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Reality Spaces for Visualizing Cancer
Databases: An Evolutionary Computation
Approach **

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Hybrid Unsupervised/Supervised Virtual Reality Spaces for Visualizing Cancer Databases: An Evolutionary Computation Approach

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Abstract. This paper introduces a multi-objective optimization approach to the problem of computing virtual reality spaces for the visual representation of relational structures (e.g. databases), symbolic knowledge and others, in the context of visual data mining and knowledge discovery. Procedures based on evolutionary computation are discussed. In particular, the NSGA-II algorithm is used as a framework for an instance of this methodology; simultaneously minimizing Sammon's error for dissimilarity measures, and mean cross-validation error on a k-nn pattern classifier. The proposed approach is illustrated with an example from cancer genomics data (e.g. lung cancer) by constructing virtual reality spaces resulting from multi-objective optimization. Selected solutions along the Pareto front approximation are used as nonlinearly transformed features for new spaces that compromise similarity structure preservation (from an unsupervised perspective) and class separability (from a supervised pattern recognition perspective), simultaneously. The possibility of spanning a range of solutions between these two important goals, is a benefit for the knowledge discovery and data understanding process. The quality of the set of discovered solutions is superior to the ones obtained separately, from the point of view of visual data mining.

1 Introduction

According to the World Health Organization (WHO) <http://www.who.int/cancer/en/>, from a total of 58 million deaths in 2005, cancer accounts for 7.6 million (or 13%) of all deaths worldwide. This places cancer as one of the leading causes of death in the world, with lung cancer (the main cancer leading to mortality) accounting for 1.3 million deaths per year. Thus the importance of understanding the mechanisms of lung cancer is clear. One approach is through the rapid quantification of the gene expression levels of samples of healthy and diseased lung tissue. This new field blending the knowledge from biologists, computer scientists and mathematicians is known as Bioinformatics and is yielding large quantities of data of a very high dimensional nature that needs to be understood.

The increasing complexity of the data analysis procedures makes it more difficult for the user (not necessarily a mathematician or data mining expert), to extract useful information out of the results generated by the various techniques. This makes

graphical representation directly appealing; for which Virtual Reality (VR) is a suitable paradigm. Virtual Reality is *flexible*, it allows the construction of different virtual worlds representing *the same* underlying information, but with a different look and feel. VR allows *immersion*, that is, the user can navigate inside the data, interact with the objects in the world. VR creates a *living* experience. The user is not merely a passive observer but an actor in the world. VR is *broad and deep*. The user may see the VR world as a whole, and/or concentrate the focus of attention on specific details of the world. Of no less importance is the fact that in order to interact with a Virtual World, no mathematical knowledge is required, and the user only needs minimal computer skills. A virtual reality technique for visual data mining on heterogeneous, imprecise and incomplete information systems was introduced in [24, 25] (see also <http://www.hybridstrategies.com>).

The purpose of this paper is to explore the construction of high quality VR spaces for visual data mining using a multi-objective optimization technique applied to the understanding of a publicly available lung cancer gene expression data set. This approach provides both a solution for the previously discussed problem, and the possibility of obtaining a set of spaces in which the different objectives are expressed in different degrees, with the proviso that no other spaces could improve any of the considered criteria individually (if spaces are constructed using the solutions along the Pareto front). This strategy clearly represents a conceptual improvement in comparison with spaces computed from the solutions obtained by single-objective optimization algorithms in which the objective function is a weighted composition involving different criteria.

2 The multi-objective approach: A hybrid perspective

In order to establish a formulation of the problem based on multi-objective optimization, a set of objective functions has to be specified, representing the corresponding criteria that must be simultaneously satisfied by the solution. The minimization of a measure of similarity information loss between the original and the transformed spaces and a classification error measure over the objects in the new space can be used in a first approximation. Clearly, more requirements can be imposed on the solution by adding the corresponding objective functions. Following a principle of parsimony this paper will consider the use of only two criteria, namely, Sammon's error (Eq-3) for the unsupervised case and mean cross-validated classification error with a k-nearest neighbour pattern recognizer for the supervised case.

