Preliminary Level Automated Classification of Brain Tumor Using PCA and PNN

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Abstract: Probabilistic neural networks (PNN) are finding many uses in the medical diagnosis application. The goal of this paper is to use Probabilistic Neural Network (PNN) with one important mathematical technique ‘Principal Component Analysis’ (PCA) for preliminary level brain tumor classification. As a result, the tested Magnetic Resonance Image (MRI) of brain is classified either benign or malignant. Automated classifications of brain tumors is performed in two stages. Feature extraction using Principal Component Analysis (PCA) and classification using Probabilistic Neural Network (PNN)

Keywords: Probabilistic Neural Network, Clustering, classification, Segmentation.

I. INTRODUCTION
PNN provide a powerful tool to help doctors to analyze, model and make sense of complex clinical data across a broad range of medical applications. Most of the applications provide the solution to the classification problem. Feature selection plays an important role in classifying systems such as PNN. The higher performance with lower computational effort is expected with this process. One of the most popular methods for dealing with this problem is the PCA method. This method transforms the existing attributes into new ones considered to be crucial. Used feature extraction methods in automated diagnosis of arterial diseases. Since classification is more accurate when the pattern is simplified through representation by important features, feature extraction and selection play an important role in classifying systems. Used image segmentation based on the soft computing for improved implementation of the brain tumor detection. The MRI brain image is acquired from patient’s database and then Image acquisition, preprocessing, image segmentation is performed for brain tumor detection. Also did the work for improving brain tumor characterization on MRI by probabilistic neural network and non-linear transformation of textural features. The PNN architecture can be used for high speed classification of remotely sensed imagery and can be applied to remotely sensed data.

II. METHODOLOGY

- Training phase
- Testing phase
- Image conversion
- Development phase

Training phase:- Mathematical technique ‘PCA’ is used to extract feature vectors of all images in the database. Back propagation algorithm is used to train the neural network.

Testing phase:- Feature vector of the test image will be computed in this phase. Euclidean Distance is also calculated to decide in which class of ‘brain tumor’ the input image is to be fitted.

Image Conversion:- MR images will be converted into matrices form using Feed Forward PNN will be used to classify MRI images. Already available function for PNN is used as it in MATLAB.

Development phase:- Performance analysis based on the result will be carried out in the development phase.

III. EXISTING SYSTEM

High-order measures of independence for the independent component analysis problem and discusses the class of Jacobi algorithms for their optimization. Several implementations are discussed. We compare the proposed approaches with gradient-based techniques from the algorithmic point of view and also on a set of biomedical data Many recent contributions to the ICA problem in the neural network literature describe stochastic gradient algorithms involving as an
essential device in their learning rule a nonlinear activation function. Other ideas for ICA, most of them found in the signal processing literature, exploit the algebraic structure of high-order mends of the observations. They are often regarded as being unreliable, inaccurate, slowly convergent, and utterly sensitive to outliers. As a matter of fact, it is fairly easy to devise an ICA method displaying all these flaws and working on only carefully generated synthetic data sets. This may be the reason that cumulant-based algebraic methods are largely ignored by the researchers of the neural network community involved in ICA. This article tries to correct this view by showing how high-order correlations can be efficiently exploited to reveal independent components. This article describes several ICA algorithms that may be called Jacobi algorithms because they seek to maximize measures of independence by a technique akin to the Jacobi method of diagonalization. These measures of independence are based on fourth-order correlations between the entries of Y. As a benefit, these algorithms evades the curse of gradient descent:

Advantages and Drawbacks for PNN

- It is usually much faster to train a PNN network than a multilayer preceptor network.
- PNN networks generally are more accurate than multilayer preceptor networks.
- PNN networks are relatively insensitive to outliers (wild points).
- PNN networks generate accurate predicted target probability scores.
- PNN networks approach Bayer’s optimal classification.
- PNN networks are slower than multilayer preceptor networks at classifying new cases.
- PNN networks require more memory space to store the model.

