

Semiautomated segmentation of bone marrow biopsies images based on texture features and Generalized Regression Neural Networks

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Abstract

This work presents preliminary results of a method for semi-automatic detection of fat and hematopoietic cells as well as trabecular surfaces in bone marrow biopsies, in order to calculate the percentage of each type of tissue or cell area in relation to the whole area.

Experimental results using selected clinical cases are presented. Twenty six biopsies were used, presenting varied distributions of cellularity and trabeculae topography. The approach is based on Digital Image Processing techniques and a Neural Network used for classification using textural features obtained from biopsies images. Results were improved with Mathematical Morphology filters.

The algorithm produces highly satisfactory results. The method was shown to be faster and more reproducible than conventional ones, like region growing, edge detection, split and merging.

The results from this computer-assisted technique are compared to others obtained by visual inspection by two expert pathologists, and differences of less than 9 % are observed.

Keywords

Digital Image Processing – Segmentation – Classification – Texture – Generalized Regression Neural Networks

1. Introduction

One of the most interesting fields in Digital Image Processing is the segmentation of an image into its different objects (Gonzalez and Woods, 1993). Segmentation plays a vital role in numerous biomedical imaging applications, such as the quantification of tissue volumes, diagnosis, localization of pathologies, study of anatomical structures and others (Glasbey 1995). The segmentation techniques can be divided into two groups: techniques based on contour detection which search for local grey level discontinuity in the image and those involving region growing which seek homogeneous image parts according to statistical measurements such as grey level and texture. The segmentation process of medical images is a difficult task to be accomplished in digital image processing (Chalama 1997).

Histopathology is an area normally considered in purely descriptive terms. However, a quantitative approach may bring it to more solid grounds. This is definitely true for bone pathologies (Frisch 1985). Changes in the amount of total bone and osteoid, together with the activity of fat and hematopoietic cells as well as trabecular surfaces, are probably the most important features in metabolic bone disease (Bullough 1990). The manual analysis of bone marrow is tedious and a time-consuming task that can be simplified by means of an automatic method (Revell 1983). With this automation better statistical information can be obtained (Clermonts 1985).

The goal of this work is the building of a tool to support pathology reports generation where the quantification and classification biopsies items are present. The use of a computational method to achieve classification and cell counting is very useful to help in qualitative and quantitative aspects of final diagnosis.

Numerous segmentation methods for cell images have been proposed, most of them trying to detect and separate different types of cells (Wu 1999, Sobrevilla 1999, De Medeiros 2001, Park 1997). Although they are useful antecedents of this work, the advantage of the method we propose is that it also produces good results for segmentation of trabecular surfaces.

The method consists in the application of a semi-automatic simple pattern classification algorithm based on the texture that images present.

Texture segmentation is a long established research field in image processing (Russ 1995). Texture means the subjective impression of the appearance of a surface. The classification based on texture feature values has become the essential technique for the treatment of some medical images (Baeg 1998, Serón 2002, Kneitz 1996, Nasser Esgiar 1998). Main research fields search for well-suited texture feature calculation methods and appropriate classification techniques.

For texture recognition, the image is divided into small sub-images. From the gray levels of each sub-image, textural features are computed. Many methods for the computation of these have been proposed so far (Singh 2001, Singh 2002). These methods can be divided into some main groups, such as statistical features, frequency-domain based features, fractal features and those derived from Wavelets and Gabor decomposition.

The results obtained with statistical features and frequency-domain based features are presented in this work (Haralick 1973). Fractal analysis using the Hurst exponent and fractal dimension (Cross

1994, Cross 1993, Dougherty 2001) was tried in an earlier stage, but the preliminary segmentation obtained was not successful in the marrow biopsies images.

Classical automatic schemes for classification like K-means and Fuzzy-C-means did not give satisfactory results when applied to marrow biopsies images. Consequently we propose a semi – automatic mapping process achieved with Generalized Regression Neural Networks (GRNN).

This work compares the results obtained using statistical (Gray level co-occurrence matrix) and frequency-domain (spectral power density) based features.

