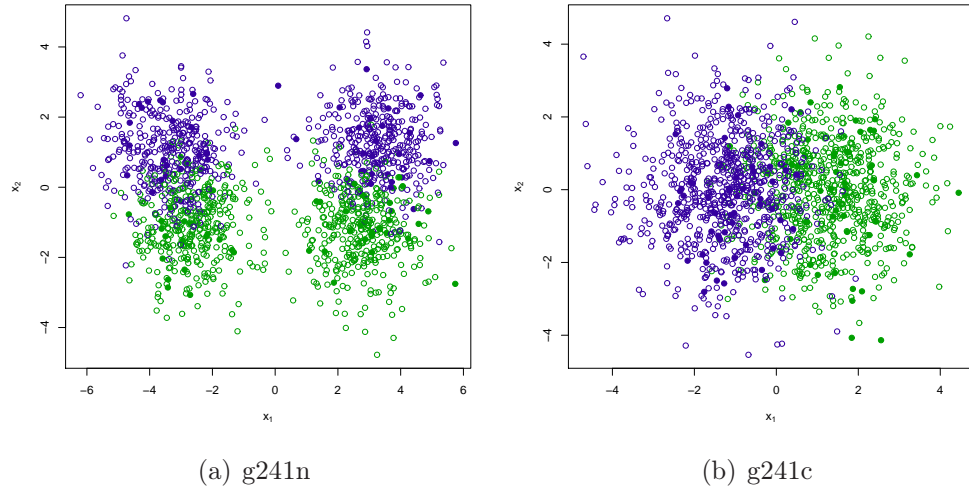


Figure 3.1: Two-dimensional projections of the benchmark datasets. Filled circles represent labeled data while empty circles represent unlabeled data.

to the images. The data will therefore lie close to a five-dimensional manifold. Figure 3.2(b), taken from [Chapelle and Schölkopf \(2006\)](#), shows one of the original images in the dataset (Figure 3.2(a)) and the transformed one (Figure 3.2)

For the protein function prediction task, the database described in section 2.2.1 was used as the set of labeled instances. Then, all the available *Embryophyta* proteins at UniProtKB/Swiss-Prot database that has no entries in the GOA project were added as the core set of unlabeled instances. As discussed in the introduction to this chapter, the proteins associated to the nodes in the functional path of a GO term, were also left as unlabeled instances regarding that classifier. Finally, 30000 unlabeled instances were randomly chosen in order to accomplish an approximate relation of ten unlabeled instances per each labeled one (remember from section 2.2.1 that the set of labeled protein sequences comprises 3368 instances). All the unlabeled sequences were characterized according to the procedure described in section 2.2.3 and the same feature selection strategy from section 2.2.5.

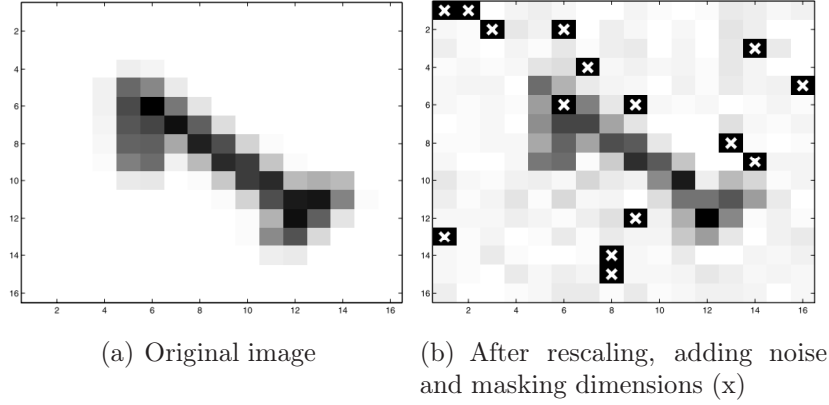


Figure 3.2: Example instance from the Digit1 dataset (taken from [Chapelle and Schölkopf \(2006\)](#))

3.2.3 Decision making

The S^3VM and Lap-SVM were used as base classifiers, both of them with the Gaussian kernel. For the Lap-SVM, the K-NN graph was selected for implementing the manifold regularization term, since there is some empirical evidence that suggests that fully connect graphs performs worse than sparse graphs ([Zhu and Goldberg, 2009](#), chapter 5).

All the parameters of the algorithms, including the dispersion of the kernels, the trade-off parameters of the SVMs, the regularization constants of both methods and the number of neighbors for constructing the graph, were tuned with a particle swarm optimization meta-heuristic. Again, the decision making was implemented following the one-against-all strategy with SMOTE oversampling for avoiding class-imbalance. Also, the 5-fold cross-validation strategy was implemented for assessing the performance of the predictors.

3.3 Results and discussion

3.3.1 Analysis of benchmark datasets

Table 4.1 shows the results obtained with each algorithm over the three benchmark datasets. Again, the geometric mean between sensitivity and specificity is used as global performance measure. The first line on each value of the table shows

Table 3.1: Performance over the three benchmark sets. Each position shows “mean \pm standard deviation” and the corresponding p-value. Highlighted values are significantly better than the supervised SVM.

Dataset	SVM	S ³ VM	Lap-SVM
g241n	0.76 \pm 0.11 –	0.80 \pm 0.12 0.43	0.80 \pm 0.07 0.34
g241c	0.61 \pm 0.07 –	0.76 \pm 0.13 0.03	0.58 \pm 0.16 0.70
Digit1	0.85 \pm 0.10 –	0.87 \pm 0.13 0.69	0.92 \pm 0.06 0.09

“mean \pm standard deviation” across the five repetitions of the cross-validation procedure. The second line shows the corresponding p-value for a paired t-test between the semi-supervised methods and the supervised SVM. Bold face values indicate which values are significantly better than the supervised SVM for a paired t-test at 90% significance level.

As expected, S³VM shows a superior performance for the dataset implementing the cluster assumption, while Lap-SVM outperforms the supervised SVM in the dataset that implements the manifold assumption. None of the semi-supervised methods is able to outperform the supervised algorithm when none of the semi-supervised assumptions holds. This is perhaps the most important issue when dealing with semi-supervised learning: previous knowledge of the nature of the data is required to select the appropriate learning model.

3.3.2 Analysis of GO prediction in *Embryophyta* plants

Figure 3.3 shows a comparison between the results with the S³VM (orange line) and the SVM method presented in chapter 2 (green line). The results with the supervised SVM are the same as in Figure 2.3, but the classes are ordered this time according to the performance of the S³VM method. Again, left plots show sensitivity, specificity and geometric mean achieved with the five-fold cross-validation procedure, while right plots depicts the corresponding p-values obtained from a paired t-test at a 95% significance level. Orange bars show the cases when the

S³VM significantly outperforms the supervised SVM and green bars show the opposite case.

The main purpose of this comparison is to verify whether or not the inclusion of the additional cluster-based semi-supervised term in the training of the SVM improves the performance of the system, thus providing information about the accomplishment of the cluster assumption when the unlabeled data is incorporated to the training process. Figure 3.3(a) shows that six out of the fourteen molecular functions considered in this ontology were significantly improved. In particular, *Receptor binding*, *Transcription factor activity* and *Enzyme regulator activity* have a special importance, considering that the SVM method was outperformed by BLASTp in those three GO terms when using the supervised model (see figure 2.3(a)). The inclusion of the cluster assumption also improved the performance on *Hydrolase activity**, *Binding** and *Protein binding**. Regarding the Cellular Component ontology (Figure 3.3(b)), eight cellular components were significantly improved, while another two (*Mitochondria* and *Cytoplasm**) also reached high p-values over 0.9. Finally, sixteen biological processes presented statistically significant improvements when including the unlabeled data with the semi-supervised cluster assumption. Only one biological process, *Lipid metabolic process*, suffered a statistically significant deterioration, which indicates that the unlabeled data is presenting a misleading cluster structure regarding this GO term.

In order to analyze how this improvements affect the system when compared to conventionally used prediction tools, Figure 3.4 shows a comparison between the results with the S³VM (orange line) and the traditional BLASTp method (blue line). It can be seen from figure 3.4(a) that the S³VM significantly outperforms BLASTp in five molecular functions, while BLASTp remains better than the S³VM only for *Transcription factor activity*. In contrast to Figure 2.3(a), BLASTp reduced the number of GO terms where it showed superiority from four GO terms to just one, while the machine learning method went from being superior in two molecular function to five. Regarding the cellular component ontology, there are only two cellular components for which there is no statistically significant difference between BLASTp and the S³VM: *Perixosome* and *Endosome*. For

