Feature Based Growth Rate Analysis and Pattern Recognition of Mesenchymal Stem Cells

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Abstract - Stem cell is a generic cell that can make exact copies of it. Stem cells are master cells and can grow into different cell types in the body during evolution. Different parts of your body are made up of different kinds of cells. Stem cells are not specific to one area in the body and can turn into skin, bone, blood or brain cells. Occasionally stem cells do not build up accurately once injecting to human body, which will lead to complicated. To avoid this problem the growth of the stem cell should be monitored with the help of morphological and textural characteristics. Meanwhile, with the rapid development of feature extraction and pattern recognition techniques, it provides people with new thought on the study on complex image retrieval while it’s very difficult for traditional machine learning method to get ideal retrieval results. Pattern Recognition is one of the very important and actively searched trait or branch of artificial intelligence. It is the science which tries to make machines as intelligent as human to recognize patterns and classify them into desired categories in a simple and reliable way. For this reason, proposed a new approach for image features based on Modified Non Subsampled Contourlet Transform (MNSCT), Radial Basis Function Kernel Principal Component Analysis (RBFKPCA), and Relevance Vector Machine (RVM). The proposed method is used to extract the salient features of stem cell images by using efficiency of Modified Non Subsampled Contourlet Transform (MNSCT) that finds the best feature points. Relevance Vector Machine (RVM) is a probabilistic model whose functional form is equivalent to the SVM. It achieves comparable recognition accuracy to the SVM, yet provides a full predictive distribution, and also requires substantially fewer kernel functions.

Keywords: Mesenchymal stem cells - Non Subsampled Contourlet Transform (NSCT) - Modified Non Subsampled Contourlet Transform (MNSCT) - Radial Basis Function Kernel Principal Component Analysis (RBFKPCA) - Support Vector Machine (SVM) - Relevance Vector Machine (RVM).

I. INTRODUCTION

Stem cells are the raw material from which all of the body’s mature differentiated cells are made. Stem cells give rise to brain cells, nerve cells, heart cells, pancreatic cells. The stem cell images are analyzed with the help of morphological and texture feature. Many features can be extracted from the image are used for the analysis. Non Sampled Contourlet Transform (NSCT) [1] eliminates the noise in redundant images. The contour let transform is a multidirectional and multiscale transform that is constructed by combining the Laplacian pyramid [2] with the directional filter bank [3] proposed. The pyramidal filter bank structure of the contour let transform has very little redundancy, which is important for compression applications. Various classification methodologies have been reported such as, rule-based systems [4] and [5], fuzzy logic systems [6], statistical methods based on Markov random fields and support vector machines [7]. Global geometric moments and their invariants are widely used in many areas of image analysis, including pattern recognition [8], image reconstruction [9], and fingerprint Recognition [10]. Principal component analysis (PCA) is deployed in JPEG2000 to provide spectral de-correlation as well as spectral dimensionality reduction [11]. Feature extraction
of hyper-spectral remote sensing data can be investigated using Principal component analysis (PCA) which has shown to be a good unsupervised feature extraction. On the other hand, this method only focuses on second orders statistics. By mapping the data onto another feature space and using nonlinear function, Kernel PCA (KPCA) can extract higher order statistics [12]. In recent years, SVM learning has found a wide range of real-world applications, including object recognition [13], speaker identification [14] and face detection in images [15]. RVM, developed by Tipping [16] is based on Bayesian estimation for classification problems. It is proved to be faster than SVM since it yields an optimum solution with fewer training samples. Relevance vector machine [17], [18] (RVM)-based regression and classification has been proposed by Tipping. To compare the performance of RVM and SVM classifications with different number of features, the steepest ascent [19], [20] algorithm is used to reduce the number of features. The proposed method is explained in section 2, Section 3 shows the Results and Discussions of this method, and finally concludes this remarks in section 4.