The proximity (or similarity) of an object to another object may be defined by a distance (or similarity) calculated over the independent variables and can be defined by using a variety of measures. In the present case a normalized Euclidean distance is chosen:

$$d_{\frac{x}{t}} = \sqrt{(1/p) \sum_{j=1}^p (x_{ij} - t_{kj})^2} \quad (1)$$

2.1 Structure Preservation: An unsupervised perspective

Examples of error measures frequently used for structure preservation are:

$$\text{S stress} = \sqrt{\frac{\sum_{i<j} (\delta_{ij}^2 - \zeta_{ij}^2)^2}{\sum_{i<j} \delta_{ij}^4}}, \quad (2)$$

$$\text{Sammon error} = \frac{1}{\sum_{i<j} \delta_{ij}} \frac{\sum_{i<j} (\delta_{ij} - \zeta_{ij})^2}{\delta_{ij}} \quad (3)$$

$$\text{Quadratic Loss} = \sum_{i<j} (\delta_{ij} - \zeta_{ij})^2 \quad (4)$$

For heterogeneous data involving mixtures of nominal and ratio variables, the Gower similarity measure [11] has proven to be suitable. The similarity between objects i and j is given by

$$S_{ij} = \frac{\sum_{k=1}^p s_{ijk} / \sum_{k=1}^p w_{ijk}}{\sum_{k=1}^p w_{ijk}} \quad (5)$$

where the weight of the attribute (w_{ijk}) is set equal to 0 or 1 depending on whether the comparison is considered valid for attribute k . If $v_k(i), v_k(j)$ are the values of attribute k for objects i and j respectively, an invalid comparison occurs when at least one them is missing. In this situation w_{ijk} is set to 0.

For quantitative attributes (like the ones of the datasets used in the paper), the scores s_{ijk} are assigned as

$$s_{ijk} = 1 - |v_k(i) - v_k(j)| / R_k$$

where R_k is the range of attribute k . For nominal attributes

$$s_{ijk} = \begin{cases} 1 & \text{if } v_k(i) = v_k(j) \\ 0 & \text{otherwise} \end{cases}$$

This measure can be easily extended for ordinal, interval, and other kind of variables. Also, weighting schemes can be incorporated for considering differential importance of the descriptor variables.

2.2 Multi-objective Optimization Using Genetic Algorithms

An enhancement to the traditional evolutionary algorithm[1], is to allow an individual to have more than one measure of fitness within a population. One way in which such an enhancement may be applied, is through the use of, for example, a weighted sum of more than one fitness value [3]. Multi-objective optimization, however, offers another possible way for enabling such an enhancement. In the latter case, the problem arises for the evolutionary algorithm to select individuals for inclusion in the next population, because a set of individuals contained in one population exhibits a Pareto Front[19] of best current individuals, rather than a single best individual. Most [3] multi-objective algorithms use the concept of dominance to address this issue.

A solution $\tilde{x}_{(1)}$ is said to dominate [3] a solution $\tilde{x}_{(2)}$ for a set of m objective functions $\langle f_1(\tilde{x}), f_2(\tilde{x}), \dots, f_m(\tilde{x}) \rangle$ if

1. $\tilde{x}_{(1)}$ is not worse than $\tilde{x}_{(2)}$ over all objectives.
For example, $f_3(\tilde{x}_{(1)}) \leq f_3(\tilde{x}_{(2)})$ if $f_3(\tilde{x})$ is a minimization objective.
2. $\tilde{x}_{(1)}$ is strictly better than $\tilde{x}_{(2)}$ in at least one objective. For example, $f_6(\tilde{x}_{(1)}) > f_6(\tilde{x}_{(2)})$ if $f_6(\tilde{x})$ is a maximization objective.

One particular algorithm for multi-objective optimization is the elitist non-dominated sorting genetic algorithm (NSGA-II) [7], [6], [5], [3]. It has the features that it *i*) uses elitism, *ii*) uses an explicit diversity preserving mechanism, and *iii*) emphasizes the non-dominated solutions.

2.3 Original Study

Gene expressions were compared in [21] for severely emphysematous lung tissue (from smokers at lung volume reduction surgery) and normal or mildly emphysematous lung tissue (from smokers undergoing resection of pulmonary nodules). The original database contained 30 samples (18 severe emphysema, 12 mild or no emphysema), with 22,283 attributes. Genes with large detection *P*-values were filtered out, leading to a data set with 9,336 genes, that were used for subsequent analysis. Nine classification algorithms were used to identify a group of genes whose expression in the lung distinguished severe emphysema from mild or no emphysema. First, model selection was performed for every algorithm by leave-one-out cross-validation, and the gene list corresponding to the best model was saved. The genes reported by at least four classification algorithms (102 genes) were chosen for further analysis. With these genes, a two-dimensional hierarchical clustering using Pearson's correlation was performed that distinguished between severe emphysema and mild or no emphysema. Other genes were also identified that may be causally involved in the pathogenesis of the emphysema. The data was obtained from http://www.ncbi.nlm.nih.gov/projects/geo/gds/gds_browse.cgi?gds=737.

