

# Dataset Analysis Using Membership-Deviation Graph

Igtel Bayarsaikhan, Jimin Lee, and Sejong Oh

**Abstract**—Classification is one of the primary themes in computational biology. The accuracy of classification strongly depends on quality of a dataset, and we need some method to evaluate this quality. In this paper, we propose a new graphical analysis method using ‘Membership-Deviation Graph (MDG)’ for analyzing quality of a dataset. MDG represents degree of membership and deviations for instances of a class in the dataset. The result of MDG analysis is used for understanding specific feature and for selecting best feature for classification.

**Keywords**— feature, classification, machine learning algorithm.

## I. INTRODUCTION

CLASSIFICATION is one of the primary themes in computational biology and bioinformatics. In service of this task, we select a specific *dataset*, and then train *classifiers* using known classified data. The accuracy of training and classification depends on the quality of the dataset. If a dataset has separable classes, the classification accuracy of the feature may be good. But it is difficult to measure the separability of a dataset.

Feature selection [1-6] is the problem of selecting the best feature that is, ideally, necessary and sufficient to describe the target concept [7]. The objective of feature selection is to obtain a dataset characterized by (1) low dimensionality, (2) retention of sufficient information, (3) enhancement of separability in feature space for examples in different categories via the removal of effects resulting from noisy features, and (4) the comparability of features among examples in the same category [8]. Most of feature selection algorithms have an evaluation function that produces scores for candidate datasets. Let us suppose that  $F_1$  and  $F_2$  are datasets and  $E(x)$  is an evaluation function, and if  $E(F_1) > E(F_2)$ , we can expect that dataset  $F_1$  will yield better training/testing accuracy than  $F_2$ .

We can use the evaluation functions to understand and analyze specific dataset. We introduce some evaluation functions in section 2. The evaluation functions produce numerical scores, and we guess the characteristics of a dataset depending on the numerical values. If we can visualize the

vales, we can easily understand characteristics of a dataset.

In this paper, we propose a graphical analysis method, called ‘Membership-Deviation Graph (MDG)’. MDG is a scatter diagram for a class of a dataset, and based on two evaluation values, degree of membership and deviations for instances in the class. From the MDG, we can observe the distributions and accuracies of each class in a dataset. It means that we can decide about the quality of the feature.

The remainder of this paper is structured as follows. Section 2 summarizes several dataset evaluation functions. Section 3 describes the proposed Membership-Deviation Graph. Section 4 describes the results of experiments concerning MDG. We apply MDG on Yeast dataset, and the conclusions of this paper are provided in Section 5.

## II. DATASET EVALUATION FUNCTIONS

The goal of dataset evaluation is to calculate the distances among classes in the dataset. Let  $D$  be a distance function and  $D(C_1, C_2) > D(C_3, C_4)$ , thus making classes  $C_1$  and  $C_2$  more separable than  $C_3$  and  $C_4$ .

The Euclidean distance is a base of other distance functions. The Euclidean distance between the two points  $p(p_1, p_2, p_3, \dots, p_n)$  and  $q(q_1, q_2, q_3, \dots, q_n)$  is defined as follows :

$$d(p, q) = \sqrt{(p_1 - q_1)^2 + (p_2 - q_2)^2 + \dots + (p_n - q_n)^2} \\ = \sqrt{\sum_{i=1}^n (p_i - q_i)^2} \quad (1)$$

The Hausdorff distance [9] measures how far two subsets of a metric space are from one another. The Hausdorff distance between set A and set B is defined as follows:

$$H(A, B) = \max\{h(A, B), h(B, A)\} \quad (2)$$

In the equation (2), The directed function  $h(A, B)$  refers to the distance between set A and set B, and  $H(A, B)$  is also the distance between sets A and B.

J. Liang *et al.* [10] previously suggested a new feature selection algorithm based on a distance discriminant (FSDD). They used the distance among center points of categories, the standard deviation of each attribute in a feature, and the intra-set distance of categories.

