

Mining Biomedical Images with Density-based Clustering

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Abstract

Density-based clustering algorithms have recently gained popularity in the data mining field due to their ability to discover arbitrary shaped clusters while preserving spatial proximity of data points. In this work we adapt a density-based clustering algorithm, DBSCAN, to a new problem domain: Identification of homogenous color regions in biomedical images. Examples of specific problems of this nature include landscape segmentation of satellite imagery, object detection and, in our case, identification of significant color regions in images of skin lesions (tumors). Automated outer and inner boundary segmentation is a key step in segmentation of structures such as skin lesions, tumors of breast, bone, and brain. This step is important because the accuracy of the subsequent steps (extraction of various features, post-processing) crucially depends on the accuracy of this very first step. In this paper, we present an unsupervised approach to segmentation of pigmented skin lesion images based on DBSCAN clustering algorithm. The color regions identified by the algorithm are compared to those identified by the human subjects and the Kappa coefficient, a statistical indicator of computer-human agreement, is found to be significant.

1. Introduction

In this work we examine the applicability of a spatial data mining method, density-based clustering, to a novel domain: Biomedical image data mining. In particular, we explore whether a specific density-based spatial clustering method, DBSCAN, can be adapted to solve one of the essential problems of image data mining, that is extraction of key features based on an accurate identification of significant regions (objects or segments) in an image. Density-based clustering methods have recently attracted interest due to their

desirable properties such as being able to discover arbitrary shapes and requiring fewer parameters at the start of the process than other comparable approaches.

Data preparation is typically the least formalized, the most domain-dependent, and the most time consuming part of the knowledge discovery process [3,7]. The first step in the preparation of biomedical image data is typically the identification (segmentation) of significant objects such as tumors or tissue fragments. Identification/segmentation is crucial for the subsequent mining tasks, because accurate extraction of significant features relies on this step.

In this paper, we present an unsupervised approach to segmentation of skin lesion images based on DBSCAN (Density Based Spatial Clustering of Applications with Noise) algorithm. Our method allows separation of the healthy skin from the lesion and identification of the subregions inside the lesion with variegated coloring. The number of color regions identified can be fed into further classification and association mining tasks.

The rest of the paper is organized as follows: We first give an introduction to the density-based clustering method. We then discuss our adaptation of this method to the particular domain of skin lesion images. We describe the preprocessing steps before the image is submitted to the clustering module. We then explain the post-processing step which produces a region allocation that is congruent with human perception. Next we describe our experimental evaluation methodology and the experimental results. After a discussion of related work we conclude the paper with a summary of lessons learned and future research.

2. Density-based Spatial Clustering

Cluster analysis has been widely used in various disciplines such as pattern recognition, computer vision, and data mining [9]. There are numerous

clustering algorithms including hierarchical, nearest-neighbor, fuzzy, density-based, grid-based etc. The reader is referred to [8] for further information on various clustering algorithms. In this study we apply a density-based clustering method, DBSCAN, to image segmentation problem. Here we briefly introduce the DBSCAN algorithm.

DBSCAN (Density Based Spatial Clustering of Applications with Noise) is a density-based clustering algorithm that is designed to discover clusters and noise in a spatial data set [4]. It has two major parameters: *Eps* and *MinPts*. The neighborhood of an object within a radius *Eps* is called the *Eps-neighborhood* of the object. If the *Eps-neighborhood* of an object contains at least *MinPts* objects, then this object is called a core object. To find a cluster, DBSCAN starts with an arbitrary object *o* in the data set. If the object *o* is a core object w.r.t. *Eps* and *MinPts*, a new cluster with *o* as the core object is created. The algorithm continues growing the clusters by adding to the clusters all objects that are density-reachable from the core object.

GDBSCAN (Generalized DBSCAN) algorithm, generalizes the two parameters of the DBSCAN algorithm so that it can cluster point objects as well as spatially extended objects according to both their spatial and non-spatial attributes [12]. The neighborhood of an object is defined by a binary predicate *NPred*. The *NPred-neighborhood* of an object is defined as the set of objects for which the predicate *NPred* is true. A second predicate *MinWeight* of a set of objects *S*, is defined such that it is true if the weighted cardinality of the set *S*, $wCard(S)$, is greater than or equal to the minimum cardinality, *MinCard*.