2.4 Experimental Settings

Each sample in this study is a vector in a high dimensional space, and therefore, direct inspection of the structure of this data, and of the relationship between the descriptor variables (the genes) and the type of sample (normal or cancer), is impossible. Moreover, within the collection of genes there is a mixture of potentially relevant genes with others which are irrelevant, noisy, etc. The need of simultaneously finding a visual representation (3D) respecting (as much as possible) the set of object interrelationships as defined by the original attributes, and the construction of a new feature space effectively differentiating the two classes of objects present, makes this problem suitable for a multi-objective optimization approach.

The collection of parameters describing the application of the NSGA-II algorithm is shown in Table-1. A modest population size and number of generations were used, with a relatively high mutation probability in order to enable richer genetic diversity. Randomization of the set of data objects was applied in order to reduce the bias in the composition of the cross-validated folds by providing a more even class distribution between successive training and test subsets. The number of folds was set in consideration of the sample size.

Table 1: Experimental settings for computing the pareto-optimal solution approximations by the multi-objective genetic algorithm (PGAPack extended by NSGA-II).

population size	100	number of generations	500
chromosome length	90 (= 3 · 30)	ga seed	101
No. new inds. in ($i + 1$ st) pop.	20	objective functions should be minimized	
chromosome data representation	real	crossover probability	0.8
crossover type	uniform (prob. 0.6)	mutation probability	0.4
mutation type	gaussian	selection type	tournament
tournament probability	0.6	mutation and crossover	yes
population initialization	random, bounded	lower bound for initialization	-2
upper bound for initialization	2	fitness values	raw
stopping criteria	maximum iterations	restart ga during execution	no
parallel populations	no		
number of objectives	2	number of constraints	0
pre-computed diss. matrix	Gower dissimilarity		
evaluation functions	mean cross-validated k-nn error and Sammon error		
cross-validation (c.v.)	5 folds	randomize before c.v.	yes
knn seed	101	k nearest neighbors	3
non-linear mapping measure	Sammon	dimension of the new space	3

2.5 Results

The set of non-dominated solutions obtained by the NSGA-II algorithm is shown in the scatter plot of Fig-1(a), where the horizontal axis is the mean cross-validated knn error and the vertical axis the Sammon error. The approximate location of the Pareto front is defined by the convex polygon joining the solutions provided by chromosomes 2, 1, 10, etc. Chromosome 2 defines a space with a perfect resolution of the supervised problem in terms of the “*no or mild emphysema*” and “*severe emphysema*” classes (knn error = 0), but at the cost of a severe distortion of the space. Whereas, chromosome 1 approximates a pure unsupervised solution (with low Sammon error). Its classification error is large indicating that few non-linear features preserving the similarity structure lacks classification power. This may be due to the large amount of attribute noise, redundancy, and irrelevancy within the set of 22, 283 original genes.

Clearly, it is impossible to represent virtual reality spaces on a static medium. However, a composition of snapshots of the VR spaces using the solutions along the Pareto front approximation is shown in Fig-1(b-d). Different mappings (even with important differences from the point of view of the mapping error) lead to similar 3D visual representations, which indicates good reproducibility of the solutions. The similarities are associated to the main distributions of the clouds of points, which are preserved, while there might be local discrepancies with respect to the placement of some objects.

A solution satisfying classification error as much as possible (actually with 0-error) is shown in Fig-1(b) where both classes are separated into 2 main clouds of points and a distinct point, Object 6, positioned separately from the clouds. It can be seen that Object 6 is positioned relatively differently in the spaces that comprise the best Sammon error Fig-1(d) and tradeoff between the classification error objective and Sammon er-

ror objective Fig-1(c). This is why, visually, the latter space represents a compromised solution between the two goals and a tradeoff between the two objective functions. It should be remembered that the class information is not used at all for computing the spaces. Chromosome 10, according to Fig-1(a) and Fig-1(c), can be considered to be the best multi-objective compromised solution in which both error criteria are simultaneously as low as possible. It shows a reasonable class discrimination with a non-large similarity structure distortion, which is a very meaningful result.