II. MATERIALS AND METHODS

Pattern recognition is used for the identification of the healthier stem cells used for analyzing the growth rate. There are various methods for finding Pattern Recognition, Relavance vector machine (RVM) is the most significant and efficient method used for Classification. In order to find the growth rate of Stems cells accurately, sparse property is implemented in RVM. Growth rate is calculated based on some texture features. The features required for the calculation of the growth rate are extracted using modified NSCT with Radial basis function of KPCA. In modified NSCT the various filters present in the filter bank are combined as a single filter, hence the filter used modified NSCT has the efficiency of all the filters in the filter bank. The features of mesenchymal stem cells are extracted by using modified NSCT and the patterns are recognized by RVM and the overall block diagram shown in given below

![Overall Architecture](image)

2.1 NON SAMPLED CONTOURLET TRANSFORM (NSCT):

Nonsubsampled Contour Transformation (NSCT) [1] is a kind of algorithm which uses the iterative non subsampled filter bank to achieve a series of multi-resolution [2], multi-direction and translation invariant frequency domain sub-images. In this algorithm, multi-scale analysis is conducted primarily on the images, then multi-direction analysis is conducted, and the layer by layer iteration is conducted separately for these two steps. The structure of NSCT is divided into nonsubsampled pyramid filter bank (NSPFB) and nonsubsampled directional filter bank (NSDFB), and its realization structural principle is as shown in Fig. 2. Its decomposition process can be attain by the nonsubsampled pyramid filter bank (NSPFB) and it is used to conduct multi-scale decomposition on the images, to obtain various high-frequency sub-band images with different frequencies and a low-frequency sub-band image, and then the nonsubsampled directional filter bank is used to conduct multi-direction decomposition for the obtained various high-frequency sub-band images, so as to obtain the sub-band images (coefficients) of different scales and different directions.
When the decomposition level of NSPFB is the low pass sub-band image and L high pass sub-band images can be obtained through decomposition. Because there is no down-sampling, all the sub-band images have the same sizes with the original images. The realization process of NSDFB is up-sampling can be conducted on the filter, and secondly the wave filtering is conducted on the decomposed sub-band images in the two channel direction of last stage, so as to achieve the more accurate directional decomposition in the frequency domain. 2 t directional sub-band images which have the same size as the original images can be obtained through 1 stage directional decomposition of NSDFB on a scale sub-band image. NSPFB is combined with NSDFB to achieve NSCT. The band-pass sub-band images, which are obtained after decomposing the original images through NSPFB, are NSDFB and the band-pass directional sub-band images can be obtained, so as to achieve the multi-scale and multi-direction decomposition of images.

2.1.1 NSCT ALGORITHM:
1. Compute the NSCT of the input image for N levels.
2. Estimate the noise standard deviation of the input image.
3. For each level DFB,
   (a) Estimate the noise variance.
   (b) Compute the threshold and the amplifying ratio.
   (c) At each pixel, compute the mean and the maximum magnitude of all directional subbands at this level, and classify it by (1) into strong edges, weak edges, or noises.
   (d) For each directional subband, use the nonlinear mapping function given in (2) to modify the NSCT coefficients according to the classification.
4. Reconstruct the enhanced image from the modified NSCT coefficients.

2.2 MODIFIED NON SUBSAMPLED CONTOURLETTRANSFORMATION (NSCT):
In former NSCT method, the directional filter bank contains one or more filters but the particular filter is used according to the input images which are undergone to decomposition. And also aliasing effect is present in the directional filter. To increase the efficiency of the decomposition filter, the filters present in the directional filter banks are combined to form a single filter to utilize the advantage of all the filters and also more filters are used in the modified NSCT. This will increase the decomposition efficiency by make use of all the filters.

Filter in MNSCT:
1. "Haar" Filters
2. McClellan Transformed Of The Filter
3. Ladder Filters
4. Ideal Filter
5. Diamond Maxflat Filters Obtained from a Three Stage Ladder filter
6. Orthogonal Filters

7. Daubechies Filter

8. Regular Linear Phase Biorthogonal Filter

When the above filter banks are used in the modified NSCT (Non Sampled Contourlet Transform). These filter banks are used to increase the decomposition and efficient values compare to NSCT. When the above filters are used and these filters are working in the series condition to decompose the mesenchymal stem cells.

