Balanced ROC analysis (BAROC) protocol for the evaluation of protein similarities

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Abstract

Identification of problematic protein classes (domain types, protein families) that are difficult to predict from sequence is a key issue in genome annotation. ROC (Receiver Operating Characteristic) analysis is routinely used for the evaluation of protein similarities, however its results—the area under curve (AUC) values—are differentially biased for the various protein classes that are highly different in size. We show the bias can be compensated for by adjusting the length of the top list in a class-dependent fashion, so that the number of negatives within the top list will be equal to (or proportional with) the size of the positive class. Using this balanced protocol the problematic classes can be identified by their AUC values, or by a scatter diagram in which the AUC values are plotted against positive/negative ratio of the top list. The use of likelihood-ratio scoring (Kaján et al, Bioinformatics, 22, 2865–2869, 2007) the bias caused by class imbalance can be further decreased.

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1. Introduction

Classification of protein sequences is a fundamental exercise in genome annotation so assessing and comparing the efficiency of sequence or structure similarity measures is a crucial task. The method of current choice is ROC (Receiver Operating Characteristic) analysis that evaluates the ranking ability of a similarity measure [1,2]. Briefly, the members of a database are ranked according to their similarity to a query using a similarity score (such as calculated by BLAST [3], Smith and Waterman [4], etc.), and a similarity score is considered efficient, if the proteins belonging to the query’s known class (true positives) are on the top of the list. The analysis is carried out by preparing a sensitivity vs. specificity plot, whose integral value, AUC (area under curve) is 1.00 if the true positives are all on the top of the list, and will tend to 0.5 if the ranking is random [1].

In the practice of bioinformatics there are two main variants of this method (Fig. 1). In the element-wise scenario, originally suggested by Gribskov and Robinson [5], one prepares a separate ranked list for all the queries, which will yield one AUC value for each query. The group-wise scenario can be applied if the database is a priori divided into distinct classes, such as domain types [6,7]. Here we prepare one single ranking for a protein group, in which we rank the members of the test set with respect to their maximal similarity to the positive train group. Then we calculate a single AUC for the given group.

If we wish to calculate a cumulative value for an entire database using either scenario, the usual method is to calculate the arithmetic average of the individual AUC values, what is equivalent to constructing a cumulative AUC plot (such as shown in Fig. 4 below) and calculating its integral value [6,7]. In this representation higher curves indicate better performance.

ROC analysis presupposes the existence of two classes, of which the query’s known class is the positive while the rest of the
database is the often much larger negative group, in other terms the positive/negative ratio ($p/n$) is often negligibly small. In order to handle this class-imbalance problem and to get a dataset of manageable sizes it is customary to cut the top list at some point, and according to Gribskov and Robinson [5] this can be done by limiting the top list so as to include $n$ negatives (where $n$ is usually taken as some plausible number like 10, 50 etc.). These are the so-called ROC$n$ (e.g. ROC10, ROC50) values that were originally proposed for the element-wise scenarios but they are also frequently used in the group-wise scenarios.

ROC analysis is considered reliable because it includes both specificity and sensitivity, so a method with a high AUC value can be expected to be a robust in a variety of conditions. While we generally agree with this view, we noticed that the ROC$n$ values can be differentially biased in various classes within a database. The result of this differential bias is that one cannot directly compare AUC values between different classes, although AUC values obtained on the same class with different methods are comparable. This is a major problem, since discovery of new protein or gene groups is perhaps the most important tasks in genome research, so there is a need to assess and compare, without class-imbalance artifacts, the predictability of protein groups. Doubtless, this assessment could be done by developing and fine-tuning classifier algorithms for each of the new candidate groups, which is, on the other hand too time consuming in view of the amounts of data to be analyzed. Our motivation was to use ROC analysis directly for the purpose. As the positive/negative ratio within a top list is analyzed. Our motivation was to use ROC analysis directly for the purpose. As the positive/negative ratio within a top list is known to influence the AUC values, we were looking for methods where this ratio can be controlled and monitored.

Here we propose a simple method wherein the number of the selected negatives included into the analysis is equal to (or proportional with) the number of the positive sample. This balanced ROC analysis provides less biased AUC values, which can be used to identify difficult groups within a database.