3 Conclusions

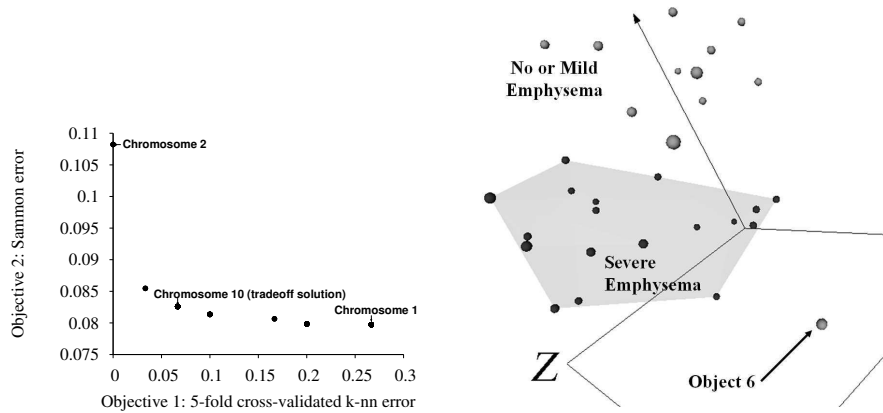
A multi-objective optimization approach was introduced for the problem of computing virtual reality spaces in the context of visual data mining and knowledge discovery applied to relational structures (e.g. databases). The multi-objective procedure was based on NSGA-II using two objective functions representative of unsupervised and supervised criteria (mean cross-validated knn error as a measure of miss-classification, and Sammon error as a measure of similarity structure loss). This methodology was applied to the analysis of high dimensional genomic data collected in the framework of Lung cancer research. A Pareto front approximation was recognizable from within the solutions provided by the final population. Selected solutions from along that approximation were used for the construction of a sequence of visualizations showing the progression from spaces with complete class separation and poor similarity preservation to spaces with reversed characteristics. A solution with a reasonable compromise between the two criteria was identified and clearly contained properties of both extreme solution spaces. These research results, although preliminary, showed large potential and further investigation is required.

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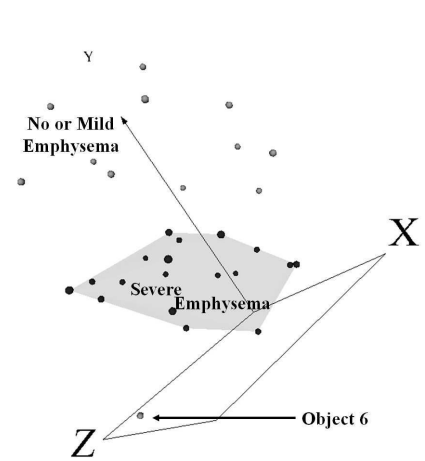
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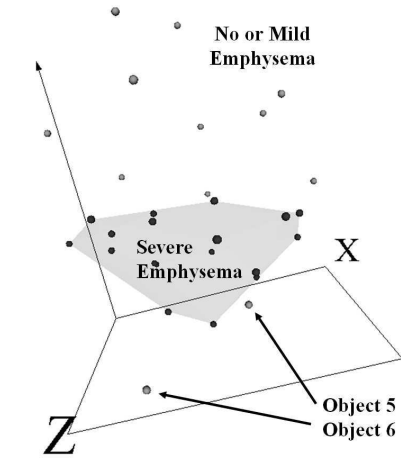


(a) Front obtained by multi-objective optimization (NSGA-II) that approximates the true Pareto Front.

(b) Chromosome 2 (5-fold CV k-nn Error: 0.0000, Sammon Error: 0.1082)



(c) Chromosome 10 (5-fold CV k-nn Error: 0.0667, Sammon Error: 0.0826)



(d) Chromosome 1 (5-fold CV k-nn Error: 0.2667, Sammon Error: 0.0797)

Fig. 1: Set of 100 multi-objective solutions. Those along the Pareto front approximation progressively span the extremes between minimum classification error and minimum dissimilarity loss. 3 solutions were selected and snapshots of VR spaces computed. Geometries: “light grey spheres” = no or mild emphysema samples, “dark grey spheres encased within a convex hull” = severe emphysema samples. Behavior = static.

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