3. Color Image Segmentation using DBSCAN

In this work, we follow a region-based segmentation approach. The goal is to partition an image into disjoint regions that correspond to healthy skin and subregions inside lesion.

Figure 1 shows the flowchart of the segmentation procedure. There are two major steps involved. First, the image is split into smaller regions until all the regions meet the homogeneity criteria determined by the threshold for splitting. Then, these regions are merged to form the final regions by the DBSCAN algorithm.

The segmentation algorithm employs an iterative, coarse-to-fine strategy. The aim of the first iteration is to identify the lesion border. During the following iterations, the algorithm moves towards the inside of

the lesion and identifies subregions with variegated coloring. This approach allows the algorithm to dynamically adjust the parameters during various stages of the procedure.

The following subsections will describe the steps involved in the segmentation procedure.

3.1. Preprocessing

Skin lesion images often contain extraneous artifacts such as skin texture and hair that make the segmentation more difficult. In order to reduce the effects of these artifacts, we have smoothed the images with a 5 x 5 median filter [6]. This filter has the advantage of preserving the edges well enough for accurate border detection [13].

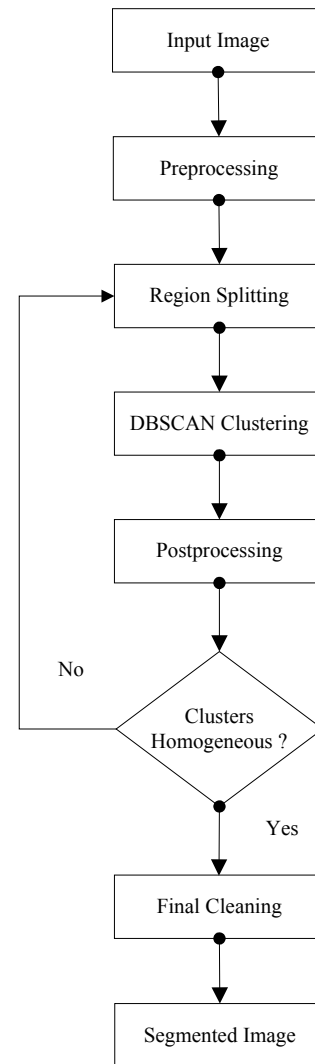


Figure 1. Flowchart of the segmentation procedure

3.2. Region Splitting

Region splitting is a top-down process. A region is split into four subregions if it fails to satisfy the homogeneity criterion. Splitting process starts from the highest-level region which is the whole image to be segmented. Each region in the image is represented by its mean color calculated in the RGB color space. To determine the homogeneity of a region, the Euclidean distances between the region and each of its four potential subregions are computed. If any of the color distances computed above is greater than the threshold for splitting, the region will be split. Otherwise, the region remains unchanged. The splitting process goes on recursively until each region is either found to be homogeneous or too small to be split further.

The threshold for splitting is a very important parameter. If it is set too high the accuracy of the segmentation will decrease. The result will show rectangular borders around the segments. On the other hand, if it is set too low, the image might be over-segmented with most of the segments being as small as a few pixels. The program automatically determines the threshold for splitting as follows.

First, we determine the color of the healthy skin. This is done by taking four windows of size 10×10 pixels from four corners of the image and calculating the median color of the pixels. We use this median color as the color of the healthy skin. Note that median color is used instead of mean to reduce the interference of hairs [16]. Then, we divide the image into 100 subregions and use each of the subregions as a sample of the image. Finally, we calculate the color distances between the healthy skin color and each of these samples and sort these values. The threshold for splitting is determined from the significant gaps among these values. In the following discussion, the distance between the healthy skin color and a sample n will be simply referred to as the value of sample n .