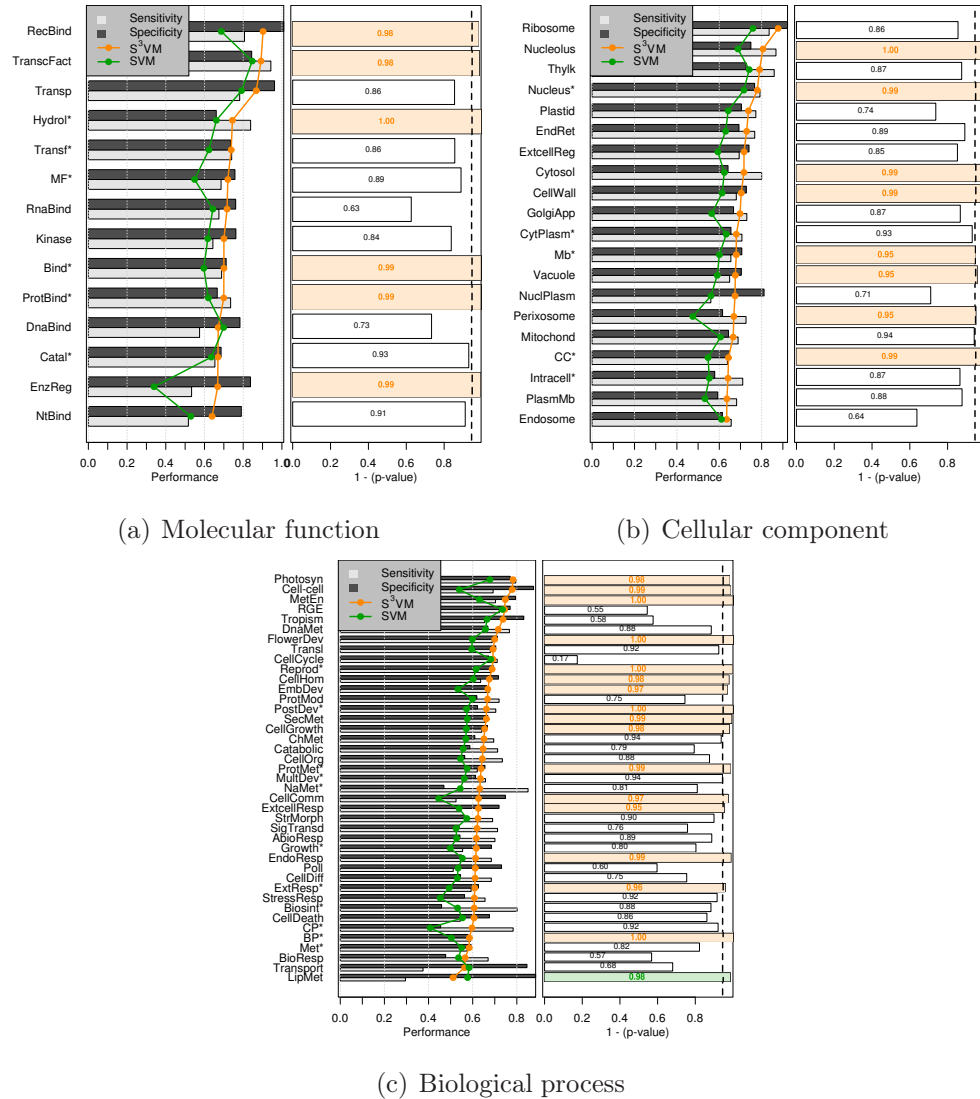


Figure 3.3: Comparison between the S³VM method and the supervised SVM. Bars in the left plots show sensitivity and specificity of the S³VM and lines depict geometric mean for S³VM (orange) and the classical supervised SVM (green). Right plots depict the p-values obtained by paired t-tests at a 95% significance level. For each ontology, the best predicted categories are ordered from top to bottom.

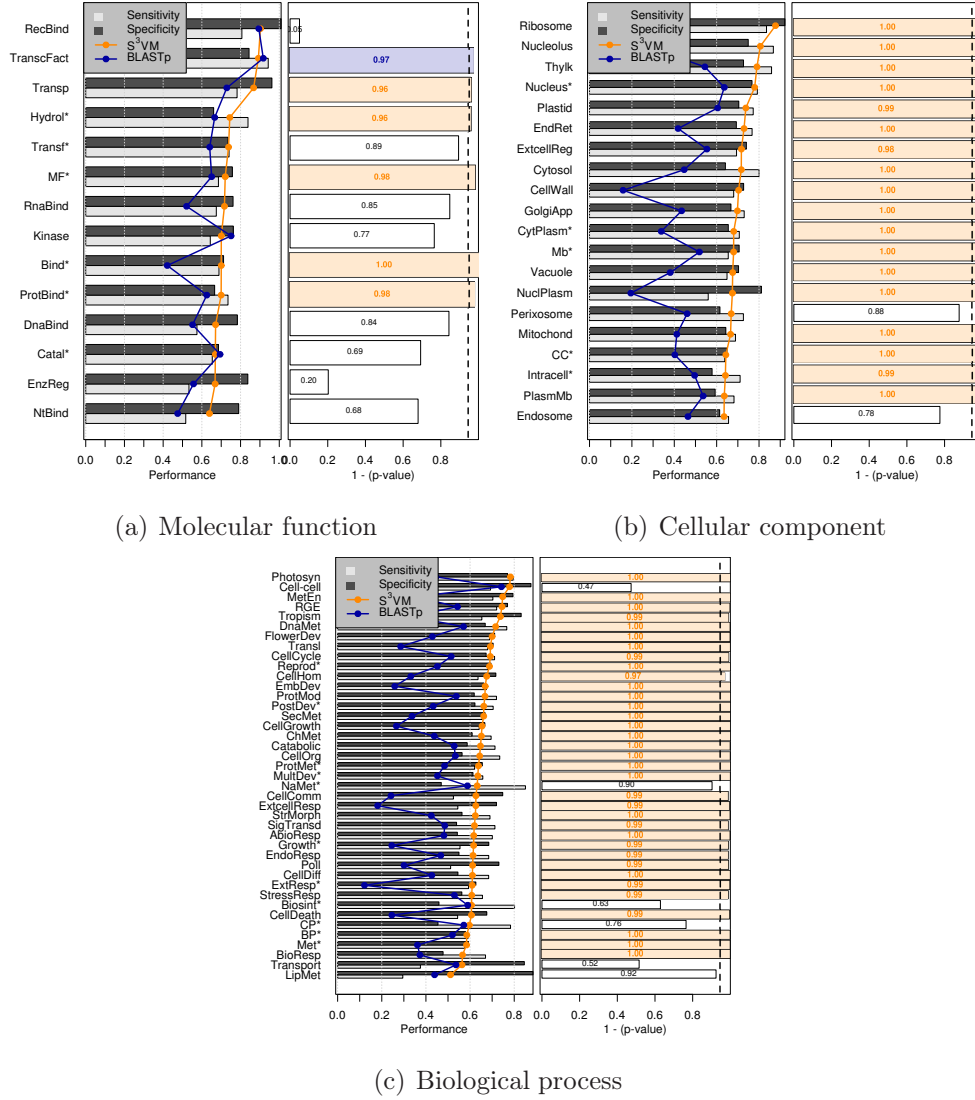


Figure 3.4: Comparison between BLASTp and the S³VM method. Bars in the left plots show sensitivity and specificity of the S³VM and lines depict geometric mean for S³VM (orange) and BLASTp (blue). Right plots depict the p-values obtained by paired t-tests at a 95% significance level. For each ontology, the best predicted categories are ordered from top to bottom.

all the remaining eighteen cellular components, the semi-supervised method obtained superior performance. A similar behavior is shown at figure 3.4(c), where the S³VM significantly outperforms BLASTp in 35 out of the 41 biological processes, while the remaining six process showed no statistical difference between the methods.

On the other hand, Figure 3.5 shows the comparison between the supervised SVM and the Laplacian-SVM. This analysis provides information about the impact of incorporating unlabeled data on the training set but, this time, by implementing the semi-supervised manifold assumption. This time, it is possible to see that there are less GO terms that have been improved by the inclusion of the unlabeled data. For the molecular function ontology (Figure 3.5(a)), only the *Nucleotide binding* and *Enzyme regulator activity* GO terms were significantly improved respecting the supervised SVM; in turn the implementation of the manifold assumption significantly degraded the performance for the GO term *Transcription factor activity*. Regarding the cellular component ontology (Figure 3.5(b)), improvements are present for *Perixosome*, *Vacuole* and the root node of the ontology, while a decrease is evinced for the *Nucleus** GO term. As for the biological process ontology, seven GO terms enhanced their prediction performance (*Embryonic development*, *Response to extracellular stimulus*, *Response to external stimulus**, *Metabolic process**, *Response to biotic stimulus*, *Cell communication* and the root node of the ontology), while another two were worsened (*Cell cycle* and *DNA metabolic process*).