In a second stage of the method, morphological filters, derived from Mathematical Morphology theory, were used to improve the segmentation. Mathematical Morphology refers to a branch of nonlinear image processing and analysis, which focuses on the geometric structures within an image (Serra 1982). It emerged as a general theory to providing a unified approach in order to deal with problems in Medicine, Biology, and many other fields and can be applied in a satisfactory way to the resolution of fat cells segmentation problems (Dougherty 1992).

The proposed method is described in the next sections.

2. Materials and Methods

2.1. Image Acquisition

The images were obtained from bone marrow biopsies. They were chosen among normal samples and bone marrow biopsies that showed different disorders.

This work is based on twenty-six images. These were obtained from an Optic Microscope Medilux-12 with a TC Plan Achromat 4X objective, N.A. 0.10 and digitized with a CCD Hitachi KP-C550 color camera. This camera has an effective pixel dimension of 682 (H) x 492 (V) and a wavelength range of 400 to 700 nm. It supports a video resolution of 430 TV scan lines which was sufficient to capture biopsy images of 640 x 480 pixels from the live video by a PC. After selecting the area of interest, a single image is recorded in each case.

The images were saved on Windows Bitmap format and converted to 8-bit gray scale.

2.2. Generalized Regression Neural Networks

The mapping process is achieved using the Generalized Regression Neural Networks (GRNN) proposed by Specht (Specht 1991). The GRNN architecture is rather similar to the Probabilistic Neural Networks, also known as Radial Basis Network Functions (RBF).

The GRNN trains faster than other multilayer architectures and they model arbitrary non – linear functions efficiently. However they demand larger computational availabilities and a longer recall lapse than the wide known multilayer perceptron.

The GRNN learning process is built within a hybrid paradigm, since it incorporates supervised and not supervised algorithm. The GRNN architecture presents three forward connection layers: a first layer of input cells, a hidden layer and an output layer. The neurons of the input layer send the

information to the intermediate layer, the cells of the hidden layer are activated with a function of the distance between the input pattern and the synaptic weights, stored in each cell (called centroid) via a radial gaussian function.

Each neuron j of the hidden layer stores a vectorial c_{ji} , the centroid, whose distances r_j it is calculated as the euclidean distance that separates to the vector of entrance x_i of the centroid:

$$r_j^2 = \|x - c_j\|^2 = \sum (x_i - c_{ji})^2 \quad \text{Ec. 1}$$

where x_i = net inputs and y_i = hidden layer outputs

The output of the neuron y_j is calculated applying a radial function, term that is applied to symmetrical functions, usually the gaussian:

$$\phi(r) = e^{-r^2 / 2\sigma^2} \quad \text{Ec. 2}$$

The output of the hidden neuron j is shown in Equation 3.

$$y_j = e^{-\sum_i (x_i - c_{ji})^2 / 2\sigma_j^2} \quad \text{Ec. 3}$$

Each node of the hidden layer specializes on an input region space and the whole set of input cells must totally cover the interest region.

The outputs of the hidden neurons, each corresponding to a space region pattern are weighted and normalized previously to be fed to the output layer. The transfer functions of the output layer neurons are lineal, and they calculate the pondered and normalized sum of the hidden layer output (Equation 4).

$$z_k = \frac{\sum_j w_{kj} y_j}{\sum_j w_{kj}} \quad \text{Ec. 4}$$

The learning process consists of two stages, given that the different layers of this network perform different tasks, so it is reasonable to separate the optimization of the hidden and output layer by using different techniques. The training process of the hidden layers is performed via a self-organized fashion, whereas the linear weights of the output layer are computed using a supervised learning rule. The network undergoes a hybrid learning process. The self-organized component of the learning algorithm serves to allocate network resources in a meaningful way by placing the centers of the radial-basis functions in only those regions of the space where significantly data are present. To select the hidden unit's centers the standard k-nearest-neighbor rule is used. For the supervised learning operation to compute the linear weights of the output layer, the error-correcting learning rule as the least mean square algorithm is commonly applied (Haykin, 1999).

2.2. Segmentation Process

The different types of tissues regions to identify are shown on Figure 1.