RELIEF [11-13] is regarded as one of the more successful feature selection algorithms. The basic idea of RELIEF is to iteratively estimate feature weights according to their ability to discriminate between neighboring instances. In each iteration, an instance  $x$  is selected randomly, and then the

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nearest instance of it is found from the same class (NH) as well as different classes (NM). Finally, the weight value is updated by the equation:

$$w_i = w_i + |x^{(j)} - NM^{(j)}(x)| - |x^{(j)} - NH^{(j)}(x)| \quad (3)$$

As we mentioned before, above evaluation functions produce numerical values, and it is difficult to imagine the characteristics of specific feature. In the next section, we describe a new visual analysis tool, Membership-Deviation Graph.

### III. MEMBERSHIP-DEVIATION GRAPH

#### A. Summary

MDG is a two-dimensional scatter diagram as shown in Fig.1. One diagram contains information for a class in a dataset, so we may generate 7 diagrams if a dataset has 7 classes. Each point on the diagram corresponds to instance of represented class. In the diagram, the axis of X represents the deviation of distance from center point (instance) of a class, and the axis of Y represents the degree of membership of each point into the class.

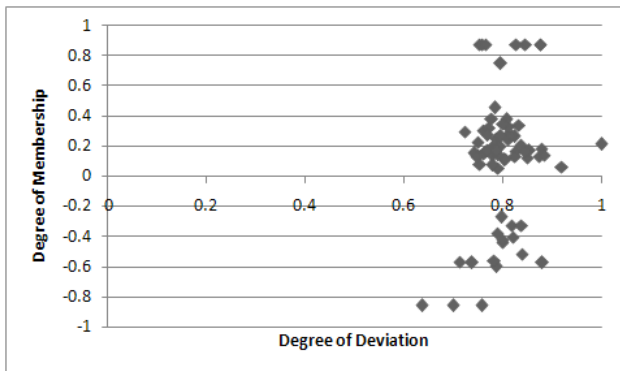


Fig. 1 An example of MDG

Fig. 2 shows the meaning of deviation. The scope of deviation of *Case A* is narrower than *Case B*. It means that the points in *Case B* are more widely scattered than one in *Case A*. Fig. 3 shows the meaning of membership. The membership of points in *Case C* is stronger than one in *Case D*. If instances in a class have strong membership and narrow scope of deviation, the class may be well-separable from other classes. We describe more detail meaning of deviation and membership in the next parts.

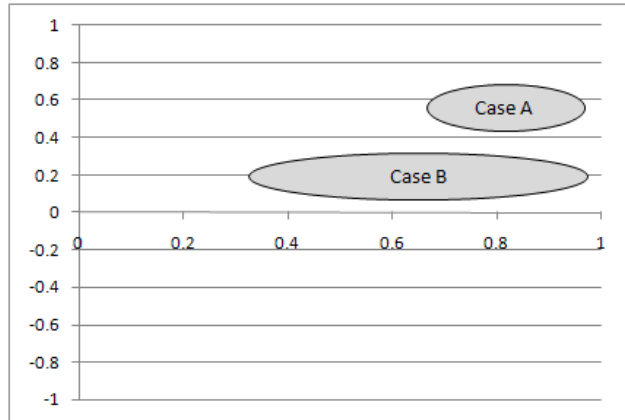


Fig. 2 Two cases of deviation

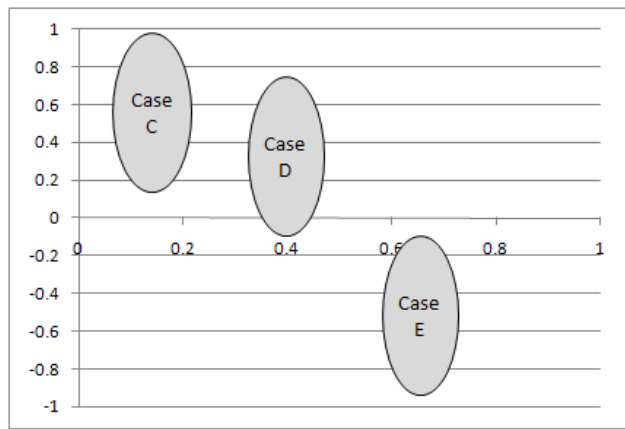


Fig. 3 Three cases of membership

#### B. Degree of Membership

A class may have many instances, but the memberships of each instance are different. In Fig. 4, for example,  $P_1$  and  $P_2$  are belongs to class  $C_1$ , but  $C_2$  is located in overlapping area with  $C_1$ . Therefore, no classifier can easily predict  $P_2$  is belongs to class  $C_1$ . We calculate degree of membership for an instance by counting the number of nearest instances that belongs to same class. If the degree of membership is 1, it means that the instance is located fully far from other classes, and can be easily classified. If the degree of membership is -1, it means that the instance is located in deeply overlapped area with other classes, and very difficult to correctly classify.