Figure 3.6 depicts a comparison between the results obtained with BLASTp and the Lap-SVM method. The first important result that can be inferred from the present analyses is that, in general terms, for the problem of protein function prediction, the semi-supervised cluster assumption holds for many more cases than the semi-supervised manifold assumption. However, the most important aspect to be analyzed here, is how the results in Figure 3.6 complement the results from Figure 3.4. Only two molecular functions presented an statistically significant superior performance with the Lap-SVM over BLASTp. One of them, *RNA binding*, did not show statistical significance when comparing BLASTp and S³VM. The same behavior is present for the *Perixosome* cellular component and for the biological processes *Transport* and *Lipid metabolic process*. This results

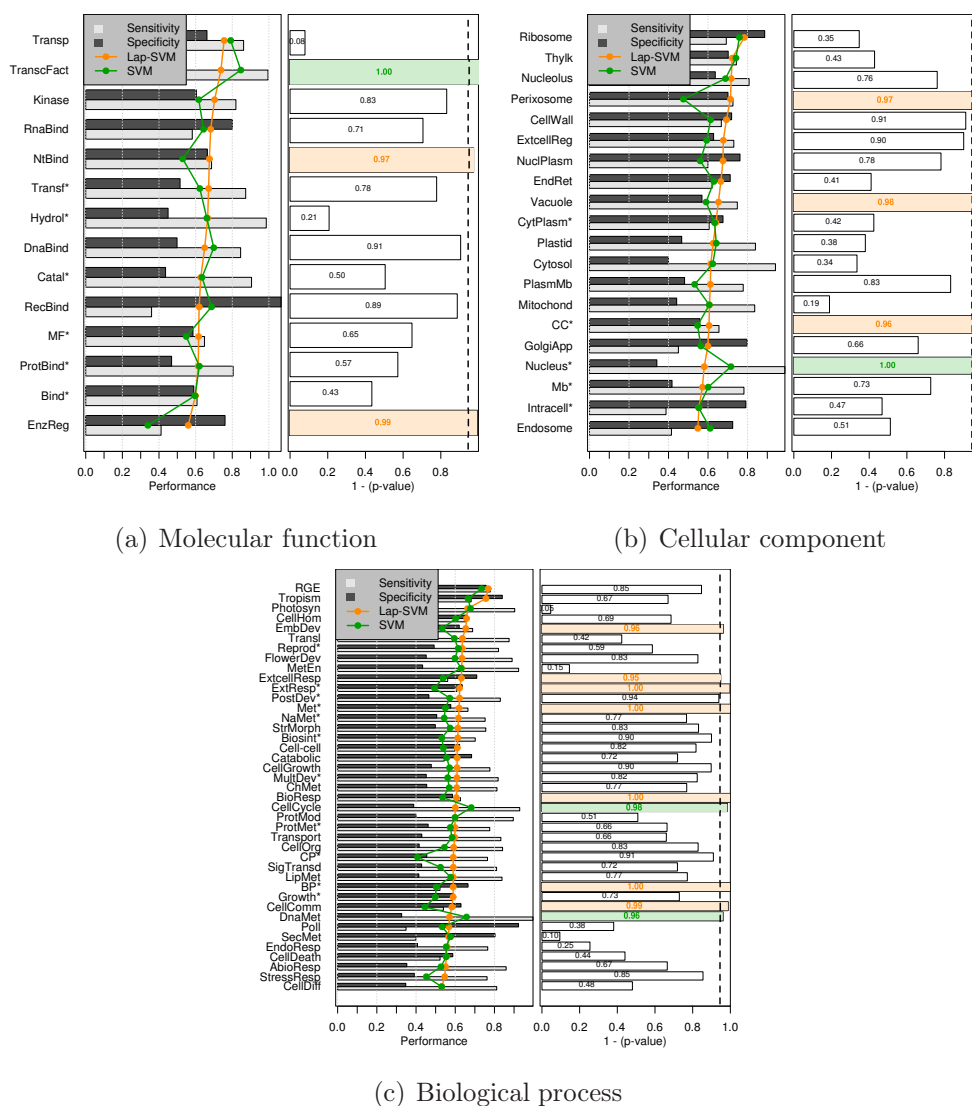


Figure 3.5: Comparison between the Lap-SVM method and the supervised SVM. Bars in the left plots show sensitivity and specificity of the Lap-SVM and lines depict geometric mean for Lap-SVM (orange) and the classical supervised SVM (green). Right plots depict the p-values obtained by paired t-tests at a 95% significance level. For each ontology, the best predicted categories are ordered from top to bottom.

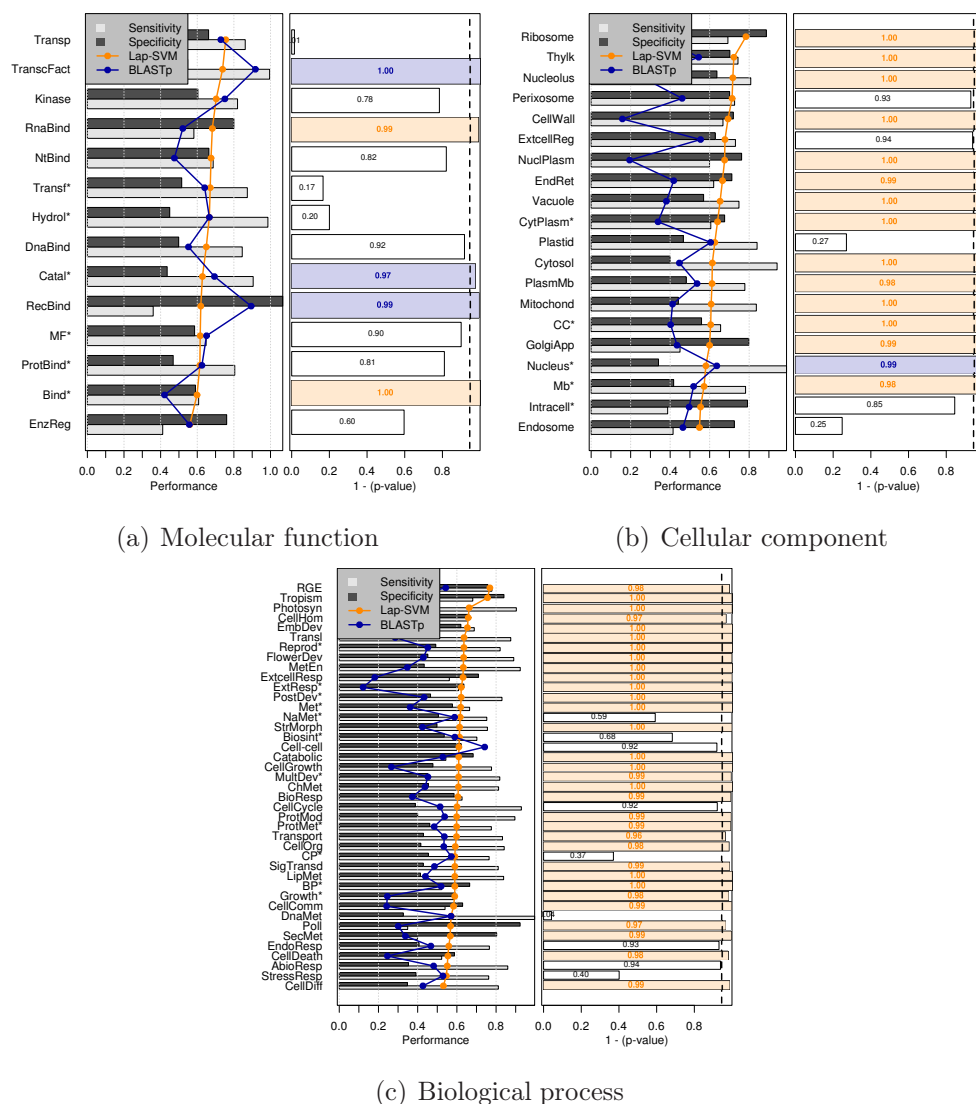


Figure 3.6: Comparison between BLASTp and the Lap-SVM method. Bars in the left plots show sensitivity and specificity of the Lap-SVM and lines depict geometric mean for Lap-SVM (orange) and BLASTp (blue). Right plots depict the p-values obtained by paired t-tests at a 95% significance level. For each ontology, the best predicted categories are ordered from top to bottom.

indicate that the manifold assumption is best suited than the cluster assumption for this particular GO terms. A few GO terms were not improved by any of the assumptions.

3.4 Concluding remarks

In this chapter, an analysis of the suitability of semi-supervised methods for the prediction of protein functions in *Embryophyta* plants was performed. A review of the state of the art of semi-supervised classifiers was presented, highlighting the different assumptions that each method does about the underlying distribution of the data. Two semi-supervised methods were chosen to perform the tests, each representing one of the main semi-supervised assumptions: cluster assumption and manifold assumption. The results show that semi-supervised learning applied to the prediction of GO terms in *Embryophyta* organisms, significantly outperforms the supervised learning approach, at the same time outperforming the commonly used sequence alignment strategy in most cases. In general terms, the highest performance were reached when applying the cluster assumption. However, several GO terms that were not significantly improved with the cluster assumption, achieved higher performance with the manifold based semi-supervised method, demonstrating that a single assumption is not enough for improving the learning process by the exploitation of the additional unlabeled data.

Chapter 4

Semi-supervised learning with multi-objective optimization

Semi-supervised learning uses unlabeled data to improve the estimation of the predictor function f_{θ} . However, there are two main concerns in order to correctly explore the information contained in the unlabeled data. First, as mentioned in the preceding chapter, each model has to make some specific assumption about the underlying structure of the data. Then, it turns evident that blindly selecting a semi-supervised learning method for a specific task will not necessarily improve performance over supervised learning. In fact, unlabeled data can lead to worse performance with the wrong assumption ([Zhu and Goldberg, 2009](#), chapter 2). Second, when integrating unsupervised and supervised information by means of an objective function, it is not usually clear what the best weighting between these components will be, and it is possible that the weighting chosen may have a significant effect on the final outcome ([Handl and Knowles, 2006](#)).

In this scenario, managing semi-supervised learning within the framework of multi-objective optimization, may provide a more flexible tool for the integration of both unsupervised and supervised components. Specifically, the use of Pareto optimization provides the means to avoid the need for hard constraints and for a fixed weighting between unsupervised and supervised objectives ([Handl, 2006](#)). Additionally, since multi-objective optimization allows the integration of multiple individual objectives, it would be possible to integrate several semi-supervised

assumptions within the same learning process, thus allowing for a wider applicability of the method.