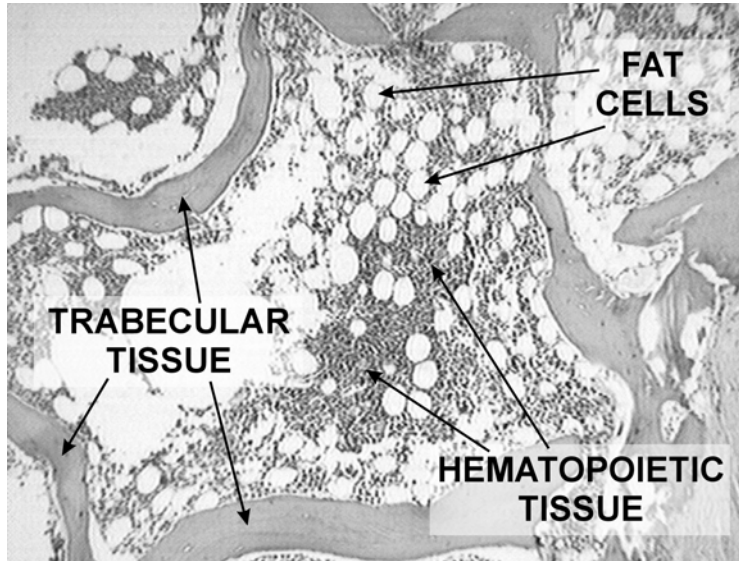


Figure 1: Typical image of a bone marrow biopsy.

The expert chooses N pixels, at least three, which correspond to each texture class to detect. The goal is to detect three classes corresponding to trabeculae surface, hematopoietic and fat cells.

Once the sample pixels are chosen, texture features corresponding to the region that surrounds each pixel are calculated. The size of the analysis region should be chosen appropriately, as it will be explained later on. The method was developed experimenting with two types of features:

- **Statistical Features:** they are based on different operations on Gray Level Co-occurrence Matrix (GLCM). After evaluating several feature combinations the vector was constructed as: $H = [cnth\ cntv\ cntd\ engh\ engv\ engd\ eph\ epv\ epd\ maxh\ maxv\ maxd\ mean]$, where:
 - **cnth, cntv, cntd:** Contrast at GLCM in horizontal (0°), vertical (90°) and diagonal (45°) direction respectively.
 - **engh, engv, engd:** Energy (order 0 differential moment) at GLCM in the same directions.
 - **eph, epv, epd:** Entropy at GLCM in the same directions.
 - **maxh, maxv, maxd:** Maximal Probability at GLCM in the same directions.
 - **mean:** Ratio of gray mean value to maximum of the range.
- **Frequency Features:** the vector in this case is composed by the values obtained after application of Fourier Transform and the later Spectral Density Power calculation. The values of the first and fourth quadrants of the transform are arranged in a vector.

A vector of features for each selected pixel is obtained, without any normalization applied. These are the input patterns that the GRNN requires in order to be created and trained. Its respective outputs will be the values of gray that will be shown for each class: 1, 127 and 256.

Subsequently, the image is processed by regions of the same size. The successive regions could be overlapped. The same type of texture features used at the sample pixels are calculated for each region.

Let x be the feature vector of the region that is being analyzed. This vector will be the input of the trained GRNN. The output given by the network will be a number in the range [1, 256] which indicates the gray level for the classification image.

Thus the classification image is created. This image will be processed to have only three gray values, making a simple transformation: the values in [1, 64] will be transformed to 1, the values in [65, 192] will be transformed to 127 and the values in [193, 256] will be transformed to 256. The final image is obtained, with 3 gray intensities.

Parameters are chosen empirically using the set of images:

- **Amount of sample pixels:** The number of prototypes patterns characterizing every class to segment.
- **Region size:** The size of the window where texture features are calculated.
- **Region Overlapping:** It defines the window displacement.
- **Features Vector:** It can be calculated by Fourier Transform of the region or by statistical analysis (co-occurrence matrix).
- **The spread value:** an internal parameter of the GRNN.

2.3. Post-processing

The segmented image may contain wrongly identified regions. This error is mainly due to the fact that in some regions more than one texture may be present. To improve the accuracy of the segmentation it is useful to apply some Mathematical Morphology filters.

The application of the different Morphological filters should be interactively indicated by the user, always matching the segmentation image with the original image.