2.3 PRINCIPAL COMPONENT ANALYSIS (PCA):

Principal Component Analysis (PCA) [11] is a statistical technique used to reduce a set of correlated multivariate measurements to a smaller set where the features are uncorrelated to each other and also it is a powerful technique for extracting structure from possibly high dimensional data sets. The goal of PCA is to reduce the dimensionality of the data while retaining as much as possible of the variation present in the dataset. Each dimensionality reduction technique finds an appropriate transformation by satisfying certain criteria.

2.3.1 PCA ALGORITHM:

1. Subtract the mean from all the data points
2. Compute the covariance matrix \( S = \sum_{n=1}^{N} x_n x_n^T \)
3. Diagonalize \( S \) to get its eigen values and eigenvectors
4. Retain \( c \) eigenvectors corresponding to the \( c \) largest eigenvalues such that \( \sum_{j=1}^{c} \lambda_j / \sum_{j=1}^{N} \lambda_j \) equals the desired variance to be captured.
5. Project the data points on the eigenvectors \( a_n = E^T \left( x_k - \frac{1}{N} \sum_{i=1}^{N} x_i \right) \)

2.4 KERNEL PRINCIPAL COMPONENT ANALYSIS (KPCA):

Kernel principal component analysis (kernel PCA) [12] is an extension of principal component analysis (PCA) using techniques of kernel methods. Using a kernel, the originally linear operations of PCA are done in a reproducing kernel Hilbert space with a non-linear mapping. Kernel based principle components analysis is a non linear PCA created using the kernel trick. KPCA maps the original inputs into a high dimensional feature space using a kernel method. To transform the current features into a high-dimensional space and the calculate eigenvectors in this space. To ignore the vectors with really low eigen values and then do learning in this transformed space. KPCA is computationally intensive and takes a lot more time compared to PCA. The reason being that the number of training data points in PCA is much higher then KPCA. There are many types of kernels like Gaussian, Polynomial and the Sigmoid, comparing to the other types of kernels, they had better advantages in the Radial Basis Function of Kernel Principal Component Analysis.

2.5 RADIAL BASIS FUNCTION KPCA (RBFKPCA):

Radial basis function is a kind of supervised neural networks. It has a design of NN as curve-fitting problem and also learning the find surface in multidimensional space best fit to training data. Generally it has been used the multidimensional surface to interpolate the test data and find the approximate function with linear combination of Radial basis functions a \( F(x) = \sum w_i h(x) \),Where \( h(x) \) is mostly Gaussian function
When the above architecture has three types of layers: Input layer, hidden layer, Output layer. The input layer has denoted by f(x), the source nodes that connect to the network to its environment and the second layer is the hidden layer, these layer provides hidden units provide a set of basis function and it has a high dimensionality layers and final layer is the Output layer, these layer has functions of x1 , x2 , x3 , x4 …xn has linear combination of hidden functions.

2.6 SVM

Support Vectors Machines (SVM) [14] has recently shown their ability in pattern recognition and classification [15]. The potential of the SVM is illustrated on a 3D object recognition task using the Coil database and on an image classification task using the Corel database. The images are either represented by a matrix of their pixel values (bitmap representation) or by a color histogram. Support vectors (SVs) contain highlighted pixels that help to create the margins or boundaries in an image. SVM is a mostly used method in pattern recognition and object recognition initially, given a set of points which belong to either of two classes; a linear SVM finds the hyper plane leaving the largest possible fraction of points of the same class on the same side, while maximizing the distance of either class from the hyper plane.

Support Vector Machines (SVM) as a classification method. It has been provide better classification results that other widely used pattern methods, such as the maximum likelihood and neural network classifiers. SVM are very attractive for the classification of remotely sensed data.