2. Materials and methods

2.1. Datasets and algorithm

The SCOP95 dataset used for modeling consisted of protein domain sequences taken from the SCOP database v:1.69, filtered for 95% identity [8]. For the comparison we used the Smith–Waterman algorithm [4] as coded in the EMBOSS program package [9] and applied predefined classification tasks for superfamly prediction. The classification tasks and the Smith–Waterman data were taken from record PBS0001 of the Protein Classification Benchmark collection [10], so the positive set contained members of a superfamly, one family being the +test set and the rest of the families were the +training set. We performed the evaluation using either the element-wise or the group-wise scenario. In each case we used supervised subdivision [12].

2.2. ROC calculation

Calculation of AUC was carried out as described [6,7,10,11], using top lists including a given number of negative samples.

The number of negatives ($n$) was expressed in multiples of the positive samples ($p$) participating in the restricted ranking. This normalized number $N_r$ is thus related to the positive/negative ratio, so at $N_r=1.0$, $p/n$ is lower than 1.0 for difficult classes, since in these cases there are positive samples that do not show up in the top lists, especially when less sensitive methods comparison methods are used.

For the balanced ROC (BaROC) protocol we selected the top list in such a way that it contained as many negative samples as the number of samples in the entire positive set. This corresponds to $N_r=1.0$, so the $p/n$ values are between zero and 1.0. It is equally possible to use a value of $N_r=2.0$ is i.e. to use twice as many negatives as there are positives, in that case the $p/n$ values will be between zero and 0.5. One can easily show that for any $N_r$, $p/n \leq 1/N_r$.

We mention that the BaROC protocol — same as the ROC$n$ principle suggested by Gribskov and Robinson [5]— is based on toplist truncation, so the AUC value expected for random ranking is lower than 0.5.

2.3. Likelihood-ratio scoring

In the practice of protein sequence classification the input variable of ROC analysis is a sequence similarity score, such as a BLAST or Smith–Waterman score. Recently it has been proposed that a simple likelihood-ratio approximant, LR, is a more efficient ranking indicator [12]. The LR score of a query is calculated as

$$LR = \frac{S_{max}^+}{S_{max}^-}$$

where $S_{max}^+$ and $S_{max}^-$ are the top similarities obtained between the query and members of the positive and negative groups, respectively. In contrast to a simple similarity score $S_{max}$ LR also contains information on the negative class. A similar
scoring measure, an approximant of the posterior probability \( p \) is used by the IBk program of the WEKA package [13].

\[
p = \frac{S^+_{\text{max}}}{S^+_{\text{max}} + S^-_{\text{max}}}.
\]

(2)

Since the likelihood ratio is defined as \( p/(1-p) \), it is easy to show that these two indices have in fact the same meaning and thus lead to identical ranking. In our comparisons we used the LR score, in addition to simple scoring that uses only \( S^+_{\text{max}} \). We point out that LR scoring is meaningful only in the group-wise scenario, the ranking of the element-wise scenario does not change upon the transformation to LR, so the ranking remains the same as with simple scoring.

3. Results

The heterogeneity of protein groups is a fundamental problem of protein classification. Protein groups vary in terms of the number of group-members, the length and variability of the sequences, the separation from the nearest non-member sequences etc. So we can expect that using a top list of a given length may influence the ROC\( n \) value.

Let’s take the globin superfamily a.1.1 of the SCOP95 data set as an example, and use the Smith–Waterman algorithm to compare sequence similarities. This superfamily has 103 members, and we will define a classification task wherein family a.1.1.1 (Globine-like, 5 members) will be the test group. We will calculate AUC in a group-wise scenario, using 10, 50 negatives (which corresponds to the generally used ROC10 and ROC50 scenarios), or the full dataset in the analysis. The results in Table 1 indeed show that the values are strongly dependent on the number of negatives taken into the consideration.

The phenomenon is shown graphically for the entire SCOP95 dataset in which the values represent the average of 246 classification tasks (Fig. 2). In this case we calculated ROC AUC in three different ways, i.e. using the element-wise scenario, the group-wise scenario with and without likelihood-ratio scoring.