The membership function  $M(P_i)$  is defined as follows:

$$M(P_i) = \frac{Cnt(P_i)}{K} \times \frac{AvgD(SP_i)}{AvgD(SP_i) + AvgD(NP_i)} \quad (4)$$

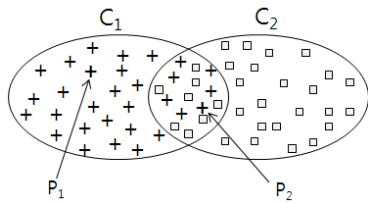


Fig. 4 Two instances in a same class

In the equation (4),  $K$  is the total number of nearest instances of instance  $P_i$ , and function  $Cnt(P_i)$  returns the number of nearest instances that belongs to same class of instance  $P_i$ . Therefore,  $\frac{Cnt(P_i)}{K}$  means the ratio of homogeneous instances of  $P_i$  over whole  $K$ -nearest neighbor instances. In the second part of equation (4),  $SP_i$  is homogeneous instances of  $P_i$  and  $NP_i$  is non-homogeneous instances of  $P_i$  over whole  $K$ -nearest neighbor instances.  $AvgD(SP_i)$  is average distance of  $SP_i$ . Important thing of  $M(P_i)$  is that it has positive value if  $P_i$  is correctly classified by KNN classifier, or it has negative value. For simplicity we normalize  $M(P_i)$  into  $[-1,1]$ .

*Degree of Deviation*

The degree of deviation for an instance has intuitive meaning and we just describe the deviation function  $DIV(P_i)$  as follows:

$$DIV(P_i) = ED(P_i, C) \tag{5}$$

In the equation (5),  $ED(P_i, C)$  is a function that returns Euclidean distance between  $P_i$  and center point of  $P_i$ 's class. Before drawing MDG, we normalize the output of  $DIV(P_i)$  into value range  $[0,1]$ .

IV. EXPERIMENTS

A. Dataset

To show usefulness of MDG, we choose Yeast dataset [14], and analyze it using MDG. Original Yeast dataset has 10 classes, but we remove a class that has a little bit instances. TABLE I summarize Yeast data set. We test  $K$ -Nearest Neighbor (KNN), Artificial Neural Network (ANN), and Support Vector Machine (SVM) classifiers, and TABLE II summarize the classification accuracies.

