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¹O. I. Chumachenko,²V. S. Gorbatiuk

SOFT CLUSTERING ALGORITHM BASED ON SEPARATING HYPERSURFACES

^{1,2}National Technical University of Ukraine “Ihor Sikorsky Kyiv Polytechnic Institute”, Kyiv, UkraineE-mails: ¹chumachenko@tk.kpi.ua, ²vladislav.horbatiuk@gmail.com

Abstract—A new “soft” clustering algorithm is proposed based on the use of artificial neural networks as models of hypersurfaces that separate clusters. The algorithm allows to solve the problem of soft clusterization as a problem of smooth nonlinear function optimization and, therefore, to apply the entire mathematical apparatus of nonlinear optimization, which has evolved significantly in recent years.

Index Terms—Clustering; artificial neural networks; soft clustering; nonlinear optimization.

I. INTRODUCTION

The clustering problem is probably the most well-known problem from the “unsupervised learning” class of problems [1] – that is problems in which certain additional information from “teacher” that can facilitate the learning process for the system is not provided for input examples. This problem arises in solving many practical problems in various areas, such as: image processing, data compression, bioinformatics and others.

II. PROBLEM STATEMENT

The clustering problem [2] that is considered in this paper can be described as follows:

Having a set of examples $X = (\bar{x}_1, \dots, \bar{x}_n)$ where each example is a vector in space R^d , and given number of clusters $K \in N$, we need to set a certain cluster number $k = 1, \dots, K$ for each example so that the resulting vector of cluster numbers $\vec{k} = [k_1, \dots, k_n]^T$ minimizes a certain criterion $CR(\vec{k}, X)$:

$$\vec{k}^* = \arg \left\{ \min_{\vec{k}} \{CR(\vec{k}, X)\} \right\}.$$

III. AN OVERVIEW OF EXISTING METHODS

As it is known, the clustering problem even for simple case when the loss function is a total Euclidean distance of the points to the center of their cluster is NP hard [3], [4] and thus there is no general algorithm that will surely find the optimal vector of clusters numbers \vec{k}^* and will not require an exponentially increasing number of calculations with an increase in the number of examples. Because of this, various heuristic methods are used to solve this problem approximately. Let us consider the main

known methods of clusterization and their disadvantages.

K-means clustering [5]. It is the most famous and one of the simplest clustering algorithms. Informally, it can be described as follows: at first, initial cluster centers are selected in certain way (often randomly), after which iterations are executed until the algorithm’s stopping condition is met, where each iteration consists of 2 steps: the step of finding the closest current cluster center for each example, and the step of calculating the new cluster center – as the mean value of all examples for which the current center of this cluster was the closest. Stopping condition is the equality of new and current clusters’ centers. Formally, the algorithm consists of the following steps:

1. The starting centers of clusters are set as randomly selected unique K examples:

$$c_j^{(0)} := x_{r_j}, j \in \{1, \dots, K\}, r_j \in \{1, \dots, n\}; \forall t \in \{1, \dots, K\},$$

$$\forall j \in \{1, \dots, K\} : t \neq j \rightarrow r_j \neq r_t$$

2. $it := 0$.

3. For each example $x_i, i \in \{1, \dots, n\}$ the center of the cluster closest to it is located and memorized:

$$nc_i := \arg \min_{j \in \{1, \dots, K\}} \{ |x_i - c_j^{(it)}| \}$$

4. New cluster centers are calculated as the mean of all examples for which the current cluster center was the closest:

$$c_j^{(it+1)} := \frac{1}{n_j} \sum_{i:nc_i=j} x_i, n_j = \sum_{i:nc_i=j} 1, j \in \{1, \dots, K\}$$

5. If at least for one value $j \in \{1, \dots, K\}$ the new center of the cluster is different from the current one –

$c_j^{(it+1)} \neq c_j^{(it)}$ – then $it := it + 1$ is assigned and a new iteration of the algorithm is performed starting from step 3. Otherwise, the algorithm stops, and the current distribution of examples between clusters along with the cluster centers are the algorithm outputs.