IV. PROPOSED SYSTEM

MRI Brain Image Classification and Tumor Detection Is Proposed Based On,
- Probabilistic Neural Network for classification.
- Clustering Algorithm (K means) for effective Segmentation.

A. Segmentation

Segmentation is the process of partitioning a digital image into multiple segments (sets of pixels, also known as super pixels)[6]. The goal of segmentation is to simplify and/or change the representation of an image into something that is more meaningful and easier to analyze. Image segmentation is typically used to locate objects and boundaries (lines, curves, etc.) in images. More precisely, image segmentation is the process of assigning a label to every pixel in an image such that pixels with the same label share certain visual characteristics.

The result of image segmentation is a set of segments that collectively cover the entire image, or a set of contours extracted from the image (see edge detection). Each of the pixels in a region is similar with respect to some characteristic or computed property, such as color, intensity, or texture. Adjacent regions are significantly different with respect to the same characteristic(s) when applied to a stack of images, typical in medical imaging, the resulting contours after image segmentation can be used to create 3D reconstructions with the help of interpolation algorithms like Marching cubes[6].

B. Clustering

Clustering can be considered the most important unsupervised learning problem, so, it deals with finding a structure in a collection of unlabeled data. A cluster is therefore a collection of objects which are “similar” between them and are “dissimilar” to the objects belonging to other clusters [10].

Clustering algorithms may be classified as listed below

- Exclusive Clustering
- Overlapping Clustering
- Hierarchical Clustering
- Probabilistic Clustering

In the first case data are grouped in an exclusive way, so that if a certain datum belongs to a definite cluster then it could not be included in another cluster. On the contrary the second type, the overlapping clustering, uses fuzzy sets to cluster data, so that each point may belong to two or more clusters with different degrees of membership. In this case, data will be associated to an appropriate membership value. A hierarchical clustering algorithm is based on the union between the two nearest clusters [10]. The beginning condition is realized by setting every datum as a cluster. After a few iterations it reaches the final clusters wanted.

C. K-Means Clustering

Cluster analysis, an important technology in data mining, is an effective method of analyzing and discovering useful information from numerous data. Cluster algorithm groups the data into classes or clusters so that objects within a cluster have high similarity in comparison to one another, but are very dissimilar to objects in other clusters [10]. Dissimilarities are assessed based on the attribute values describing the objects. Often, distance measures are used. As a branch of statistics and an example of unsupervised learning, clustering provides us an exact and subtle analysis tool from the mathematic view K-means algorithm belongs to a popular partition method in cluster analysis. The most
widely used clustering error criterion is squared-error criterion, it can be defined as

\[ J_c = \sum_{j=1}^{c} \sum_{k=1}^{n} \left\| x^{(j)}_k - m_j \right\|^2 \]

where \( J_c \) is the sum of square-error for all objects in the database, \( x_k \) is the point in space representing a given object, and \( m_j \) is the mean of cluster \( C_j \). Adopting the squared-error criterion, K-means works well when the clusters are compact clouds that are rather well separated from one another and are not suitable for discovering clusters with no convex shapes or clusters of very different size. For attempting to minimize the square-emir criterion, it will divide the objects in one cluster into two or more clusters. Aiming at the dependency to initial conditions and the limitation of K-means algorithm that applies the square-error criterion to measure the quality of clustering, this paper presents a new improved K-means algorithm that is based on effective techniques of multi-sampling and once-clustering to search the optimal initial values of cluster centers. Our experimental results demonstrate the new algorithm can obtain better stability and excel the original K-means in clustering results.

### D. Pseudo Code for K-Means

In this section, we briefly describe the original K-means algorithm.

Original K-means(s,k), s={\( x_1, x_2, \ldots, x_n \)}.

Input: the number of clusters \( K \) and a dataset containing \( n \) objects \( (x_i) \).