The SVM approach seeks to find the optimal separating hyper plane between classes by focusing on the training cases that are placed at the edge of the class descriptors. These training cases are called support vectors. Training cases other than support vectors are discarded. This way, not only is an optimal hyper plane fitted, but also less training samples are effectively used; thus high classification accuracy is achieved with small training sets. This feature is very advantageous, especially for remote sensing datasets and more specifically for Object-based Image Analysis, where object samples tend to be less in number than in pixel-based approaches.

An SVM classifies data by finding the best hyper plane that separates all data points of one class from those of the other class. The best hyper plane for an SVM means the one with the largest margin between the two classes. Margin means the maximal width of the slab parallel to the hyper plane that has no interior data points. The support vectors are the data points that are closest to the separating hyper plane; these points are on the boundary of the slab.
2.6.1 SVM PROBLEMS:

When the SVM had number of required support vectors typically grows linearly with the size of the training set and also it has Non probabilistic predictions. It requires estimation of error and margin trade off parameters.

2.7 RVM:

Relevance Vector Machine (RVM) [17] as a probabilistic sparse kernel model based on the support vector machine theory. Each of the model's weights has associated a prior that is characterized by a set of hyper parameters whose values are determined during the learning process. Relevance Vector Machine (RVM) makes the probabilistic predictions and yet that keeps the excellent predictive performance of the support vector machine. It also can make the preservation of the sparseness property of the SVM. In fact, for a wide variety of test problems it actually results in models which are dramatically sparser than the corresponding SVM, while sacrificing little if anything in the accuracy of prediction. RVM is used for detecting the cancer cells and the classification of DNA sequences. It utilizes fewer kernel functions than SVM.

Relevance vector machine (RVM) is a special case of a sparse linear model, where the basis functions are formed by a kernel function $\phi$ centered at the different training points While this model is similar in form to the support vector machines (SVM), the kernel function here does not need to satisfy the Mercer’s condition, which requires $\phi$ to be a continuous symmetric kernel of a positive integral operator.

Multi-kernel RVM is an extension of the simple RVM model. It consists of several different types of kernels. The sparseness property enables automatic selection of the proper kernel at each location by pruning all irrelevant kernels, though it is possible that two different kernels remain on the same location.

III. RESULTS AND ANALYSIS

3.1 FEATURE EXTRACTION RESULTS

There are two types of methods to extract the features. The initial one is NSCT (Non Sampled Contour let Transform), PCA (Principal Component Analysis), SVM (Support Vector Machine) and the other methods of MNSCT (Modified Non Sampled Contour let Transform), KPCA (Kernel Principal Component Analysis), and RVM (Relevance Vector Machine). Features like energy, entropy, auto correlation, contrast, homogeneity and dissimilarity of the images are extracted. In the proposed method, using the MNSCT and RVM results are compared to the NSCT and SVM for the best results. In MNSCT, the original image is decomposed by using the matlab function, in this function there are many filter banks is used to decompose the segmented image to a fine detailed image so that the features can be extracted more accurately. After the decomposition process the decomposed image is reconstructed by using the function “mnsctrec”. The function mnsctrec will be doing the opposite process of the mnsctdec and the reconstruction should be perfect until the process is carried out.
The eight frames of images can be reclaiming from the 72-hour video lapse of distinctive hours subsequently. When the above Mesenchymal Stem Cells are progenitor connective tissues cells. The height and width of stem cells is 720 pixels. When the above images can be calculated from the different hours of growing the stem cells to analyze the growth rate.