The algorithm tries to minimize the criterion of total mean distance between points in one cluster:

$$CR(\vec{k}, X) = \sum_{i=1}^K \sum_{\vec{x}_j: k_j=i} \|\vec{x}_j - \vec{\mu}_i\|^2.$$

The main drawbacks of the algorithm are:

1) susceptible to getting stuck in a local optimum – depending on the initial cluster centers the algorithm can stop at various local optima, not finding globally optimal centers (although it is clear that no algorithm that does not require exponentially increasing number of calculations while increasing the number of examples can find the globally optimal centers unless $P = NP$);

2) the algorithm prefers clusters with approximately same number of examples;

3) it is clear that the application of the algorithm to the data set where clusters do not meet the algorithm “expectations” – i.e. is not similar to the spherical areas that are separated in space, usually gives poor results.

Hierarchical clustering algorithms [6]. Algorithms of this group are building a certain hierarchy of clusters; there are 2 main approaches for building a hierarchy: **agglomerative** when the algorithm starts with each example is in its own cluster and the clusters are gradually merged, and **divisive** when at the beginning all examples belong to one cluster and the division of one cluster into several smaller clusters is gradually done.

The main disadvantages of this algorithms family are:

A. It is not always possible to clearly define a global criterion that is minimized by this algorithm.

B. A certain locally optimal clustering is often obtained because of the “greedy” nature of most of the algorithms of this family.

The Kohonen self-organizing map [7] (SOM). Self-organizing map is an artificial neural network [8] “suitable” for unsupervised learning, and during learning the network tries to somehow approximate distribution of input examples. This approximation is achieved via “positioning” a certain number of network’s output neurons in the space of input examples so that all neurons tend to be positioned closer to given input examples – often finding centers of examples’ clusters in the process. Thus, after all output neurons are positioned, the clustering of input ex-

amples can be carried out by finding the closest neuron for each example, and assigning to one cluster all examples for which one certain common neuron turned out to be the closest one.

The main shortcomings of the SOM are:

1) For the original SOM “algorithm” it is unclear what optimization criteria is minimized while network neurons are positioned in space of input examples.

2) The result of the SOM algorithm depends on certain parameters the values of which need to be carefully chosen.

3) In terms of clustering problem there is no evidence that the SOM gives fundamentally better results in comparison with other clustering algorithms, such as k -means.

IV. SOFT CLUSTERING ALGORITHM BASED ON SEPARATING HYPERSURFACES

A new clustering algorithm with the following properties is proposed:

a) it has a clearly defined criterion which is minimized during the execution of the algorithm;

b) the optimization criterion is a differentiable function of the parameters according to which the optimization is performed;

c) it theoretically allows to find clusters that are separated by the arbitrarily complex hypersurface.

The algorithm can be applied when the criterion $CR(\vec{k}, X)$ has the following form:

$$CR(\vec{k}, X) = \sum_{i=1}^K f(X, \vec{k}, i),$$

where $f(X, \vec{k}, i)$ is the function which is either completely continuous function of X or its only discontinuity comes from using the indicator function “ $1(\vec{k}, \vec{x}_j, i) = 1$, if $k_j = 1, 0$ otherwise”. For example, if we consider the criterion of total mean distance between points in one cluster which is used by the k -means algorithm, then its $f(X, \vec{k}, i)$ can be written as follows:

$$f(X, \vec{k}, i) = \frac{1}{\sum_{\vec{x}_j} 1(\vec{k}, \vec{x}_j, i)} \cdot \sum_{\vec{x}_q, \vec{x}_w} 1(\vec{k}, \vec{x}_q, i) \cdot 1(\vec{k}, \vec{x}_w, i) \cdot \|\vec{x}_q - \vec{x}_w\|^2.$$

Let us consider the simple case where we have only two clusters. We perform the “softening” of the original problem [9] by introducing the continuous function $k(\vec{x}) \in [0, 1]$ which for each example \vec{x} will

return the probability of this example belonging to the cluster 1; respectively, the value $1-k(\vec{x})$ represents the probability of this example belonging to the cluster 0. In this case, the indicator functions are replaced by the value $k(\vec{x})$ and, for example, the “softened” variant of the total mean distance between points in one cluster will be as follows:

$$CR(k, X) = \frac{1}{\sum_{\vec{x}_j} [1 - k(\vec{x}_j)]} \cdot \sum_{\vec{x}_q, \vec{x}_w} \left\{ [1 - k(\vec{x}_q)][1 - k(\vec{x}_w)] \|\vec{x}_q - \vec{x}_w\|^2 \right\} + \frac{1}{\sum_{\vec{x}_j} k(\vec{x}_j)} \sum_{\vec{x}_q, \vec{x}_w} \left\{ k(\vec{x}_q)k(\vec{x}_w) \|\vec{x}_q - \vec{x}_w\|^2 \right\},$$

wherein the first component determines the contribution of the cluster with number 0 and the second – the cluster with number 1.