Output: a set of \( k \) clusters \( c_j \) that minimize the squared-error criterion

**Begin**

\( m=1 \);

Initialize \( k \) prototypes \( Z_j \), \( j \in [1,K] \);

repeat

  for \( i=1 \) to \( n \) do

    Begin

    for \( j=1 \) to \( k \) do

        compute \( D(X_i,Z_j) = |X_i - Z_j| \);

        if \( D(X_i,Z_j) \) min \{ \( D(X_i,Z_j) \) \} then \( X_i \in C_j \);

    end;

    if \( m=1 \) then \( J_c(m) = \sum_{j=1}^{k} \sum_{x \in C_j} |X_i - Z_j|^2 ; \)

    \( m=m+1 \)

    For \( j=1 \) to \( k \) do

      \( Z_j = 1/n_j \sum_{j=1}^{n_j} x^{(j)}_i \sum_{x \in j} x^{(i)}_i \)

      \( J_c(m) = \sum_{j=1}^{k} \sum_{x \in C_j} |X_i - Z_j|^2 ; \)

  Until \( J_c(m) - J_c(m-1) < \epsilon \)

**End**

The computational complexity of original K-means algorithm is \( O(ndk) \), where \( n \) is the total number of objects, \( k \) is the number of clusters, and \( d \) is the dimensions of datasets.

### E. Algorithm Flow diagram

![Algorithm Flow diagram](image-url)
V. SYSTEM ARCHITECTURE

As the complexity of systems increases, the specification of the system decomposition is critical. Moreover, subsystem decomposition is constantly revised whenever new functionality, new issues are addressed. Subsystems are merged into alone subsystem, a complex subsystem is split into parts, and some subsystems are added to take

A. Modules description

- GLCM Feature Extraction.
- PNN Training and Classification.
- Clustering Method for Tumor Detection.

Texture Analysis

Texture is that innate property of all surfaces that describes visual patterns, each having properties of homogeneity. It contains important information about the structural arrangement of the surface, such as; clouds, leaves, bricks, fabric, etc. It also describes the relationship of the surface to the surrounding environment. In short, it is a feature that describes the distinctive physical composition of a surface.

Texture properties include:
- Coarseness
- Contrast
- Directionality
- Line-likeness
- Regularity
- Roughness

Texture is one of the most important defining features of an image. It is characterized by the spatial distribution of gray levels in a neighborhood [8]. In order to capture the spatial dependence of gray-level values, which contribute to the perception of texture, a two-dimensional dependence texture analysis matrix is taken into consideration. This two-dimensional matrix is obtained by decoding the image file; jpeg, bmp, etc.

B. Methods of Representation

There are three principal approaches used to describe texture; statistical, structural and spectral.

- Statistical techniques characterize textures using the statistical properties of the grey levels of the points/pixels comprising a surface image. Typically, these properties are computed using: the grey level co-occurrence matrix of the surface, or the wavelet transformation of the surface.
- Structural techniques characterize textures as being composed of simple primitive structures called “Texel’s” (or texture elements). These are arranged regularly on a surface according to some surface arrangement rules.
- Spectral techniques are based on properties of the Fourier spectrum and describe global periodicity of the grey levels of a surface by identifying high-energy peaks in the Fourier spectrum.

For optimum classification purposes, what concern us are the statistical techniques of characterization.[1] This is because it is these techniques that result in computing texture properties. The most popular statistical representations of texture are:
- Co-occurrence Matrix
- Tamura Texture
- Wavelet Transform
C. Co-occurrence Matrix

Originally proposed by R.M. Haralick, the co-occurrence matrix representation of texture features explores the grey level spatial dependence of texture [2]. A mathematical definition of the co-occurrence matrix is as follows [4]:

- Given a position operator \( P(i,j) \),
- let \( A \) be an \( n \times n \) matrix
- Whose element \( A[i][j] \) is the number of times that points with grey level (intensity) \( g[i] \) occur, in the position specified by \( P \), relative to points with grey level \( g[j] \).
- Let \( C \) be a \( n \times n \) matrix that is produced by dividing \( A \) with the total number of point pairs that satisfy \( P \). \( C[i][j] \) is a measure of the joint probability that a pair of points satisfying \( P \) will have values \( g[i], g[j] \).
- \( C \) is called a co-occurrence matrix defined by \( P \).