The options are:

- **Dilation of one of the classes:** it allows enlarging some of the segmented regions without losing their shape.
- **Erosion of one of the classes:** it achieves the opposite effect to the dilation, diminishing the area covered by the class that is indicated.
- **Closing:** if during the detection erroneous holes appear within one of the classes, this operation will close them.
- **Elimination of small areas:** it is made by combination of morphological filters to eliminate objects of a certain connectivity whose area is smaller than a given threshold.

2.4. Area Calculation

Once the segmentation image is obtained, the calculation of relative areas is made according to the number of identified pixels of each class. User intervention is allowed again to select a region of

interest (ROI) of the image where there are not artifacts, and the segmentation has been satisfactory. The ROI have a polygonal shape.

Given the region of interest, the class area percentages are calculated (trabecular tissue, fat cells, hematopoietic tissue), according to the next expression:

$$\% \text{ Class } i \text{ Area} = \frac{\text{Amount of class } i \text{ pixels}}{\text{Amount of ROI pixels}} \times 100$$

2.5. Implementation

The proposed technique was implemented on MatLab[®] 6.5. The Morphological Filters was extracted from the SDC Morphology Toolbox[®].

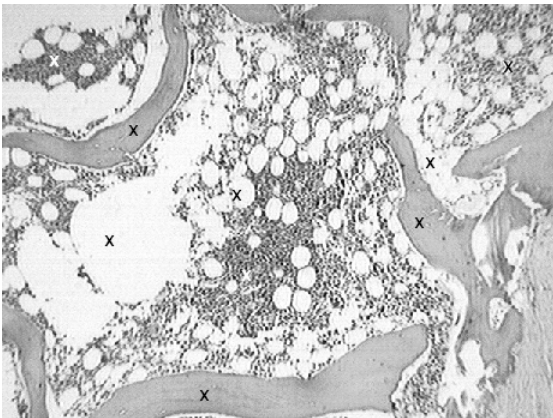
To facilitate the test stage and to have an appropriate way to interact with pathologists, a user-friendly interface was constructed. This shows the original image, the segmentation image and the results of the application of every filter.

The pathologist observes morphological filters with appropriate medical sentences that are easily understood by the operator such as “Close of holes the trabeculae region”, “Increase fat area” and “Decrease cellularity area”.

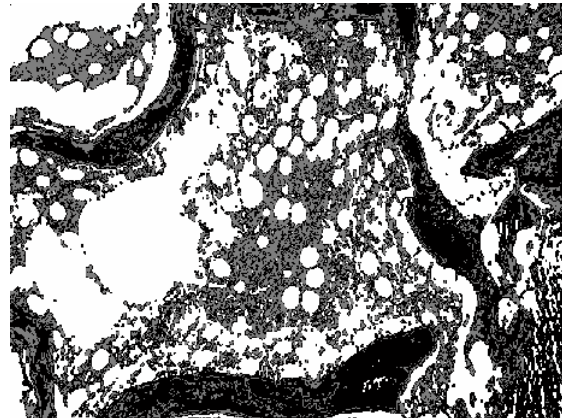
3. Results

Figure 2 shows the step sequence of the proposed method for one sample image. Figure 2 (a) shows the original image, with the pixels of three classes selected by a user. Figure 2 (b) shows the segmented image using frequency texture features. Figure 2 (c) presents the improvement achieved by morphological methods. Figure 2 (d) shows the chosen region for the calculation of areas. The numerical results obtained for this example are:

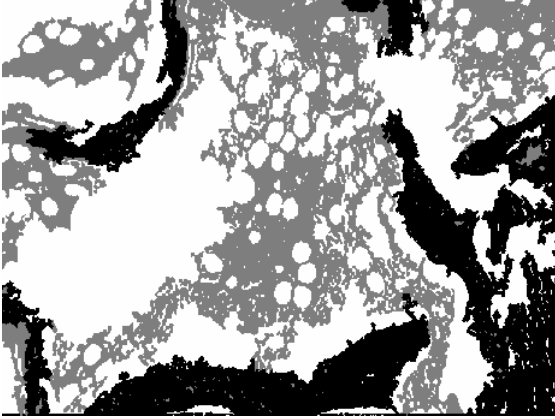
Trabeculae Tissue	= 19 %
Hematopoietic Tissue	= 50 %
Fat Cells	= 31 %