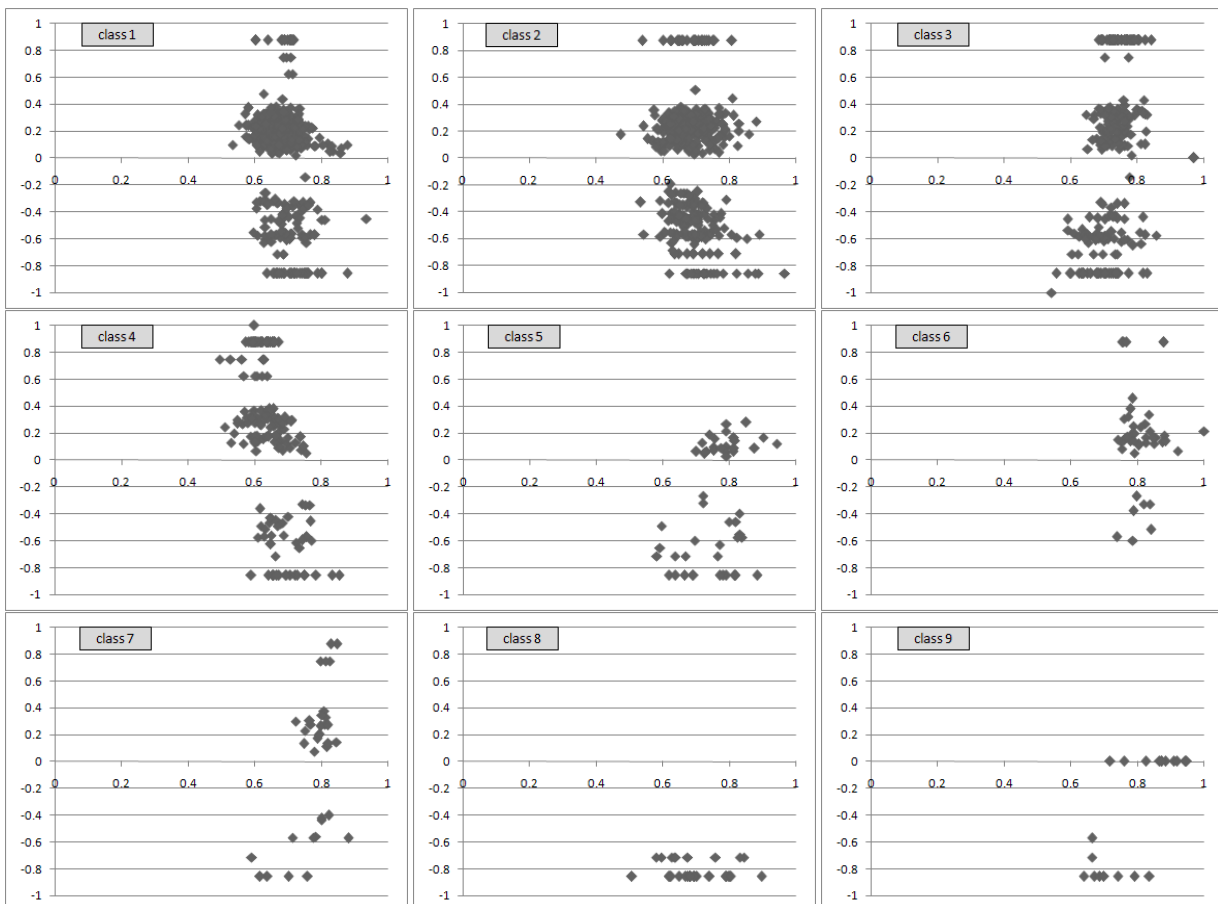


Fig. 5 Result of MDG analysis for Yeast dataset

### B. MDG Analysis Results

Fig. 5 summarizes MDG analysis results. In the case of *class 8*, membership degrees of all instances have large negative values; it means that all instances of *class 8* are located in overlapped area with other classes. Therefore, the classification accuracies produced by 3 classifiers for *class 8* are all zero. In the case of *class 6*, the ratio of instances in overlapped area is relatively small and also the scope of deviation is relatively narrow; 3 classifiers produce high classification accuracy. Classification accuracy of *class 1* is greater than *class 2* because *class 2* has more points that have negative membership value.

TABLE I  
GENERAL INFORMATION OF YEAST DATASET

No of instances	No of Classes	No of attributes
1484	9	9

TABLE II  
CLASSIFICATION ACCURACIES FOR YEAST DATASET

Class	KNN	ANN	SVM
<i>Class 1</i>	0.69	0.16	0.86
<i>Class 2</i>	0.47	0.38	0.3
<i>Class 3</i>	0.53	0.43	0.5
<i>Class 4</i>	0.65	0.66	0.68
<i>Class 5</i>	0.23	0.27	0.0
<i>Class 6</i>	0.82	0.68	0.59
<i>Class 7</i>	0.61	0.39	0.0
<i>Class 8</i>	0.0	0.0	0.0
<i>Class 9</i>	0.6	0.1	0.6
<i>Total</i>	0.57	0.31	0.53

From the MDG, we can see distributions of instances in a class, and we can compare each class in a feature. Now MDG is just observation tool, but if we process the result of MDG, we may be able to get more useful information for feature selection.

### V. CONCLUSION

MDG is a scatter diagram to show distributions of instances in a class. It contains information about degree of membership and deviation for an instance in a class. We can observe each class and compare classes of a dataset using MDG. We also can guess the approximate quality of the dataset. Comparison of the MDG results from different datasets is one of further works.

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