The values are plotted against the size of the negative set expressed in multiples of the positive list. This was carried out by preparing the ranked list in the usual way and truncating it from the top when the necessary number of negatives was reached. It is conspicuous that the AUC values calculated by any of the three methods show a steeply increasing tendency if the number of negatives is below the size of the positive group. The methods using simple scoring (i.e. when the Smith–Waterman score was used for ROC analysis either in the group-wise or in the element-wise scenario) continue an increasing tendency above this threshold, but apparent that the curve of likelihood-scoring (in a group-wise scenario) does not, its value is seemingly independent of the number of negative samples in the top list. In addition, the latter method has substantially lower standard deviation values than the other two methods. In principle, the comparison of the scoring methods should not depend on the dataset, and in fact we see that the ranking order of the methods is consistently the same as we reach values of \( N^*_r > 1 \). On the other hand, the curves are crossing each other below \( N^*_r = 1.00 \), which shows that the comparison of the methods is not consistent if
We propose to use a number of negatives that is equal to or double of the number of the positives participating in the classification tasks included in the Protein Classification Benchmark Collection [10], with identical results (data not shown) which gives us hope that this protocol will be applicable to other classification methods that rank the objects according to similarities since in those cases not all the positives will show up in the top list. In the present dataset we see many such cases since the SCOP dataset is difficult for a sequence comparison method such as Smith Waterman. Simply put, regions in the BAROC scatter diagram allows one to distinguish various classes within a database. High AUC, high p/n ratio indicates tight groups, with high similarities between the members and well separated from the rest. High AUC low p/n is characteristic of less well-separated or less tight groups in which the within-group similarity is not sufficiently strong. Finally, low AUC, low p/n ratio is indicative of problem groups that are difficult to predict.

4. Discussion

Generally speaking, for statistical evaluation such as a ROC AUC calculation, it is recommended to have the same number of positive and negative samples [1,2]. This condition is never met in bioinformatics databases, we have much less positives than negatives. A correct solution would be to randomly select equal numbers from the two classes and to construct many classifiers for all cases, which is clearly too time consuming given the number of classes and data. Instead, bioinformaticians resort to the truncation of the top lists, which – as we saw in Table 1 – does not necessarily helps one to decide whether or not a low AUC value is indicative of a problematic object class or it is a class-imbalance artifact. As a way out, we propose the balanced ROC scenario which adjusts the number of negatives to the level of the size of the positive set. In this manner, classes, where the positives are within the “balanced” top list, will have a high p/n value as well as a high AUC. These classes are the easy cases for which the comparison measure (in our case the Smith–Waterman similarity score) works efficiently. In the case of difficult classes, we will have fewer positives in the top list. So we can detect the problem classes, using a scatter plot like the one shown in Fig. 4, where they will be in the region of low AUC ranking. Fig. 3 shows that the balanced ROC analysis calculated in this manner is a more stringent test than AUC50 or AUCC10, at least for the SCOP95 dataset in which there are many relatively small groups.

The balancing effect is clearly shown in Fig. 4 where we plotted the AUC value for 246 groups against the p/n ratio. In the case of ROC50, the p/n ratio is way below one (A, top). In the case of the balanced ROC (B, bottom), p/n gets near one for a number of the groups, but it can be lower in the case of distant similarities since in those cases not all the positives will show up in the top list. In the present dataset we see many such cases since the SCOP dataset is difficult for a sequence comparison method such as Smith Waterman. Simply put, regions in the BAROC scatter diagram allows one to distinguish various classes within a database. High AUC, high p/n ratio indicates tight groups, with high similarities between the members and well separated from the rest. High AUC low p/n is characteristic of less well-separated or less tight groups in which the within-group similarity is not sufficiently strong. Finally, low AUC, low p/n ratio is indicative of problem groups that are difficult to predict.
5. Simplified description of the method

We propose a ROC analysis protocol that makes it possible to single out classes in a database that are likely to be difficult to predict. The method, termed Balanced ROC (BAROC), consists of calculating an AUC value for a ranked top list which truncated so as to contain as many (or twice as many) negative objects as there are positive objects in the entire analysis. In this manner each class will be analyzed with a top list whose length depends on the size of the class. The difficult groups can then be identified by their low AUC values and/or their low positive/negative ratio within the top list. The identification is helped by a scatter plot of AUC vs. positive/negative ratio, as well as by the use of a likelihood-ratio scoring scheme (Eq. (1)), that can be efficiently used in the BAROC protocol instead of the simple similarity scores.

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References