In this chapter, a new method for semi-supervised learning within the framework of multi-objective optimization is proposed. This method combines supervised learning with the semi-supervised cluster and manifold assumptions, obtaining a flexible strategy that with low dependency on the data distribution. The efficiency of the method is tested over several toy problems and is finally employed on the prediction of GO terms for protein function prediction in *Embryophyta* plants.

4.1 Proposed method

Most density-based methods and graph-based methods reviewed in sections 3.1.2 and 3.1.3, respectively follow a common principle in order to incorporate the unlabeled samples into a supervised learner: they include an additional regularizer term into the optimization problem. As shown in equation (3.8), S³VM defines a regularization term which penalizes functions which vary in high-density regions:

$$J_{S^3VM}(\boldsymbol{\theta}) = \frac{1}{2} \|\boldsymbol{\theta}\|^2 + C \sum_{i=1}^L \ell(f_{\boldsymbol{\theta}}(\mathbf{x}_i)y_i) + \lambda \sum_{i=L+1}^{L+U} \ell(|f_{\boldsymbol{\theta}}(\mathbf{x}_i)|)$$

where λ is a regularization coefficient that must be set by the user. In the same way, the optimization criterion defined on equation (3.13) by the Lap-SVM method, uses a regularizer which prefers functions which vary smoothly along the manifold:

$$J_{LapSVM}(\boldsymbol{\theta}) = \frac{1}{L} \sum_{i=1}^L \ell(f_{\boldsymbol{\theta}}(\mathbf{x}_i)y_i) + \lambda_A \|\boldsymbol{\theta}\|^2 + \frac{\lambda_I}{(L+U)^2} \mathbf{f}_{\boldsymbol{\theta}}^T \Delta \mathbf{f}_{\boldsymbol{\theta}}$$

where λ_A and λ_I are again two regularization constants that must be fixed by the user. It is quite straightforward to notice that regularization can be viewed as a special case of a multi-objective optimization problem, where several objective functions are being linearly combined by the introduction of linear weights (regularization constants). The aforementioned equations can be viewed as a special

case of a multi-objective optimization problem where there are a number of possibly conflicting objective functions $\{J_i(\boldsymbol{\theta})\}_{i=1}^M$, and they are linearly combined in a unique objective function by the introduction of linear weights:

$$J(\boldsymbol{\theta}) = \lambda_1 J_1(\boldsymbol{\theta}) + \dots + \lambda_M J_M(\boldsymbol{\theta}) \quad (4.1)$$

Solving the regularized optimization problem for a unique combination of weights λ_i , yields to a solution that focuses on the objective functions with the highest weights. Many authors have demonstrated the inability of the method to capture Pareto optimal points that lie on non-convex portions of the Pareto optimal curve (Chen, 1998; Huang et al., 2007). Besides, it is well known that the method does not provide an even distribution of points in the Pareto optimal set, but only a linear approximation of the preference function (Marler and Arora, 2009).

A more flexible solution can be obtained by managing the optimization problem as a multi-objective optimization task, where the objective function produces a vectorial output of length M :

$$\mathbf{J}(\boldsymbol{\theta}) = \{J_1(\boldsymbol{\theta}), \dots, J_M(\boldsymbol{\theta})\} \quad (4.2)$$

In this setting, the optimization algorithm does not search for a unique solution, but for the set of all Pareto-optimal solutions with non-convex trade-off surfaces. Tackling the semi-supervised classification problem within the framework of multi-objective optimization provide a more flexible framework for the integration of both unsupervised and supervised components. Specifically, the use of Pareto optimization provides the means to avoid the need for hard constraints and for a fixed weighting between unsupervised and supervised objectives. Consequently, one would expect a multi-objective approach to semi-supervised classification to perform more consistently across different data sets, and to be less affected by model assumptions.

4.1.1 Objective functions

Once again, consider a set of labeled instances $\mathcal{X}_L = \{\mathbf{x}_i\}_{i=1}^L$ for which labels $\{y_i\}_{i=1}^L$ are provided, and a set of unlabeled instances $\mathcal{X}_U = \{\mathbf{x}_i\}_{i=L+1}^{L+U}$ the labels of which are not known. In principle, a linear decision function will be considered:

$$f_{\boldsymbol{\theta}}(\mathbf{x}) = \langle \boldsymbol{\theta}, \mathbf{x} \rangle \quad (4.3)$$

Three objective functions are proposed in a multi-objective minimization framework. One for reflecting the error achieved in the classification of the labeled training data, and the other two for exploiting the information contained on the unlabeled samples. A separate objective is defined for each semi-supervised assumption: low density assumption (and implicitly the cluster assumption) and the manifold assumption, covering the complete landscape of semi-supervised assumptions. The proposed objectives are:

$$J_{sup}(\boldsymbol{\theta}) = \frac{1}{2} \|\boldsymbol{\theta}\|^2 + C \sum_{i=1}^L \ell(f_{\boldsymbol{\theta}}(\mathbf{x}_i) y_i) \quad (4.4)$$

$$J_{clus}(\boldsymbol{\theta}) = \sum_{i=L+1}^{L+U} \ell(|f_{\boldsymbol{\theta}}(\mathbf{x}_i)|) \quad (4.5)$$

$$J_{man}(\boldsymbol{\theta}) = \sum_{ij} W_{ij} (f_{\boldsymbol{\theta}}(\mathbf{x}_i) - f_{\boldsymbol{\theta}}(\mathbf{x}_j))^2 \quad (4.6)$$

where $\ell(t) = \max(0, 1-t)$ is the hinge loss function, C is a trade-off parameter for regulating the complexity of the model and W_{ij} is the graph weight connecting the i -th and j -th instances. The first objective function, J_{sup} , is the standard SVM objective function that is minimized when the decision boundary maximizes the margin between the classes in the labeled instances. The second objective, J_{clus} , is minimized when the evaluations $f_{\boldsymbol{\theta}}(\mathbf{x}_i)$ of the unlabeled points \mathbf{x}_i are away from 0, that is, when the decision boundary is away from the unlabeled instances. This ensures the accomplishment of the low density assumption and, consequently, the cluster assumption. Finally, the third objective function, J_{man} , is an energy function that is minimized when the labels change slowly through

the graph, which accomplishes the manifold assumption.

Since these three objectives are not necessarily minimized for the same vector of parameters $\boldsymbol{\theta}$ simultaneously, it is necessary to provide a Pareto optimal set of solutions $\{\boldsymbol{\theta}_{*i}\}_{i=1}^P$ reflecting the trade-offs among the objectives. Then, a single solution can be selected according to its coherence with the application at hand.

4.1.2 Non-linear mapping

Traditional supervised SVMs can be extended to the non-linear case by using the kernel trick described in section . This procedure implies mapping the data into a high dimensional Hilbert space \mathcal{H} through a mapping $\Phi : \mathcal{X} \mapsto \mathcal{H}$, where a linear decision boundary is able to correctly assign the class labels. The mapping Φ can be explicitly computed or only implicitly through the use of a kernel function K such that $K(\mathbf{x}_1, \mathbf{x}_2) = \langle \Phi(\mathbf{x}_1), \Phi(\mathbf{x}_2) \rangle$. A square kernel matrix \mathbf{K} can thus be computed, which elements are given by $K_{ij} = K(\mathbf{x}_i, \mathbf{x}_j)$.

However, since the inclusion of the additional objectives does not allow to express the whole optimization problem in terms of dot products, the optimization must be performed directly on the feature space. For this purpose is necessary to consider that, even in the case of a high dimensional feature space \mathcal{H} , a finite training set of size N (for the semi-supervised setting $N = L + U$), when mapped to this feature space, spans a vectorial subspace $\mathcal{E} \subset \mathcal{H}$ whose dimension is at most N (Belkin et al., 2006).

Let $\{\mathbf{v}_i\}_{i=1}^N$ be an orthonormal basis of \mathcal{E} , where the p -th vector \mathbf{v}_p can be expressed as:

$$\mathbf{v}_p = \sum_{i=1}^N A_{ip} \Phi(\mathbf{x}_i) \quad (4.7)$$

where \mathbf{A} is a matrix that necessarily has to satisfy $\mathbf{A}^T \mathbf{K} \mathbf{A} = \mathbf{I}$. A possible solution (among several choices) can be obtained by the eigendecomposition of \mathbf{K} as $\mathbf{K} = \mathbf{U}^T \boldsymbol{\Lambda} \mathbf{U}$, which provides $\mathbf{A} = \mathbf{U} \boldsymbol{\Lambda}^{-1/2}$. Then, a mapping $\Psi : \mathcal{X} \mapsto \mathbb{R}^N$

can be computed as:

$$\Psi_p(\mathbf{x}) = \sum_{i=1}^L A_{ip} K(\mathbf{x}, \mathbf{x}_i), \quad p = 1, 2, \dots, N \quad (4.8)$$

This mapping satisfies the relation $\langle \Psi(\mathbf{x}_i), \Psi(\mathbf{x}_j) \rangle = K(\mathbf{x}_i, \mathbf{x}_j)$ and can be used for obtaining a feature space where the linear function f_{θ} provides a non-linear decision boundary in the original space \mathcal{X} .