If we have a certain model of the hypersurface that separates the clusters in the form of a function $k(\vec{x}; \vec{w}) \in [0, 1]$ that is a differentiable function of a certain parameters vector \vec{w} – then criterion $CR(k, X)$ will also be a differentiable function of vector \vec{w} and thus for its minimization it is possible to use the entire apparatus of nonlinear continuous differentiable functions minimization which has recently been developing very rapidly. Thus, the “softened” version of clustering problem for 2 clusters can be solved as a continuous nonlinear optimization problem (if the input criteria satisfies the described above condition), for example through the use of a certain gradient descent algorithm modification.

To solve the softened version of clustering problem into K clusters we can use the “one versus all” approach – first we divide all the examples into 2 clusters, after which we select a cluster with a larger “partial criterion” value $f(X, \vec{k}, i)$, and divide it into 2 clusters and so on until we get the required number of clusters.

Perhaps the simplest function that can be selected as a model of a hypersurface separating the clusters is

a logistic sigmoid function: $k(\vec{x}; \vec{w}) = \frac{1}{1 + e^{-\vec{w}^T \vec{x}}}$. In

essence, such a model will be a certain approximation of the separating hyperplane which is determined by the parameters vector \vec{w} - for examples $\vec{x} : \vec{w}^T \vec{x} > b, b > 0$ the value of the model will be approximately equal to 1; for examples $\vec{x} : \vec{w}^T \vec{x} < -b$ – approximately equal to 0, and for examples that are “close” to the hyperplane – i.e. those for which

$-b < \vec{w}^T \vec{x} < b$ the value of the model will smoothly grow from 0 to 1 with the “transition” from one side to the other (and for all examples $\vec{x} : \vec{w}^T \vec{x} = 0$ which are located on the hyperplane, the model value is equal to 0.5). That is, using such a model when minimizing the criterion, we will try to separate all the examples by the “almost linear” hypersurface into 2 clusters so that the total mean distance of these clusters will be minimal.

Obviously, such a model is very simple and will not work well if the clusters in the existing set of examples are not linearly separate. In this case, we need more complicated models of the separating hypersurface, and we know what suits very well as such models – the neural networks. The only limitation is that the network's output should be in range $[0, 1]$ but to achieve this it is enough to pass the network output through the aforementioned logistic sigmoid function.

Thus, we obtain the following soft clustering general algorithm based on a certain separating hypersurface model $k(\vec{x}; \vec{w}) \in [0, 1]$.

Algorithm inputs:

- examples set $X = (x_1, \dots, x_n)$, $x_i \in R^d$, $i = 1, \dots, n$;
- number of clusters K , into which we need to split all the examples;
- some neural network that defines the model of hypersurface separating the clusters $k(\vec{x}; \vec{w}) \in [0, 1]$;
- criterion $CR(\vec{k}, X)$, which needs to be minimized, and which satisfies the above described condition.

Steps of the algorithm:

- 1) Obtaining the softened criterion $CR(\vec{k}, X) \rightarrow SCR(\vec{w}, X)$.
- 2) The whole set of examples is split into 2 clusters. In order to do this:
 - the initial vector of network parameters \vec{w}_0 is randomly generated;
 - a certain modification of the gradient descent is performed to minimize the value of the criterion $SCR(\vec{w}, X)$;
 - as a result, we get the configured parameters vector \vec{w}_f ;
 - all examples are divided into 2 clusters – those examples for which the model values $k(\vec{x}; \vec{w}_f) < 0.5$ are selected into cluster 0, all other examples (i.e. those for which $k(\vec{x}; \vec{w}_f) \geq 0.5$) – in cluster 1.
- 3) If the current number of clusters $< K$ then for both clusters the “partial” criterion value $f(X, \vec{k}, i)$

is calculated and the cluster having bigger value $f(X, \vec{k}, i)$ is chosen as a new set of examples for further separation into clusters, and the algorithm's execution continues from step 1. Otherwise, obtained K clusters are returned as the result of algorithm's execution.

V. CONCLUSION

A new soft clustering algorithm is introduced, which allows to:

- solve the original problem by minimizing certain differentiable criteria, thus making it possible to use all the recent developments in nonlinear optimization;
- find clusters that are separated by some complex hypersurface.