Fig: 4 Input Images Mesenchymal Stem Cells

<table>
<thead>
<tr>
<th>Input</th>
<th>Hours</th>
<th>Autocorrelation</th>
<th>Contrast</th>
<th>Dissimilarity</th>
<th>Energy</th>
<th>Homogenity</th>
<th>Entropy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Image 1</td>
<td>8 hours</td>
<td>45.9456</td>
<td>4.6422</td>
<td>0.9417</td>
<td>0.5787</td>
<td>0.8215</td>
<td>3.6407</td>
</tr>
<tr>
<td>Image 2</td>
<td>16 hours</td>
<td>42.5495</td>
<td>4.6406</td>
<td>1.0063</td>
<td>0.5787</td>
<td>0.8134</td>
<td>3.7506</td>
</tr>
<tr>
<td>Image 3</td>
<td>24 hours</td>
<td>38.5282</td>
<td>5.7792</td>
<td>1.1490</td>
<td>0.4938</td>
<td>0.7956</td>
<td>4.8811</td>
</tr>
<tr>
<td>Image 4</td>
<td>32 hours</td>
<td>32.3821</td>
<td>6.4195</td>
<td>1.3116</td>
<td>0.4128</td>
<td>0.7752</td>
<td>4.0047</td>
</tr>
<tr>
<td>Image 5</td>
<td>40 hours</td>
<td>29.8924</td>
<td>7.3105</td>
<td>1.4031</td>
<td>0.3901</td>
<td>0.7638</td>
<td>4.0463</td>
</tr>
<tr>
<td>Image 6</td>
<td>48 hours</td>
<td>24.0953</td>
<td>8.3983</td>
<td>1.5304</td>
<td>0.3773</td>
<td>0.7479</td>
<td>4.0811</td>
</tr>
<tr>
<td>Image 7</td>
<td>56 hours</td>
<td>20.1897</td>
<td>8.3983</td>
<td>1.5429</td>
<td>0.3968</td>
<td>0.7463</td>
<td>4.0600</td>
</tr>
<tr>
<td>Image 8</td>
<td>64 hours</td>
<td>17.4842</td>
<td>8.1784</td>
<td>1.5115</td>
<td>0.4224</td>
<td>0.7503</td>
<td>4.0252</td>
</tr>
</tbody>
</table>

Table 1: Tabulation of Non Sampled Contourlet Transform

<table>
<thead>
<tr>
<th>Input</th>
<th>Hours</th>
<th>Autocorrelation</th>
<th>Contrast</th>
<th>Dissimilarity</th>
<th>Energy</th>
<th>Homogenity</th>
<th>Entropy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Image 1</td>
<td>8 hours</td>
<td>47.6566</td>
<td>4.3021</td>
<td>1.0289</td>
<td>1.1699</td>
<td>1.3589</td>
<td>3.8350</td>
</tr>
<tr>
<td>Image 2</td>
<td>16 hours</td>
<td>44.2605</td>
<td>4.7547</td>
<td>1.0935</td>
<td>1.0929</td>
<td>1.3508</td>
<td>3.9451</td>
</tr>
<tr>
<td>Image 3</td>
<td>24 hours</td>
<td>40.2392</td>
<td>5.7531</td>
<td>1.2362</td>
<td>1.0080</td>
<td>1.3330</td>
<td>4.0754</td>
</tr>
<tr>
<td>Image 4</td>
<td>32 hours</td>
<td>34.0931</td>
<td>6.8917</td>
<td>1.3988</td>
<td>0.9270</td>
<td>1.3126</td>
<td>4.1990</td>
</tr>
<tr>
<td>Image 5</td>
<td>40 hours</td>
<td>31.6034</td>
<td>7.5320</td>
<td>1.4903</td>
<td>0.9043</td>
<td>1.3012</td>
<td>4.2406</td>
</tr>
<tr>
<td>Image 6</td>
<td>48 hours</td>
<td>25.8063</td>
<td>8.4230</td>
<td>1.6176</td>
<td>0.8915</td>
<td>1.2853</td>
<td>4.2754</td>
</tr>
<tr>
<td>Image 7</td>
<td>56 hours</td>
<td>21.9007</td>
<td>8.5108</td>
<td>1.6301</td>
<td>0.9110</td>
<td>1.2837</td>
<td>4.2543</td>
</tr>
<tr>
<td>Image 8</td>
<td>64 hours</td>
<td>19.1952</td>
<td>8.2909</td>
<td>1.5987</td>
<td>0.9366</td>
<td>1.2877</td>
<td>4.2195</td>
</tr>
</tbody>
</table>

Table 2: Tabulation of Modified Non Sampled Contourlet Transform

When the above table 1 NSCT (Non Sampled Contourlet Transform) and the table MNSCT (Modified Non Subsampled Contourlet Transform) has eight input images can be calculated for 64 hours of images and find the
features like autocorrelation, contrast, dissimilarity, energy, entropy and homogeneity values should be decreased in the NSCT compare to the MNSCT values.