4.2 Proposed method: multi-objective semi-supervised learning for predicting GO terms in *Embryophyta* plants

4.2.1 Selected multi-objective strategy: cuckoo search

Due to their population-based nature, meta-heuristic algorithms are able to approximate the whole Pareto front of a multi-objective optimization problem in a single run. Nowadays, real world applications of large scale multi-objective optimization are feasible thanks to the developments of this kind of algorithms, and several review papers and books have been recently published in this matter (Talbi, 2009; Yang, 2010; Zhou et al., 2011).

Recently, a new meta-heuristic search algorithm called Cuckoo Search (CS) has been developed by Yang and Deb (2009), and its multi-objective extension, Multi-Objective Cuckoo Search (MOCS) was lately proposed (Yang and Deb, 2011). MOCS has been tested against a subset of well-chosen test functions, and have been applied to solve design optimization benchmarks in structural engineering showing a clear superiority in comparison with other algorithms (Yang and Deb, 2013). This superiority can be attributed to the fact that cuckoo search uses a combination of vectorized mutation, crossover by permutation and Lévy flights and selective elitism among the best solutions. In addition, the not-so-good solutions can be replaced systematically by new solutions, and new solutions are often generated by preferring quality solutions in the solution sets. Thus, the mechanism of the overall search moves is more subtle and balanced, compared

with the simple mechanism used by other meta-heuristics such as particle swarm optimization or genetic algorithms.

Cuckoo Search is based on the parasitic behavior exposed by some species of Cuckoo birds. Its natural strategy consists on leaving eggs in host nests created by other birds. This eggs presents the particularity to have a big similitude with host eggs, the more similar they are, the greater is its chance of survival. Based on this statement, Cuckoo Search uses three idealized rules (Yang and Deb, 2009):

1. Each cuckoo lays one egg at a time, and dumps it in a randomly chosen nest.
2. The best nests with high quality of eggs (solutions) will carry over to the next generations.
3. The number of available host nests is fixed, and a host can discover an alien egg with a probability $p_a \in [0, 1]$. In this case, the host bird can either throw the egg away or abandon the nest so as to build a completely new nest in a new location.

The first assumption can be achieved by the generation of new random solutions from the existing ones. Such procedure is driven by Lévy flights of the form:

$$\theta_i^{(t+1)} = \theta_i^{(t)} + \alpha \oplus \text{Lévy}(\gamma) \quad (4.9)$$

where α is the step size and the random step is drawn from a Lévy distribution:

$$\text{Lévy} \sim u = t^{-\gamma}, (1 < \gamma \leq 3) \quad (4.10)$$

which has an infinite variance with an infinite mean. The steps form a random walk with a power-law step-length distribution with a heavy tail. Figure 4.1 shows an example of three Lévy flights generated from the same starting point at $(0, 0)$.

On the other hand, the last assumption can be approximated by replacing a fraction p_a of the Q nests by new nests (with new random solutions at new locations). Parameters p_a and α are directly concerned with the efficiency of

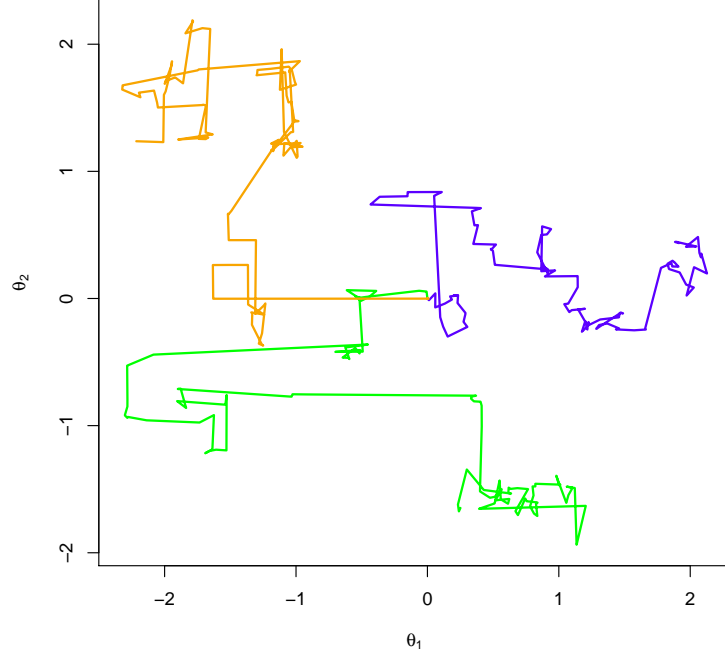


Figure 4.1: Example of three Lévy flights of length 100 starting from the origin

the search, allowing to find globally and locally improved solutions, respectively. Additionally, these parameters directly influence the convergence rate of the optimization algorithm. For instance, if the value of p_a tends to be small and α value is large, the algorithm will increment the number of iterations to converge to an optimal value. On the other hand, if the value of p_a is large and α is small, the convergence speed of the algorithm tends to be very high but it is more likely to converge to a local optimum. [Valian et al. \(2011\)](#) proposed an improvement which consists on using a range of p_a and α to change dynamically in each iteration, following the equations:

$$p_a(t) = p_{max} - \frac{t}{T}(p_{max} - p_{min}) \quad (4.11)$$

$$\alpha(t) = \alpha_{max} e^{(cT)} \quad (4.12)$$

Algorithm 2 Multi-objective Cuckoo Search

Require: Objective function $\mathbf{J}(\boldsymbol{\theta}) = [J_1(\boldsymbol{\theta}), J_2(\boldsymbol{\theta}), \dots, J_M(\boldsymbol{\theta})]$

Require: Initial population of P host nests $\{\boldsymbol{\theta}_i\}_{i=1}^P$, each one with M eggs

Require: Maximal number of iterations T

Initialize the iterations counter $t = 0$

Initialize an empty set of Pareto-optimal solutions $\mathcal{P}^{(0)} = \emptyset$

while $(|\mathcal{P}| < P) \vee (t < T)$ **do**

 Get a cuckoo randomly (say i) by Lévy flights

 Evaluate and check if the solution is Pareto optimal

 Choose a nest among P (say j) randomly

 Evaluate M solutions for nest j

if New solutions of nest j dominate those of nest i **then**

 Replace nest i by the new solution of nest j

end if

 Abandon a fraction p_a of worst nests

 Sort and find the current Pareto optimal solutions $\{\boldsymbol{\theta}^*\}^{(t)}$ and update the set of global Pareto optimal solutions $\mathcal{P}^{(t+1)} = \mathcal{P}^{(t)} \cup \{\boldsymbol{\theta}^*\}^{(t)}$

end while

return \mathcal{P}

where t is current iteration, T is the maximum number of iterations, and c is a constant given by:

$$c = \frac{1}{T} \log \left(\frac{\alpha_{min}}{\alpha_{max}} \right) \quad (4.13)$$

The MOCS algorithm is summarized in Algorithm 2. Notice that the only free parameters that must be directly chosen by the user are the size of the population and the maximal number of iterations. While the latter is not of significant importance as it only provides an “emergency stop” in the cases when the algorithm does not converge, the former is directly linked to the number of points in the Pareto front that will be obtained. In order to obtain a good resolution and maximizing the precision on the estimation of the Pareto front, this number must be as high as possible. However, it will increase the computational time for the training. This parameter was empirically fixed to $P = 1000$ for all the analysis in this chapter.

4.2.2 Decision making

As in the previous analyses, the Gaussian kernel and the K-NN graph were selected as base tools for implementing the objective functions. There are only three free parameters to be tuned this time: the dispersion of the kernel, the trade-off parameter of the SVM in the supervised objective and the number of nearest neighbors for constructing the graph in the manifold-based objective. Such parameters were, once again, tuned the particle swarm optimization strategy. It is important to remark that, unlike the analyses in the previous chapter, all the regularization parameters are unnecessary here.

Again, the decision making was implemented following the one-against-all strategy with SMOTE oversampling for avoiding class-imbalance and 5-fold cross-validation was implemented for assessing the performance of the predictors. However, there is one additional step that is necessary for the multi-objective strategy: selecting the most appropriate solution among the set of Pareto-optimal trade-offs. This task is usually left to be carried out by a human expert, depending on his knowledge of the application at hand. In the case of protein functional prediction, the chosen solution must reflect the biological knowledge about the problem, which is contained in the set of GO terms associated to each sequence.

To this end, an additional cross-validation step is performed, in order to select the solution with the highest prediction accuracy, which in turn constitutes the most biologically feasible solution. Figure 4.2 depicts the complete methodology for estimating the performance of the proposed method. Every 80/20 split represents a cross-validation procedure (80/20 is the partitioning for a single fold in a 5-fold cross-validation schema).