Examples for the operator \( P \) are: “\( i \) above \( j \)”, or “\( i \) one position to the right and two below \( j \)”, etc. This can also be illustrated as follows… Let \( t \) be a translation, then a co-occurrence matrix \( C_t \) of a region is defined for every grey-level \((a, b)\) by [1]:

\[
C_t(a, b) = \text{card}\{ (s, s+t) \in R^2 | A[s] = a, A[s+t] = b \}
\]

Here, \( C_t(a, b) \) is the number of site-couples, denoted by \((s, s + t)\) that are separated by a translation vector \( t \), with \( a \) being the grey-level of \( s \), and \( b \) being the grey-level of \( s + t \). For example; with an 8 grey-level image representation and a vector \( t \) that considers only one neighbor, we would find [1]

\[
\begin{array}{cccccccc}
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
1 & 0 & 1 & 2 & 0 & 0 & 0 & 0 \\
2 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\
3 & 0 & 0 & 1 & 1 & 0 & 0 & 0 \\
4 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\
5 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
6 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
7 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
\end{array}
\]

Figure: Classical Co-occurrence matrix

At first the co-occurrence matrix is constructed, based on the orientation and distance between image pixels. Then meaningful statistics are extracted from the matrix as the texture representation. Haralick proposed the following texture features:

1. Energy
2. Contrast
3. Correlation
4. Homogeneity
5. Entropy

Hence, for each Haralick texture feature, we obtain a co-occurrence matrix. These co-occurrence matrices represent the spatial distribution and the dependence of the grey levels within a local area. Each \((i,j)\) entry in the matrices, represents the probability of going from one pixel with a grey level of \( i \) to another with a grey level of \( j \) under a predefined distance and angle. From these matrices, sets of statistical measures are computed, called feature vectors.

**Energy:** It is a gray-scale image texture measure of homogeneity changing, reflecting the distribution of image gray-scale uniformity of weight and texture.

\[
E = \sum_{x} \sum_{y} p(x,y)^3
\]

\( p(x,y) \) is the GLCM

**Contrast:** Contrast is the main diagonal near the moment of inertia, which measure the value of the matrix is distributed and images of local changes in number, reflecting the image clarity and texture of shadow depth.

\[
I = \sum_{x} \sum_{y} (x-y)^2p(x,y)
\]

**Entropy:** It measures image texture randomness, when the space co-occurrence matrix for all values is equal, it achieved the minimum value.

\[
S = -\sum_{x} \sum_{y} p(x,y) \log p(x,y)
\]

**Correlation Coefficient:** Measures the joint probability occurrence of the specified pixel pairs.

**Correlation:**

\[
\frac{\text{sum(\text{sum}(x-\mu_x)(y-\mu_y)p(x,y)/\sigma_x/\sigma_y))}}{
\text{sum(\text{sum}(x-\mu_y)p(x,y)/\sigma_y))}}
\]

**Homogeneity:** Measures the closeness of the distribution of elements in the GLCM to the GLCM diagonal.
Homogeneity = \text{sum(sum}(p(x, y)/(1 + |x-y|))\text{)}

\textbf{D. Discrete Wavelet Transform (DWT)}

In numerical analysis and functional analysis, a discrete wavelet transform (DWT) is any wavelet transform for which the wavelets are discretely sampled. As with other wavelet transforms, a key advantage it has over Fourier transforms is temporal resolution: it captures both frequency and location information (location in time).

\textbf{Types of DWT}

There are two types of DWT. They are

- One dimensional DWT (1D DWT)
- Two Dimensional DWT (2D DWT)

\textbf{One Dimensional DWT (1-D)}

The DWT of a signal \( x \) is calculated by passing it through a series of filters. First the samples are passed through a low pass filter with impulse \( g \) resulting in a convolution of the two:

\[
y[n] = (x * g)[n] = \sum_{k=-\infty}^{\infty} x[k]g[n-k].
\]

The signal is also decomposed simultaneously using a high-pass filter \( h \). The outputs giving the detail coefficients (from the high-pass filter) and approximation coefficients (from the low-pass). It is important that the two filters are related to each other and they are known as a quadrature mirror filter.