Let us demonstrate the threshold selection using Figure 2. Here, there is a significant gap between the values of sample n and sample $n+1$. If the amount of this gap is greater than a threshold T_G and the value of sample n is also greater than a threshold T_C , we let the threshold for splitting to be the average of values of sample n and sample $n+1$. The thresholds T_G and T_C are determined empirically from a set of representative images as described in the Experimental Results section.

Once the splitting process is completed, a graph is constructed to represent the regions and their connectivity. Two regions are considered connected if

they share a common boundary. The graph is represented by an adjacency matrix.

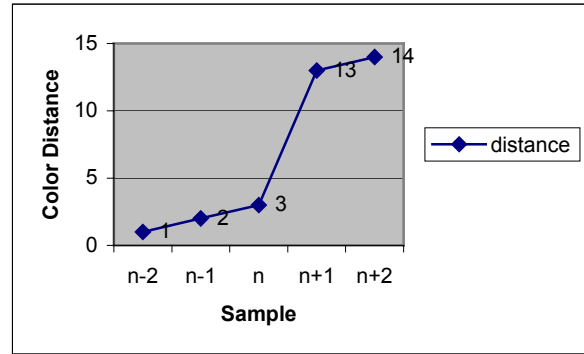


Figure 2. Demonstration of threshold selection

3.3. DBSCAN Clustering

While GDBSCAN generalizes DBSCAN to a higher level in order to be able to deal with a wider range of problems, we will carry out a specialization of GDBSCAN so that it fits our purposes. In our case, the clustering algorithm is applied to a set of regions generated as a result of splitting process and our objective is to group these regions to form larger regions that have certain amount of color homogeneity.

We define the two parameters of GDBSCAN algorithm, $NPred$ and $MinWeight$, as follows. Since each region has a non-spatial attribute, color mean, and a spatial attribute, location in the image, the binary predicate $NPred$ is defined as “color distance between two regions is smaller than Eps AND the two regions are connected” where Eps is a threshold of color distance determined from a set of representative images. For instance, given a region $R1$, a region $R2$ is in the $NPred$ -neighborhood of $R1$ iff the color distance between $R1$ and $R2$ is smaller than Eps AND $R1$ and $R2$ are connected. The function $wCard(.)$ is defined to return the total number of pixels in the $NPred$ -neighborhood. $MinCard$ is defined as a threshold of number of pixels, which is obtained experimentally. The predicate $MinWeight$ is therefore defined as “total number of pixels in the $NPred$ -neighborhood, given by $wCard(.)$, is greater than $MinCard$ pixels”.

The DBSCAN algorithm starts from an arbitrary region R . First, using the $NPred$ predicate, all connected regions that have a color distance less than Eps to R are determined. These regions form the $NPred$ -neighborhood of R . The total number of pixels in the $NPred$ -neighborhood is calculated using $wCard(.)$. If the total number of pixels is greater than $MinCard$, a new cluster with region R as the core object is created. Then, all directly density-reachable

regions that are reachable from the regions in the $NPred$ -neighborhood are added to the current cluster. The algorithm continues iteratively until no new region satisfies the predicates.

A sample lesion and the result of the initial DBSCAN clustering on this lesion are shown in Figure 3. The black regions in Figure 3b) represent the noise. They cannot be integrated into any cluster because they fail the predicate test. Notice that most of the noise is along the border of the lesion. This can be explained as follows. There is a much higher color variation in the thin region along the border when compared to the other regions. Because of this, during the region splitting, this region is divided into much smaller subregions. Since they have a high color distance to the neighboring regions, these tiny subregions are unlikely to pass the predicate test to form a new cluster or to join an existing one.