4.3 Results and discussion

In this section, the same datasets described in the preceding chapter (section 3.2.2) are used for obtaining an estimation of the performance of the proposed method. In order to provide a more detailed analysis on the interpretation of the Pareto-optimal sets that are obtained from the multi-objective strategy, several solution points from the same Pareto front are analyzed in the case of the

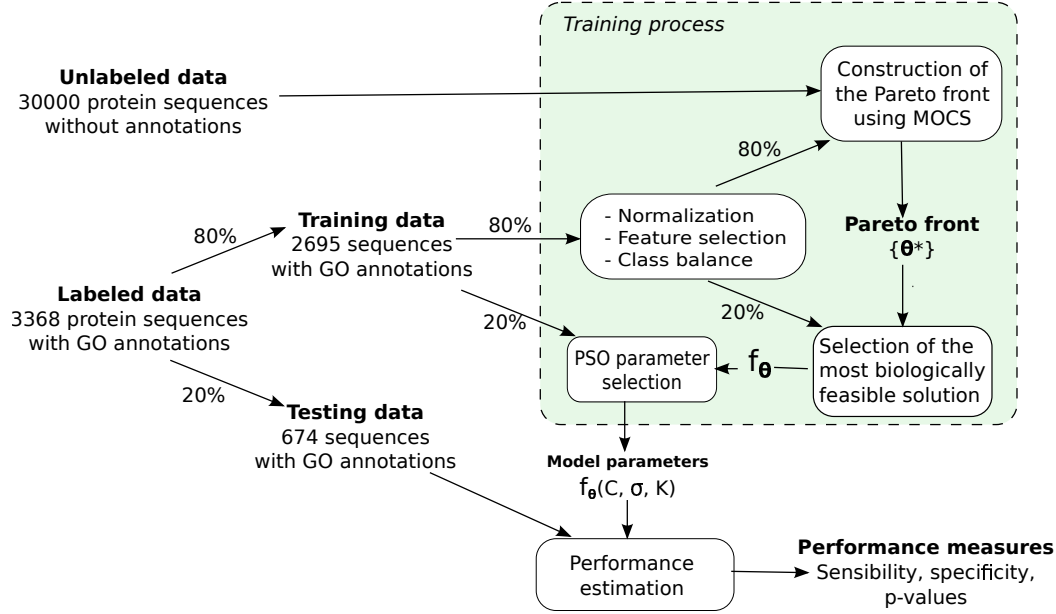


Figure 4.2: Flow diagram of the proposed methodology. The green area highlights the training process

benchmark datasets. Later, the method is tested over the *Embryophyta* proteins database and the results are compared with those of the supervised approach.

4.3.1 Analysis with the benchmark datasets

Figures 4.3, 4.4 and 4.5 show the Pareto fronts obtained for each one of the benchmark datasets g241n, g241c, and Digit1, respectively. For each figure, the left plot depicts a three-dimensional reconstruction of the Pareto front, while the right plots show the two-dimensional projections for a better comprehension. Each Pareto front has four highlighted solutions: the three minima for each objective function (green for the supervised objective, yellow for the cluster-based objective and orange for the manifold-based objective) and the most feasible solution found by the proposed method (red).

It is important to note that, since the cross-validation procedure implies the training and testing of several classifiers (one per each fold), the depicted Pareto fronts correspond only to the first fold of each problem. Nevertheless, in general terms there is a low variability among the results. Table 4.1 shows the global

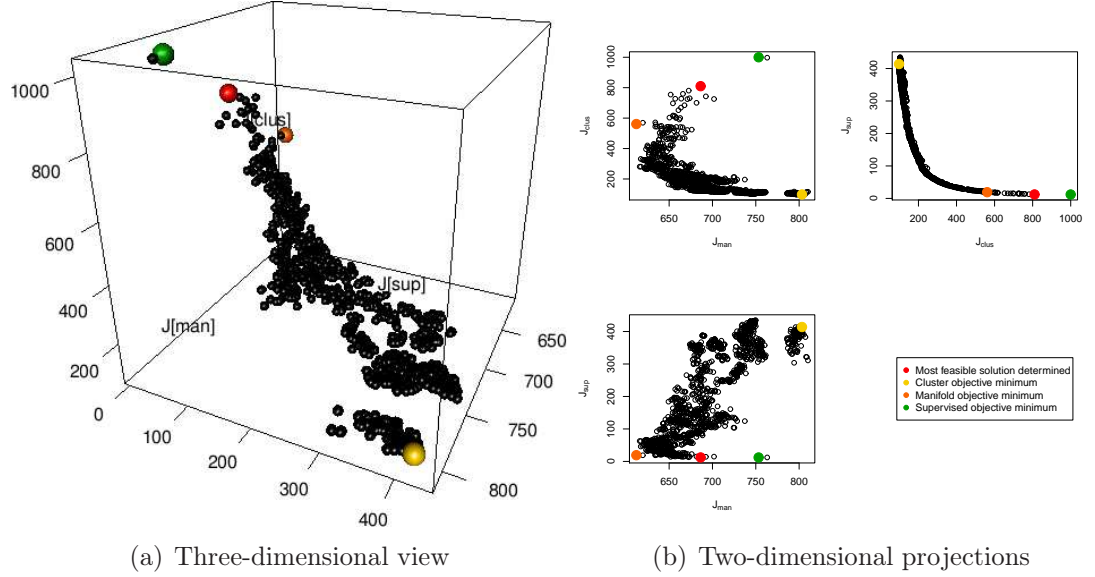


Figure 4.3: Pareto front for the g241n benchmark dataset. The yellow, orange and green dots depict the minima for the the manifold, cluster and supervised objectives, respectively. The red dot depicts the most feasible solution according to the class labels, found by the cross-validation procedure.

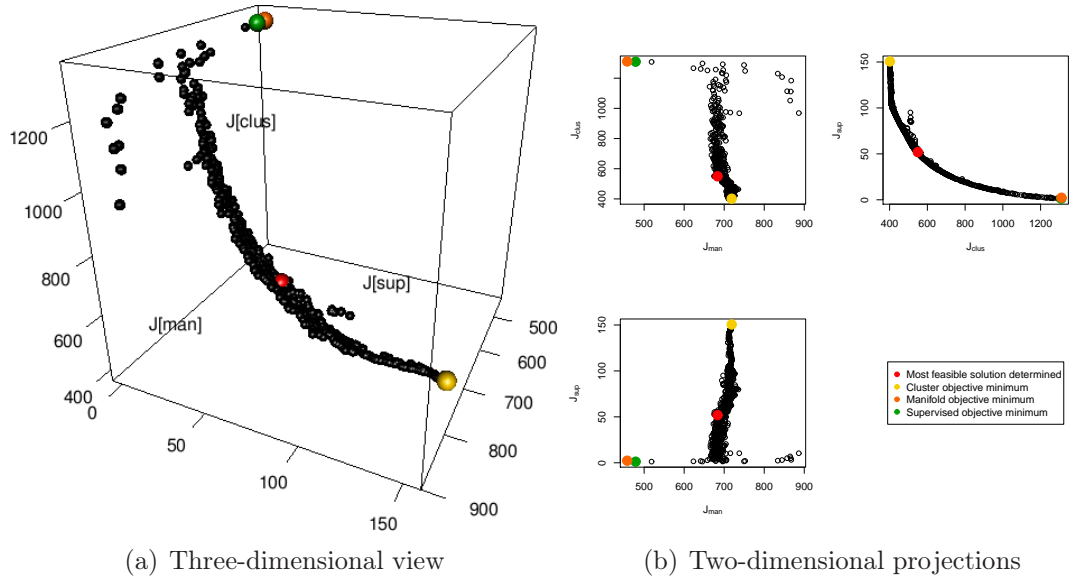


Figure 4.4: Pareto front for the g241c benchmark dataset. The yellow, orange and green dots depict the minima for the the manifold, cluster and supervised objectives, respectively. The red dot depicts the most feasible solution according to the class labels, found by the cross-validation procedure.

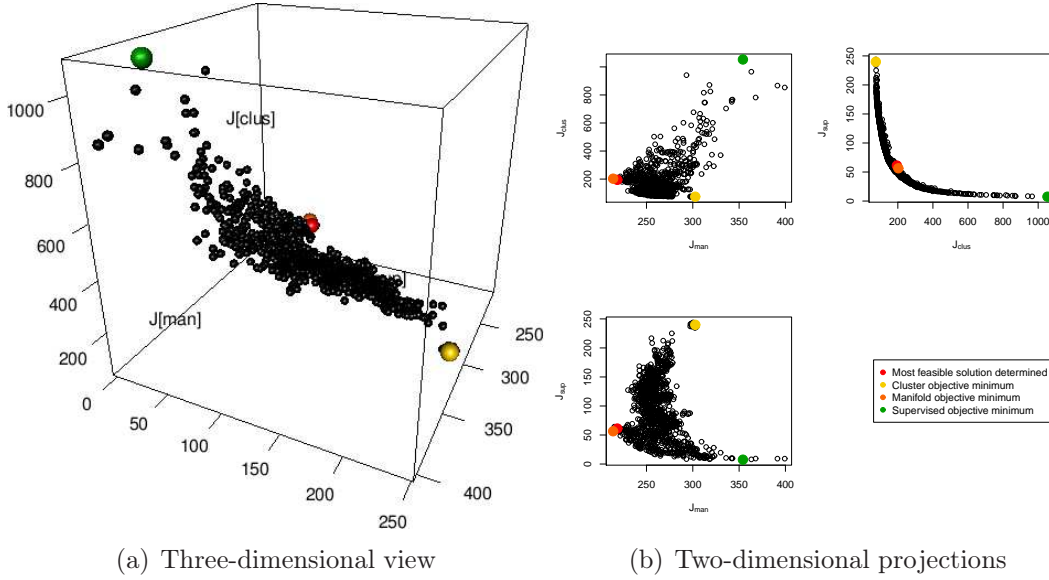


Figure 4.5: Pareto front for the *Digit1* benchmark dataset. The yellow, orange and green dots depict the minima for the the manifold, cluster and supervised objectives, respectively. The red dot depicts the most feasible solution according to the class labels, found by the cross-validation procedure.

performance results for the three benchmark problems, with each one of the highlighted solutions. Each position in the table shows the mean and standard deviation across the ten folds of the cross-validation procedure.