3.2 PATTERN RECOGNITION RESULTS

The above graph is denoted by using NSCT+PCA+SVM. SVM (Support Vector Machine) is a binary classifier. It is used to separates the classified data and it is a non-probabilistic decision machine. SVM has been growing linearly with the size of training set, they had estimate the error is minimized and maximize the margin of the lines. Compare to the MNSCT+KPCA+RVM, the above graph performance is poor because SVM had some of the limitation has cross validation of parameters is $c$ and $\sum$ and the Decisions at outputs instead of probabilities then they had a large number of kernels (in spite of sparsity) and also the kernels must satisfy Mercer criterion.

Table 3: comparison of SVM/RVM classifier

<table>
<thead>
<tr>
<th>Input</th>
<th>Hours</th>
<th>SVM Classifier</th>
<th>RVM Classifier</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Efficiency</td>
<td>Growth rate</td>
</tr>
<tr>
<td>Image 1</td>
<td>8 hours</td>
<td>0.5932</td>
<td>24046</td>
</tr>
<tr>
<td>Image 2</td>
<td>16 hours</td>
<td>0.5256</td>
<td>37015</td>
</tr>
<tr>
<td>Image 3</td>
<td>24 hours</td>
<td>0.6202</td>
<td>60061</td>
</tr>
<tr>
<td>Image 4</td>
<td>32 hours</td>
<td>0.5932</td>
<td>88515</td>
</tr>
<tr>
<td>Image 5</td>
<td>40 hours</td>
<td>0.5797</td>
<td>111830</td>
</tr>
<tr>
<td>Image 6</td>
<td>48 hours</td>
<td>0.5797</td>
<td>150892</td>
</tr>
<tr>
<td>Image 7</td>
<td>56 hours</td>
<td>0.6202</td>
<td>165432</td>
</tr>
<tr>
<td>Image 8</td>
<td>64 hours</td>
<td>0.5527</td>
<td>174841</td>
</tr>
</tbody>
</table>

Fig: 6 NSCT+PCA+SVM Graphs

Fig: 7 MNSCT+KPCA+RVM Graphs
The above table 3 denote the SVM and RVM classifiers. Compare to the SVM, RVM is a Bayesian sparse kernel technique for regression and classification. When the efficiency and growth rate of the SVM classifier values are decreased compare to the RVM classifier because they had some of the peculiar properties of RVM had the error rates compare to the SVM is less. They had no cross validation of parameters. It is one of the applicable to wide choice of basis function multi-class classification and the Probabilistic outputs dramatically fewer kernels (by an order of magnitude) but, slower to train than SVM.

3.3 OVERALL PERFORMANCE

When the above graph denotes the overall performance of graphs. The x-axis denotes the images and the y-axis denotes the growth rate. The red color denotes the NSCT, PCA, SVM performance and the blue color denotes the MNSCT, KPCA, RVM performance. The performance of modified NSCT, KPCA, and RVM is better performance compare to the SVM performance.

IV. CONCLUSIONS

In this paper, introduced an efficient feature-based extraction for mesenchymal stem cell images, which is employing with MNSCT + KPCA + RVM, transform. As a result, the proposed method gives promising results compare to the NSCT+PCA+SVM. According to the experiments, we can conclude that the method proposed is maintaining good performance; The different classifier models analyzed in this report show that they are very promising to achieve higher efficiency and speed in lesser time results. Also the KPCA is used for dimensionality reduction. The RVM classifier performs well and provides high recognition rate compared to SVM. In future using the multikernel RVM for getting the better accuracy of the images.

REFERENCES