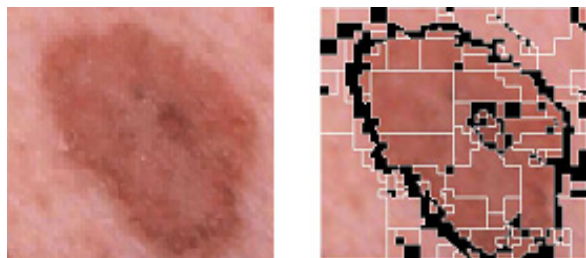


Figure 3. a) Sample image b) Before postprocessing

DBSCAN is designed to discover noise as well as clusters in a spatial data set. Also, as can be seen in Figure 3b), the number of clusters obtained from the first iteration is much more than the number of color regions that can be identified by humans. Therefore, some postprocessing is necessary.

First, the noise pixels are merged with neighboring clusters that are closest in color. Then, clusters that are closer (in color) to each other than a threshold are merged. The value of this threshold is determined by multiplying the threshold for splitting by a factor. This factor is proportional to the amount of color variation present in the initial regions. The amount of color variation in a region can be represented by the color mean absolute deviation (MAD). A region with less color variation has a smaller color MAD. In such a region, the threshold value is smaller and the merging criterion is more strict which allows the algorithm to detect subtle color variation.

3.4. More Iterations of DBSCAN

As a result of the first iteration of DBSCAN, we have identified the lesion border. Figure 4(a) shows the

image in Figure 3(a) after the first iteration. Our next goal is to go inside the lesion and identify subregions with color variegation if they exist.

If a cluster satisfies any of the following three conditions, more iterations of DBSCAN is performed: (i) the size of the cluster is larger than $1/10^{th}$ of the original image (ii) the color distance between the cluster and the healthy skin is larger than a threshold T_H (iii) the amount of color variation in the cluster is larger than a threshold T_V . The thresholds T_H and T_V are determined empirically from a set of representative images as described in the Experimental Results section.

In order to avoid over-segmentation, more than two extra iterations of DBSCAN is not allowed.

3.5. Final Cleaning

Despite the postprocessing performed after each iteration of DBSCAN, the resulting images may still have more regions than humans can perceive. Therefore, one final cleaning step is necessary. During this step, if a region is smaller than $1/100^{th}$ of the original image it is merged with the nearest cluster. Figure 4(b) shows the final segmentation result of the image in Figure 3(a).

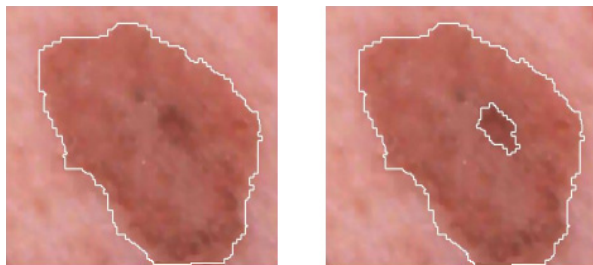


Figure 4. a) After the 1st iter. b) Final segmentation result

4. Experimental Results

In order to fine-tune the algorithm parameters and test the effectiveness of the proposed segmentation method, experiments are performed on a set of 135 clinical images with dimensions subsampled to 256×256 .

First, to fine-tune the algorithm parameters, 18 representative images are selected from the above set. After fine-tuning the parameters, the algorithm is applied to the remaining 117 images. An expert dermatologist visually examined the segmentation results. In 80% of the images the lesion borders were detected successfully. Figure 5 shows a sample of the segmentation results.

To further evaluate the effectiveness of the segmentation, an experiment is performed to compare the number of color regions identified by the algorithm and by the human subjects. Five subjects participated in the experiment. They were graduate students, ranging in age from 22 to 25. No subject had seen the segmentation results before he/she performed the experiment. All subjects had normal or corrected-to-normal vision.

The number of colors/subregions inside the lesion is used as the evaluation criterion. The kappa statistic of Cohen [2] is chosen as the measure of agreement between the segmentation results and the aggregate human observation obtained by majority vote.

The kappa coefficient equals +1 when there is a complete agreement among the observers. When the observed agreement exceeds chance agreement, kappa is positive, with its magnitude reflecting the strength of agreement. Negative values indicate that the observed agreement is less than chance agreement.