There are several interesting results to observe from Table 4.1. First of all, it must be noted that the proposed method is able to automatically select the best solution on the Pareto front, implicitly assigning a different weight to each objective, that is consistent with the nature of the problem. Remember from section 3.2.2 that the *g241n* dataset has a misleading structure where unlabeled data are neither organized on well defined clusters nor on an underlying manifold structure. In this case, it is possible to observe (see Figure 4.3) that the best solution found (the red one) is closer to the minimum of the supervised objective (green dot) than to any other minima. That means that the method assigned a higher importance to the supervised objective than to the other two. Regarding the *g241c* dataset, which has a well defined cluster structure, the best solution found is consistently closer to the yellow dot (see Figure 4.4) that represents the minimum of the cluster-based objective. The proper observed on Figure 4.5,

Table 4.1: Performance over the three benchmark sets for several solutions from the Pareto fronts. Each position shows “mean \pm standard deviation”. Highlighted values on the right are the highest among the three individual objectives.

Dataset	Proposed solution	Supervised	Cluster	Manifold
g241n	0.63 \pm 0.008	0.60 \pm 0.031	0.45 \pm 0.068	0.56 \pm 0.014
g241c	0.725 \pm 0.002	0.56 \pm 0.256	0.67 \pm 0.033	0.15 \pm 0.198
Digit1	0.93 \pm 0.005	0.71 \pm 0.264	0.70 \pm 0.159	0.92 \pm 0.011

where the best solution found is very close to the minimum of the manifold-based objective (orange dot). This is consistent with the nature of the `Digit1` dataset which was designed to have an underlying manifold structure.

The second aspect that must be noted here is that minimizing one single objective is not enough to achieve the best performance results on any problem. On all cases, the best results were achieved for a solution on the Pareto front, that does not exactly matches any of the single-objective minima. Even in the case of the `Digit1` dataset (Figure 4.5), where the best found solution (red) is very close to the minimum of the manifold-based objective (orange), there is a slight difference between those solutions that is reflected in the performance on Table 4.1.

Finally, it is worth to note the low standard deviation values achieved by the proposed method, which demonstrates the reliability of the method in spite of the random component of the underlying optimization strategy employed.

4.3.2 Analysis of GO prediction in *Embryophyta* plants

Figure 4.6 shows a comparison between the results with the proposed method (orange line) and the classical supervised SVM method presented in chapter 2 (green line). Again, left plots show sensitivity, specificity and geometric mean achieved with the five-fold cross-validation procedure, while right plots depicts the corresponding p-values obtained from a paired t-test at a 95% significance level. Orange bars show the cases when the proposed multi-objective method significantly outperforms the supervised SVM and green bars show the opposite case. This time, Figure 4.6(a) shows that nine molecular functions reached

significantly better prediction performance with the proposed method over the supervised SVM. Also, on the five remaining molecular functions, both methods achieved statistically similar results, presumably since the unlabeled data is not contributing to improve the predictions, but in no case the inclusion of this data degraded the performance of the predictor as it happened with the Lap-SVM in the preceding chapter (Figure 3.5). A similar behavior was also obtained for the Cellular Component ontology in Figure 4.6(b), where fifteen out of the 21 considered cellular components were statistically better predicted by the proposed method. The same happened also for the Biological Process ontology (Figure 4.6(c)) where only nine out of the whole set of 41 biological processes did not show an statistically significant improvement.

Finally, Figure 4.7 shows a comparison between the proposed method and BLASTp. From the whole set of 75 GO terms included in the database, only *Transcription Factor Activity* showed an statistically better prediction performance with BLASTp. This can be due to the characterization step and could be the subject of future studies.

4.4 Concluding remarks

In this chapter, a multi-objective optimization-based method for performing semi-supervised prediction of protein functions in *Embryophyta* plants was proposed. The method includes three independent objectives covering the main assumptions made by most semi-supervised methods in the literature, and combines them on a multi-objective optimization framework. This framework provides a set of trade-off solutions that can be analyzed to select the most feasible from the biological perspective, by applying a cross-validation strategy. The results show that this method is able to retrieve a prediction model that achieves equal or in most cases performance than the classical supervised SVM. Also, it outperforms the predictions of the commonly used BLASTp method for most of the GO terms analyzed in *Embryophyta* proteins.

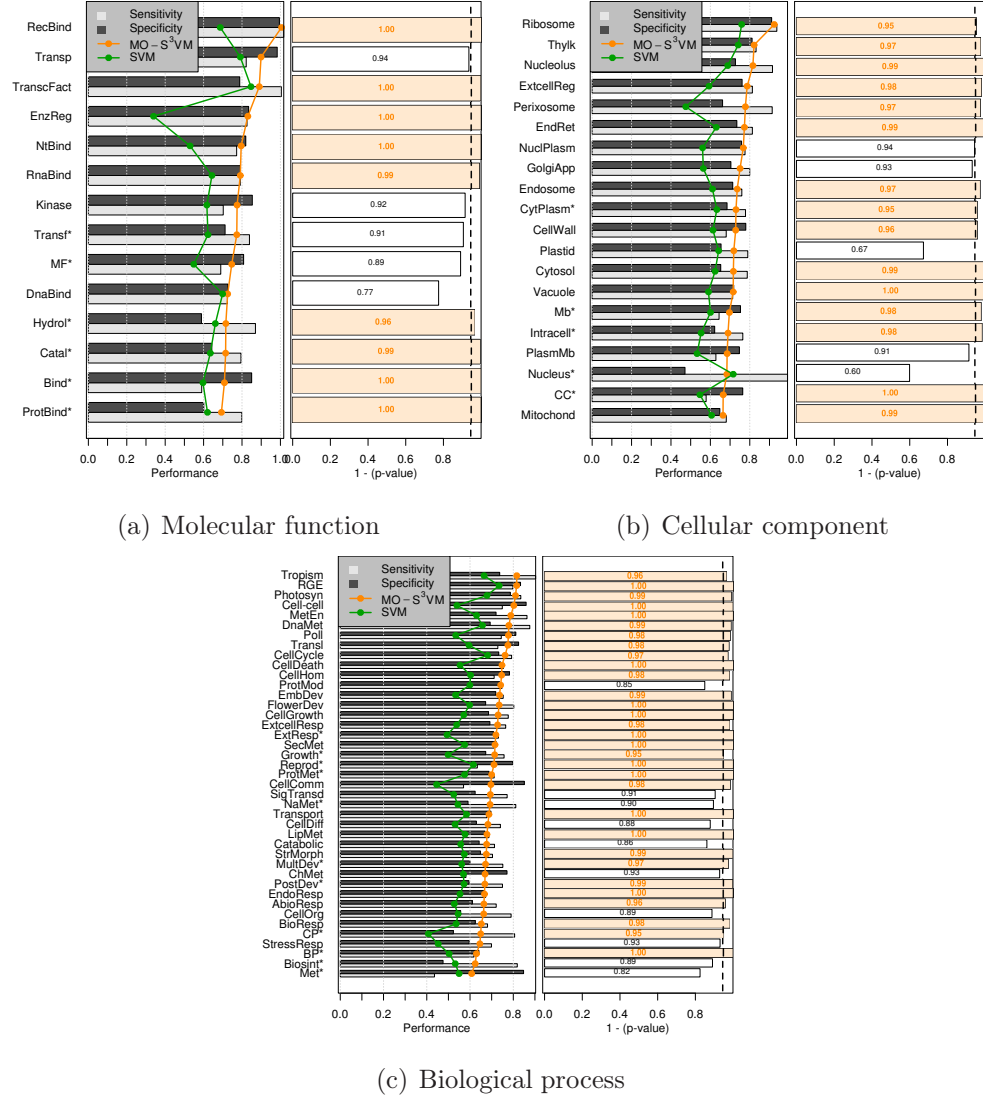


Figure 4.6: Comparison between the proposed multi-objective S^3VM method and the supervised SVM. Bars in the left plots show sensitivity and specificity of the proposed method and lines depict geometric mean for it (orange) and the classical supervised SVM (green). Right plots depict the p-values obtained by paired t-tests at a 95% significance level. For each ontology, the best predicted categories are ordered from top to bottom.