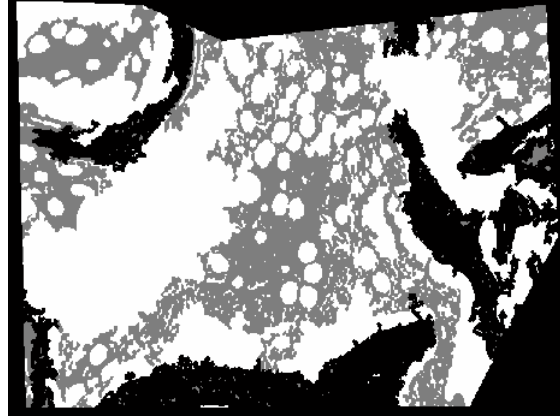
(a) Original image and sample pixels.



(b) Segmented image using frequency texture features.



(c) Improvement achieved by morphological methods.



(d) Chosen region for calculation of areas.

Figure 2: Steps sequence of the proposed method.

The same process was applied to a whole set of images. Table I shows the obtained values by the proposed method and the results that the pathologists have reported. For comparison, the average of the quantities they gave is taken, since these quantities varied as much as 35 %, which shows the subjectivity of the visual analysis.

To make a comparison between the results of the pathologists, the ones obtained using statistical features and the ones obtained using frequency features, three vectors were constructed for each image, where its components are the percentages obtained for each class (trabeculae, fat and hematopoietic cells).

In order to evaluate the difference between the pathologists results vector and the results obtained with the presented method, Euclidean vectorial distances were calculated. To make this value a significant index, it was necessary to normalize it. The normalization was calculated as:

$$SE \% = \frac{d(V_P, V_{SF})}{d(V_{Min}, V_{Max})} \%$$

$$FE \% = \frac{d(V_P, V_{FF})}{d(V_{Min}, V_{Max})} \%$$

where:

- V_P = Vector of results suggested by the pathologist.
- V_{SF} = Vector of results obtained using statistical features.
- V_{FF} = Vector of results obtained using frequency features.
- V_{Min} = Vector of minimal values of the range (0, 0, 0).
- V_{Max} = Vector of maximal values of the range (100, 100, 100).

The errors are shown in the last two columns in Table I. Empty cells indicate that the method didn't work suitably.

		Pathologist	Statistical Features	Frequency Features	SE %	FE %
Image 1	Trabeculae	11	16	17	5%	5%
	Fat	87	80	80		
	Hematopoietic	2	4	3		
Image 2	Trabeculae	18	26	24	6%	5%
	Fat	79	73	73		
	Hematopoietic	3	1	3		
Image 3	Trabeculae	18		24		5%
	Fat	77		71		
	Hematopoietic	5		5		
Image 4	Trabeculae	33		32		2%
	Fat	60		63		
	Hematopoietic	7		5		
Image 5	Trabeculae	70	54	75	13%	4%
	Fat	5	4	5		
	Hematopoietic	25	42	20		
Image 6	Trabeculae	64	49	66	14%	3%
	Hematopoietic	6	3	2		
	Fat	30	48	32		
Image 7	Trabeculae	60	57		2%	
	Hematopoietic	8	10			
	Fat	32	33			
Image 8	Trabeculae	35	36	38	4%	5%
	Hematopoietic	35	39	39		
	Fat	30	25	23		
Image 9	Trabeculae	26	22	27	3%	2%
	Hematopoietic	54	57	51		
	Fat	20	21	22		
Image 10	Trabeculae	24	17	23	7%	1%
	Hematopoietic	52	62	52		
	Fat	24	21	25		
Image 11	Trabeculae	26		15		22%
	Hematopoietic	53		33		
	Fat	21		52		
Image 12	Trabeculae	19	15	22	4%	2%
	Hematopoietic	47	46	47		
	Fat	34	39	31		
Image 13	Trabeculae	20	32	17	9%	6%
	Hematopoietic	57	48	51		
	Fat	23	20	32		
Image 14	Trabeculae	21	24	20	10%	9%
	Hematopoietic	39	49	50		
	Fat	40	27	30		
Image 15	Trabeculae	17	18	15	8%	4%
	Hematopoietic	41	50	46		
	Fat	42	32	39		
Image 16	Trabeculae	19	19	12	16%	9%
	Hematopoietic	27	47	39		
	Fat	54	34	49		
Image 17	Trabeculae	15		20		6%
	Hematopoietic	45		49		
	Fat	40		31		
Image 18	Trabeculae	19	16	16	6%	2%
	Hematopoietic	35	44	38		
	Fat	46	40	46		
Image 19	Trabeculae	12	21	30	14%	18%
	Hematopoietic	20	31	27		
	Fat	68	48	43		
Image 20	Trabeculae	10	18	15	13%	6%
	Hematopoietic	18	29	21		
	Fat	72	53	64		