\textbf{Two Dimensional DWT (2-D)}

However, since half the frequencies of the signal have now been removed, half the samples can be discarded according to Nyquist’s rule. The filter outputs are then subsample by 2 (Mallat’s and the common notation is the opposite, \( g \) - high pass and \( h \) - low pass):

\[
y_{\text{low}}[n] = \sum_{k=-\infty}^{\infty} x[k] g[2n-k].
\]

\[
y_{\text{high}}[n] = \sum_{k=-\infty}^{\infty} x[k] h[2n-k].
\]

This decomposition has halved the time resolution since only half of each filter output characterizes the signal. However, each output has half the frequency band of the input so the frequency resolution has been doubled.

\textbf{E. 2-D Transform Hierarchy}

The generic form for a two-dimensional (2-D) wavelet transform is shown in Figure.

\textbf{The 1-D wavelet transform can be extended to a two-dimensional (2-D) wavelet transform using separable wavelet filters. With separable filters the 2-D transform can be computed by applying a 1-D transform to all the rows of the input, and then repeating on all of the columns.}

\textbf{Figure: Sub band Labeling Scheme for a one level, 2-D Wavelet Transform}
The original image of a one-level (K=1), 2-D wavelet transform, with corresponding notation is shown in the above figure. The example is repeated for a three-level (K =3) wavelet expansion in the below figure. In all of the discussion K represents the highest level of the decomposition of the wavelet transform.

![Sub-band labeling Scheme for a Three Level, 2-D Wavelet Transform](image)

The 2-D sub-band decomposition is just an extension of 1-D sub-band decomposition. The entire process is carried out by executing 1-D sub-band decomposition twice, first in one direction (horizontal), then in the orthogonal (vertical) direction. For example, the low-pass sub-bands (Li) resulting from the horizontal direction is further decomposed in the vertical direction, leading to LHi and LHi sub-bands. Similarly, the high pass sub-band (Hi) is further decomposed into HLi and HHi. After one level of transform, the image can be further decomposed by applying the 2-D sub-band decomposition to the existing LHi sub-band. This iterative process results in multiple “transform levels”. In Fig. 2.14 the first level of transform results in LH1, HL1, and HH1, in addition to LL1, which is further decomposed into LH2, HL2, HH2, LH2 at the second level, and the information of LL2 is used for the third level transform. The sub-band LLL is a low-resolution sub-band and high-pass sub-bands LHi, HLi, HHi are horizontal, vertical, and diagonal sub-band respectively since they represent the horizontal, vertical, and diagonal residual information of the original image.

F. Probabilistic Neural Networks (PNN):

Probabilistic (PNN) and General Regression Neural Networks (GRNN) have similar architectures, but there is a fundamental difference: Probabilistic networks perform classification where the target variable is categorical, whereas general regression neural networks perform regression where the target variable is continuous. If you select a PNN/GRNN network, DTREG will automatically select the correct type of network based on the type of target variable [3].

G. Architecture of a PNN

![Architecture of a PNN](image)

All PNN networks have four layers:

1. **Input layer** — There is one neuron in the input layer for each predictor variable. In the case of categorical variables, N-1 neurons are used where  is the number of categories. The input neuron (or processing before the input layer) standardizes the range of the values by subtracting the median and dividing by the interquartile range. The input neurons then feed the values to each of the neurons in the hidden layer [3].

2. **Hidden layer** — This layer has one neuron for each case in the training data set. The neuron stores the values of the predictor variables for the case along with the target value. When presented with the x vector of input values from the input layer, a hidden neuron computes the Euclidean distance of the test case from the neuron’s center point and then applies the RBF kernel function using the sigma value(s). The resulting value is passed to the neurons in the pattern layer[3].