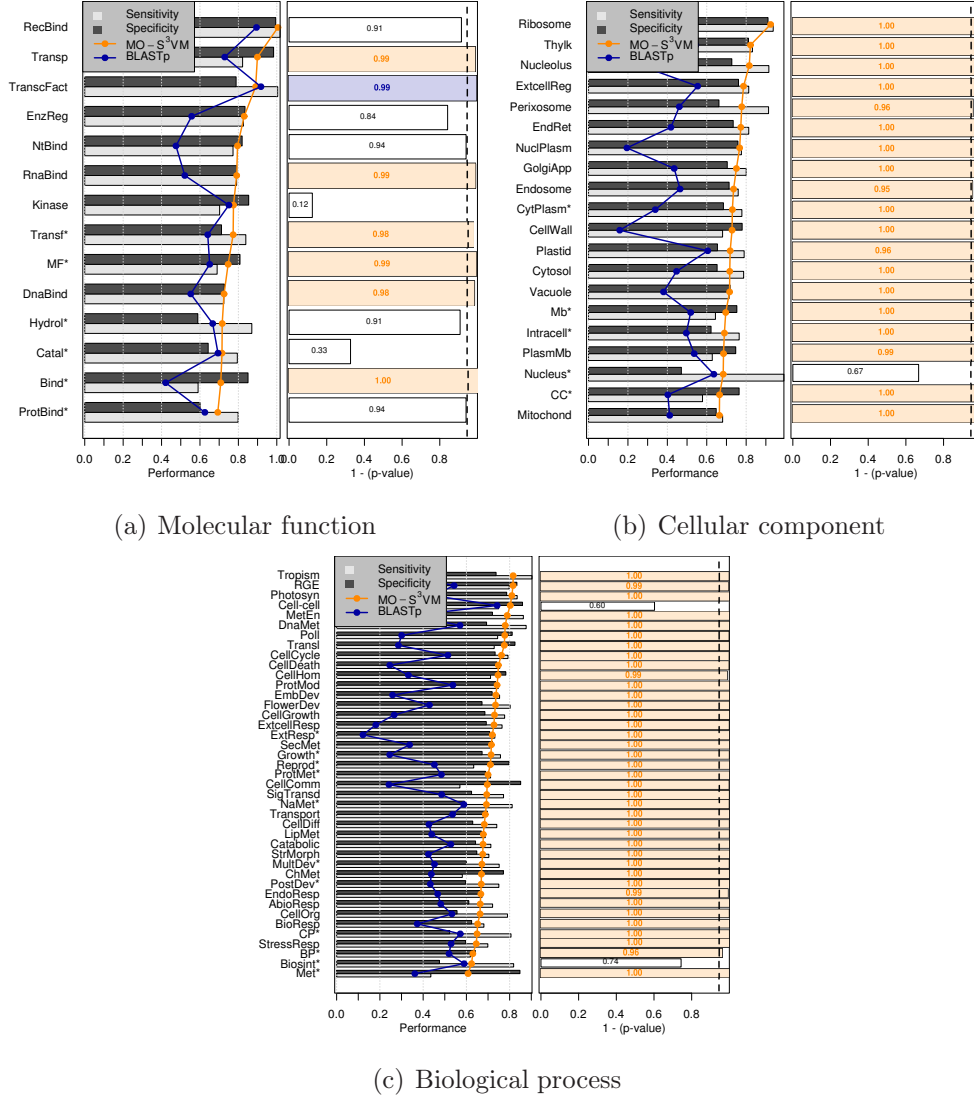


Figure 4.7: Comparison between the proposed multi-objective S^3VM method and BLASTp. Bars in the left plots show sensitivity and specificity of the proposed method and lines depict geometric mean for it (orange) and BLASTp (blue). Right plots depict the p-values obtained by paired t-tests at a 95% significance level. For each ontology, the best predicted categories are ordered from top to bottom.

Chapter 5

Conclusions

5.1 Main contributions

Protein function prediction is one of the most important challenges in bioinformatics. The most common approach to perform this task is by using strategies based on annotation transfer from homologues. The annotation process centers on the search for similar sequences in databases of previously annotated proteins, by using sequence alignment tools such as BLASTp. However, high similarity does not necessarily implies homology, and there could be homologues with very low similarity. As an alternative to alignment-based tools, more recent methods have used machine learning techniques trained over feature spaces of physical-chemical, statistical or locally-based attributes, in order to design tools that can be able of achieving high prediction performance when classical tools would certainly fail.

The present work lies on the framework of machine learning applied to protein function prediction, through the use of a modern paradigm called semi-supervised learning. This paradigm is motivated on the fact that in many real-world problems, acquiring a large amount of labeled training data is expensive and time-consuming. Because obtaining unlabeled data requires less human effort, it is of great interest to include it in the learning process both in theory and in practice. A high number of semi-supervised methods have been recently proposed and have demonstrated to improve the accuracy of classical supervised approaches in a vast number of real-world applications.

Nevertheless, the successfulness of semi-supervised approaches greatly depends on prior assumptions they have to make about the data. When such assumptions does not hold, the inclusion of unlabeled data can be harmful to the predictor. Here, the main approaches to perform semi-supervised learning were analyzed on the problem of protein function prediction, and their underlying assumptions were identified and combined in a multi-objective optimization framework, in order to obtain a novel learning model that is less dependent on the nature of the data.

All the experiments and analyses were focused on land plants (*Embryophyta*), which constitutes an important part of the national biodiversity of Colombia, including most agricultural products. Consequently, the Gene Ontology slim for plants was used to define the target functions to be predicted. However, it is very important to clarify that the methods and analyses performed in this thesis can be replicated without major concerns over other target groups of study as far as there is enough data to train the learning models.

Listed below are the main original contributions of this thesis:

- Construction of a comprehensive database comprising all the available *Embryophyta* protein sequences with at least one annotation in the Gene Ontology Annotation project (GOA). The dataset includes 3368 protein sequences from 189 different land plants and, for each sequence, a set of 438 physical-chemical and statistical attributes was computed. This database does not include annotations associated to evidence codes from automatically-assigned annotations and all the sequences belong to the reviewed part of Uniprot. The database is publicly available at:
<http://www.biomedcentral.com/1471-2105/14/68/>.
- Definition of a full methodology for training machine learning models in the prediction of Gene Ontology terms from the information contained in the primary structure of proteins. Unless most published studies, this methodology carefully describes of all the necessary steps from the conformation of the database, filtering, characterization, normalization, class balance, feature selection, and decision making, in order to obtain unbiased estimations of predictability.

- To the best of our knowledge, this thesis contains the most complete analysis on the predictability of GO terms from primary sequence information in land plants. While most analyses in the state of the art comprise only a few selected terms and single ontologies, the analysis presented in this thesis includes 75 GO terms covering molecular functions, cellular components and biological processes. This analysis provides a valuable guide for researchers interested on further advances in protein function prediction on *Embryophyta* plants.
- A complete analysis on the applicability of semi-supervised learning methods to the problem of protein function prediction over *Embryophyta* organisms. This analysis is focused on the successfulness achieved by the different assumptions made by the most common semi-supervised algorithms. It revealed that the cluster assumption is, in general terms, the most suitable for the problem at hand. However, it was also found that the manifold assumption is more successful on the prediction of some GO terms, showing that the assumptions can be regarded as complimentary to each other.
- Development of a novel method for semi-supervised learning that incorporates three independent objectives in a multi-objective optimization framework. This approach provides less dependence on the assumptions to be made about the data, which is the main drawback of the current semi-supervised methods. The multi-objective optimization strategy generates a set of trade-off solutions regarding the three proposed objectives and the most biologically feasible solution can be chosen through its coherence with the information contained on the GO labels. The results show that this method achieves comparable and in most cases superior prediction performance than both machine learning-based and alignment-based methods for protein function prediction.

Additionally, several works were derived from the ideas on this thesis, originating three master thesis that are being developed under the co-supervision of the author of this work. These three works are mainly focused on expanding machine learning approaches in several dimensions for dealing with GO term prediction from protein sequences:

- *Methodology for multi-class classification applied to bioinformatics in the prediction of protein functions in plants* by Andrés Felipe Giraldo Forero.
- *Methodology for the characterization and classification of proteins using wavelet-based approaches and dissimilarity learning* by Gustavo Alonso Arango Argoty.
- *Class-balance methodology for the prediction of protein functions oriented towards the automation of protein annotation process* by Sebastián García L’opez.

As a side result, this line of works have constituted the main core of the bioinformatics research at Universidad Nacional de Colombia, sede Manizales, leading to the development of the project “Predicción de términos de la Ontología Genética a partir de métodos de caracterización dinámica y clasificación semi-supervisada de secuencias de aminoácidos, aplicada a la clasificación funcional de proteínas de café (*Coffea arabica*)”, jointly with the Centro Nacional de Investigaciones en Café (CENICAFÉ), and currently founded by COLCIENCIAS. Besides, this line of work has also contributed to the creation of a masters program on bioinformatics at Universidad Nacional de Colombia, sede Manizales, and has generated a high number of papers and conference proceedings.

5.2 Future research directions

In spite of the good results achieved in this thesis, there are several subjects that can be further explored. Although there can be a vast number of them, there are four main subjects that we identify as the most important future research directions:

- Along this thesis, machine learning methods were considered as an alternative to traditional alignment-based method. However, the information derived from this study could be used to get further improvement in prediction performance by combining machine learning classifiers with annotation transfer methods.

-
- Classification performance are strongly influenced by the initial characterization stage that extracts numerical attributes from the protein sequences and converts them into feature vectors. In this thesis, the influence of a vast number of features was analyzed and automatic feature selection algorithms were used in order to select the most discriminant features for each GO term. However, as this work only considered global features of the whole sequence, future research directions may be focused on the dynamic characterization of the sequence in order to better explode the information of the primary structure.
 - The Cuckoo Search meta-heuristic employed in this work, can be subjected to several improvements in the selection of its hyper-parameters and stop conditions. Any improvement in this sense can contribute to a most efficient implementation of the proposed method.
 - Finally, the most recent semi-supervised algorithms are being focused on changing the data representation instead of using accurate assumptions about the actual distribution of it. Another future line of research can be focused on the proposal of novel kernels that accomplish several semi-supervised assumptions at once, in order to modify the feature space.

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