Image 21	Trabeculae	30	35	37	14%	19%
	Hematopoietic	15	29	34		
	Fat	55	36	29		
Error Mean					8.7%	6.8%

Table I: Numerical results

4. Discussion

The automatic bone marrow microscope analysis presents several difficulties due to the diversity of elements appearing in this kind of images (Baak 1991). The different types of tissues and cells frequently are overlapped and the presence of artifacts complicates the problem. In most situations, this difficulty forces the use of very specific segmentation algorithms for each task. This makes bone marrow image segmentation a difficult and challenging problem (Liu 1999).

The accuracy of segmentation mainly depends on the selection of good parameters. After experiencing with images, some criteria can be determined for the choice of parameters.

- **Amount of sample pixels:** It is chosen keeping in mind that a big quantity of sample pixels possibly improves the classification, but it will increase the processing time.
- **Size of the region:** if it is small, processing time will decrease, but it is risky in the sense that different textures can not be segmented correctly. On the other hand, if it is too big, apart from increasing the calculation time considerably, the probability of the analysis region containing more than one texture increases, surely leading to classification errors.
- **Region Overlapping:** if the window of analysis is moved in a single pixel step manner, a lengthy processing time will be required, and the result will not be notably better than the result obtained using bigger displacements. For example, moving the window by two pixels decreases calculations by one half, and the classified regions will have a size of 2 x 2 pixels, what allows an equally successful segmentation.
- **Calculated Features:** features calculated in the frequency domain leads to a noisier segmentation, but after a simple post-processing, acceptable results are achieved. The unquestionable advantage of the frequency domain is the fact that processing time decreases drastically if compared with co-occurrence features. The statistical features based on gray level co-occurrence matrix results in a clearer classification, but processing time increases considerably.

As mentioned before, the results obtained were compared to the reports obtained by visual inspection performed by two pathologists. The discrepancy between the two pathologists' reports justifies the use of the method we propose which gives differences smaller than 9 % if compared with the average of the results given by experts.

A method for texture segmentation using two different features vectors was proposed. This method improves considerably the efficiency in the recognition of the quantity of trabecular tissue, fat and hematopoietic cells present in microscopic images from biopsies. The method has been evaluated on twenty-six images collected from a Pathology Laboratory. The obtained results showed a good level of accuracy in the 81 % of them.

A tool providing such capabilities can reduce counting error produced by a subjective analysis and can shorten the time taken by manual counting. Besides that, it avoids inter and intra observer variability.

A need for automation is evident. But once optimized the sequence to follow for the processing of an image, a quick segmentation and area calculation process is obtained. It would facilitate the professional's work, decreasing diagnosis errors and minimizing subjectivity in bone marrow analysis.

For future work, it will be of interest to make new comparisons applying the presented method to images obtained at different magnifications and stained with different techniques.