3. **Pattern layer / Summation layer** — The next layer in the network is different for PNN networks and for GRNN networks. For PNN networks there is one pattern neuron for each category of the target variable. The actual target category of each training case is stored with each hidden neuron; the weighted value coming out of a hidden neuron is fed only to the pattern neuron that corresponds to the hidden neuron’s category. The pattern neurons add the values for the class they represent (hence, it is a weighted vote for that category)[3].

   For GRNN networks, there are only two neurons in the pattern layer. One neuron is the denominator summation unit the other is the numerator summation unit. The denominator summation unit adds up the weight values coming from each of the hidden neurons. The numerator summation unit adds up the weight values multiplied by the actual target value for each hidden neuron.

4. **Decision layer** — The decision layer is different for PNN and GRNN networks.

   For PNN networks, the decision layer compares the weighted votes for each target category accumulated in the pattern layer and uses the largest vote to predict the target category.
For GRNN networks, the decision layer divides the value accumulated in the numerator summation unit by the value in the denominator summation unit and uses the result as the predicted target value[4].

H. How PNN network work

Although the implementation is very different, probabilistic neural networks are conceptually similar to K-Nearest Neighbor (k-NN) models. The basic idea is that a predicted target value of an item is likely to be about the same as other items that have close values of the predictor variables [3]. Consider this figure:

Assume that each case in the training set has two predictor variables, \( x \) and \( y \). The cases are plotted using their \( x,y \) coordinates as shown in the figure. Also assume that the target variable has two categories, positive which is denoted by a square and negative which is denoted by a dash. Now, suppose we are trying to predict the value of a new case represented by the triangle with predictor values \( x=6, y=5.1 \). Should we predict the target as positive or negative?

Notice that the triangle is position almost exactly on top of a dash representing a negative value. But that dash is in a fairly unusual position compared to the other dashes which are clustered below the squares and left of center. So it could be that the underlying negative value is an odd case.

The nearest neighbor classification performed for this example depends on how many neighboring points are considered. If 1-NN is used and only the closest point is considered, then clearly the new point should be classified as negative since it is on top of a known negative point. On the other hand, if 9-NN classification is used and the closest 9 points are considered, then the effect of the surrounding 8 positive points may overbalance the close negative point.

A probabilistic neural network builds on this foundation and generalizes it to consider all of the other points. The distance is computed from the point being evaluated to each of the other points, and a radial basis function (RBF) (also called a kernel function) is applied to the distance to compute the weight (influence) for each point. The radial basis function is so named because the radius distance is the argument to the function [9].

Weight = RBF (distance)
The further some other point is from the new point, the less influence it has.

Radial Basis Function

Different types of radial basis functions could be used, but the most common is the Gaussian function:

VI. CONCLUSION

This study was undertaken to develop a PNN to classify stage of the brain tumor images and detect the Tumor using clustering technique. Grey level index values were assigned to the pixels of the indexed image and used as PNN inputs. There were 15 images, for training, and 8 images for testing. Probabilistic Neural Network with image and data processing techniques was employed to implement an automated Brain Tumor classification [7]. Decision making was
performed in two stages: feature extraction using GLCM and the classification using Probabilistic Neural Network (PNN). This paper presents a segmentation method, K-Means clustering algorithm, for segmenting Magnetic Resonance images to detect the Brain Tumor in its early stages. Although the study was limited by the available computational resources and training data, the results indicate the potential of ANNs for fast image recognition and classification. Fast image recognition and classification can be useful in the control of real-world, site-specific herbicide application.

- The paper has been appreciated by all the users in the organization.
- It is easy to use, since it uses the GUI provided in the user dialog.
- User friendly screens are provided.
- It also provides the user with variable options in customizing the packet capture.
- It has been thoroughly tested and implemented.

The presented samples demonstrate that the initial aim of the library was achieved - it is flexible, reusable, and it is easy to use it for different tasks. Although, there is still much work to do, because of a great range of different neural network architectures and their learning algorithms, but still - the library can be used for many different problems, and can be extended to solve even more[7]. I hope the library will become useful not only in my further research work, but other different researchers will find it interesting and useful.

REFERENCES