We convert the evaluation criterion (the number of colors/subregions inside the lesion) to ranks as shown in Table 1. In the experiment, no subject observed more than 3 colors/subregions in a lesion. Furthermore, only 6 out of 117 images in the segmentation results have more than 3 identified colors/subregions. Therefore, we assigned rank 3 to the cases where the lesion has 3 or more colors/subregions.

Table 1. Conversion of evaluation criterion to ranks

Number of Colors/Sub-regions	Rank
1	1
2	2
3+	3

Table 2 shows the evaluation results. The kappa coefficient, calculated from this table using SAS® System v8.0, is 0.28 (95% confidence interval), showing a significant agreement between the segmentation results and human perception.

Table 2. Evaluation results

		Human Observation			Total
		1	2	3	
Segmentation Result	1	2	3	1	6
	2	3	58	5	66
	3	2	28	15	45
Total		7	89	21	117

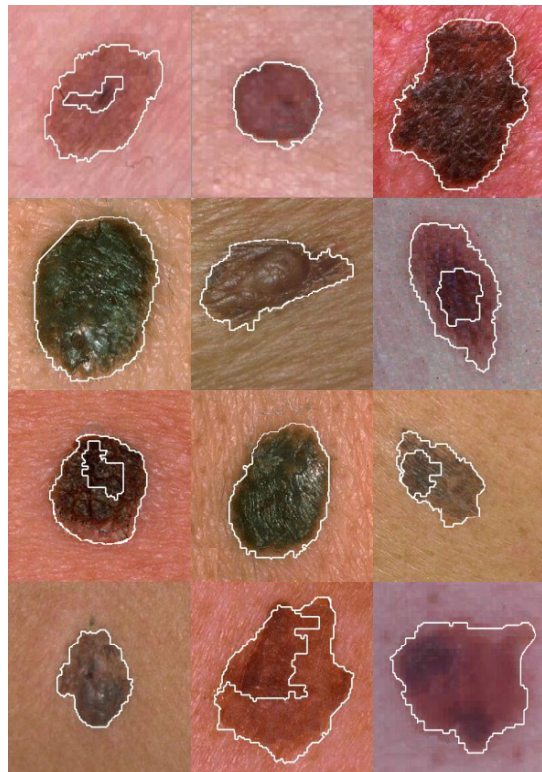


Figure 5. A sample of the segmentation results

5. Related Work

Several methods have been developed for segmentation of pigmented skin lesions, most of them being focused on border detection. In this section we review some of the recent approaches. An extensive survey of the earlier approaches can be found in [15].

Xu et al. [16] proposed a two-stage approach for unsupervised border detection. First, the color image is reduced to an intensity image which is then thresholded to obtain an approximate border. Then, the border detection is refined using edge information in the neighborhood of the initial border. Zhang et al. [17] presented an automatic scheme based on two-rounds of radial search technique originally introduced in [5]. Chung and Sapiro [1] proposed a partial-differential equations-based approach using the geodesic active contours model. Metz et al. [10] presented a semi-automatic method very similar to Xu et al.'s method that incorporates an interactive selection step to clip the region of interest. She and Fish [14] proposed a technique based on edge focusing. A fast snake is used to determine the optimum border by minimizing the energy calculated by the Laplacian of Gaussian (LoG) edge detector. Recently, Rajab et al. [11] proposed an approach based on optimal thresholding using an isodata algorithm.

Our approach differs from the previous ones in that it allows not only precise border detection but also identification of the subregions inside the lesion with variegated coloring.

6. Conclusions and Future Work

We have presented an unsupervised segmentation method based on a density-based clustering algorithm (DBSCAN) which allows separation of the healthy skin from the lesion and identification of the subregions inside the lesion with variegated coloring.

The proposed algorithm accurately detects the lesion borders in 80% of the test images. In addition, the agreement between the number of color regions identified by the algorithm and by the human subjects is found to be significant by the popular Kappa statistical agreement test.

The proposed method is not limited to the identification of color regions. By defining new distance measures for texture, for instance, the method can easily be adapted to homogenous texture region identification.

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8. References

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