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6. References

- Baak J (1991): Manual of Quantitative Pathology in Cancer Diagnosis and Prognosis. New York, Springer-Verlag.
- Baeg S, Popov A, Kamat V, Batman S, Sivakumar K, Kehtarnavaz N, Dougherty ER, Shah RB (1998): Segmentation of Mammograms into Distinct Morphological Texture Regions. IEEE Symposium on Computer-Based Medical Systems, Lubbock, June 1998.
- Bullough PG, Bansal M, DiCarlo EF (1990): The tissue diagnosis of metabolic bone disease. Role of histomorphometry. *Orthop Clin North Am*; 21:65-79.
- Chalama V, Kim Y (1997): A methodology for evaluation of boundary detection algorithms on medical images. *IEEE Trans Med Imag*; 16:642-652.
- Clermonts E, Birkrnhager-Frenkel D (1985): Software for bone histomorphometry by means of a digitizer. *Comput Methods Programs Biomed*; 21:185-94.
- Cross S (1994): The Application of Fractal Geometric Analysis to Microscopic Images. *Micron*; 25 (1):101-113
- Cross S, Rogers S, Silcocks PB, Cotton DWK (1993): Trabecular bone does not have a fractal structure on light microscope examination. *J Pathol*; 170:311-313.
- De Medeiros Martins A, Duarte Dória Neto A, Medeiros Brito A, Sales A, Jane S (2001): Texture based segmentation of cell images using neural networks and mathematical morphology. *Proceedings of IJCNN'2001, Washington, D.C., IEEE / Omnipress*, pp. 2489-2494.
- Dougherty E (1992): *An Introduction to Morphological Image Processing*. Washington, SPIE.
- Dougherty G, Henebry, G (2001): Fractal signature and lacunarity in the measurement of the texture of trabecular bone in clinical CT images. *Med Eng Phys*; 23:369-380.
- Frisch B, Lewis SM, Burkhardt R, Bartl R (1985): *Biopsy Pathology of Bone and Bone Marrow*. New York, ed. Raven Press.

- Glasbey C, Horgan, G (1995): *Image Analysis for the Biological Sciences*. New York, John Wiley & Sons.
- Gonzalez R, Woods R (1992): *Digital Image Processing*. USA, Addison – Wesley, pp. 458-465.
- Haralick RM, Shanmugan K, Dinstein I (1973): Textural features for image classification. *IEEE Transactions on Systems, man and Cybernetics*; SMC-3:610-621.
- Haykin S (1999): *Neural Networks: A Comprehensive Foundation*. Prentice-Hall.
- Kneitz S, Ott G, Albert R, Schindewolf T, Muller-Hermelink HK, Harms H (1996): Differentiation of Low Grade Non-Hodgkin's Lymphoma by Digital Image Processing. *Analyt Quant Cytol Histol*; 18:121-128.
- Liu Z, Liew HL, Clement JG, Thomas DL (1999): Bone Image Segmentation. *IEEE Trans Biomed Eng*; 46 (5):565-573.
- Nasser Esgiar A, Raouf N, Naguib G, Bennett MK, Murray A (1998): Automated Feature Extraction and Identification of Colon Carcinoma. *Analyt Quant Cytol Histol*; 20:297-301
- Park JS, Keller JM (1997): Fuzzy Patch Label Relaxation in Bone Marrow Cell Segmentation. *Proc. IEEE International Conference on Systems, Man, and Cybernetics, Orlando, FL., October 1997*, pp. 1133-1138.
- Revell P (1983): Histomorphometry of bone. *J Clin Pathol Review*; 36: 1323-1331.
- Russ, JC (1995): *The Image Processing Handbook*. USA, CRC Press, pp. 361-367
- Serón D, Moreso F, Gratin C, Vitriá J, Condom E, Grinyó JM, Alsina J (2002): Automated Classification of Renal Interstitium and Tubules by Local Texture Analysis and a Neural Network. *Analyt Quant Cytol Histol*; 24:147-153.
- Serra J (1982): *Image Analysis and Mathematical Morphology*. London, Academic Press.
- Singh M, Singh S (2001): Evaluation of Texture Methods for Image Analysis. *Proc. 7th Australian and New Zealand Intelligent Information Systems Conference, Perth, November 2001*.
- Singh M, Singh S (2002): Spatial Texture Analysis: A Comparative Study. *Proc. 15th International Conference on Pattern Recognition (ICPR'02), Quebec, August 2002*.
- Sobrevilla P, Keller J, Montseny E (1999): White Blood Cell detection in Bone Marrow images. *Proc. of 18th NAFIPS International Conference, New York, USA, June 1999*, pp 403-407.
- Specht, D (1991): A General Regression Neural Network. *IEEE Transactions on Neural Networks*, 2 (6): 588-576.
- Tou J, Gonzalez R (1974): *Pattern Recognition Principles*. New York, Addison – Wesley.
- Wu, H-S; Gil, J; Jeremiah, R; Barba, J (1999): Unsupervised approach for segmentation of textured cytologic images. *Electronics Letters, IEE*, April 1999; 35 (8):630-631.