For further development of the algorithm, the application of softmax-layer [10] to perform clustering immediately into K clusters instead of using the “one versus all” approach should be considered. In addition, the number of clusters K is not known beforehand in many practical problems, and some way to automatically find “best” number of clusters would definitely be a promising improvement of current algorithm.

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Chumachenko Olena. Candidate of Science (Engineering). Associate Professor. Technical Cybernetic Department, National Technical University of Ukraine “Ihor Sikorsky Kyiv Polytechnic Institute,” Kyiv, Ukraine.

Education: Georgian Polytechnic Institute, Tbilisi, Georgia, (1980).

Research area: system analysis, artificial neuron networks.

Publications: more than 60 papers.

E-mail: chumachenko@tk.kpi.ua

Gorbatiuk Vladyslav. Post-graduate student.

Technical Cybernetic Department, National Technical University of Ukraine “Ihor Sikorsky Kyiv Polytechnic Institute,” Kyiv, Ukraine.

Education: National Technical University of Ukraine “Ihor Sikorsky Kyiv Polytechnic Institute” (2017).

Research area: deep learning, forecasting methods.

Publications: 10.

E-mail: vladislav.horbatiuk@gmail.com

О. І. Чумаченко, В. С. Горбатюк. Алгоритм м'якої кластеризації на основі розділяючих гіперповерхонь

Запропоновано новий алгоритм «м'якої» кластеризації на основі використання штучних нейронних мереж як моделей гіперповерхонь, що розділяють кластери. Алгоритм дозволяє розв'язувати задачу м'якої кластеризації як задачу оптимізації гладкої нелінійної функції, а отже, застосовувати весь математичний апарат нелінійної оптимізації, який суттєво розвинувся за останні роки.

Ключові слова: кластеризація; штучні нейронні мережі; м'яка кластеризація; нелінійна оптимізація.

Чумаченко Олена Іллівна. Кандидат технічних наук. Доцент.

Кафедра технічної кібернетики, Національний технічний університет України «Київський політехнічний інститут ім. Ігоря Сікорського», Київ, Україна.

Освіта: Грузинський політехнічний інститут, Тбілісі, Грузія, (1980).

Напрямок наукової діяльності: системний аналіз, штучні нейронні мережі.

Кількість публікацій: більше 60 наукових робіт.

E-mail: chumachenko@tk.kpi.ua

Горбатюк Владислав Сергійович. Аспірант.

Національний технічний університет України «Київський політехнічний інститут ім. Ігоря Сікорського», Київ, Україна.

Освіта: Національний технічний університет України «Київський політехнічний інститут ім. Ігоря Сікорського», Київ, Україна, (2017).

Напрямок наукової діяльності: глибоке навчання, методи прогнозування.

Кількість публікацій: 10.

E-mail: vladislav.horbatiuk@gmail.com

Е. И. Чумаченко, В. С. Горбатюк. Алгоритм мягкой кластеризации на основе разделяющих гиперповерхностей

Предложен новый алгоритм «мягкой» кластеризации на основе использования искусственных нейронных сетей как моделей гиперповерхностей, разделяющих кластеры. Алгоритм позволяет решать задачу мягкой кластеризации как задачу оптимизации гладкой нелинейной функции, а следовательно, применять весь математический аппарат нелинейной оптимизации, который существенно развился за последние годы.

Ключевые слова: кластеризация; искусственные нейронные сети; мягкая кластеризация; нелинейная оптимизация.

Чумаченко Елена Ильинична. Кандидат технических наук. Доцент.

Кафедра технической кибернетики, Национальный технический университет Украины «Киевский политехнический институт им. Игоря Сикорского», Киев, Украина.

Образование: Грузинский политехнический институт, Тбилиси, Грузия, (1980).

Направление научной деятельности: системный анализ, искусственные нейронные сети.

Количество публикаций: более 60 научных работ.

E-mail: chumachenko@tk.kpi.ua

Горбатюк Владислав Сергеевич. Аспірант.

Национальный технический университет Украины «Киевский политехнический институт им. Игоря Сикорского», Киев, Украина.

Образование: Национальный технический университет Украины «Киевский политехнический институт им. Игоря Сикорского», Киев, Украина, (2017).

Направление научной деятельности: глубокое обучение, методы прогнозирования.

Количество публикаций: 10.

E-mail: vladislav.horbatiuk@gmail.com