Diving Deep onto Discriminative Ensemble of Histological Hashing & Class-Specific Manifold Learning for Multi-class Breast Carcinoma Taxonomy*

Sawon Pratiher§ and Subhankar Chattoraj‡

§Department of Electrical Engineering, Indian Institute of Technology Kharagpur, WB, India †Department of Electronics & Communication Engineering, Techno India University, WB, India

Abstract—Histopathological images (HI) encrypt resolution dependent heterogeneous textures & diverse color distribution variability, manifesting in micro-structural surface tissue convolutions & inherently high coherency of cancerous cells posing significant challenges to breast cancer (BC) multi-classification. As such, multi-class stratification is sparsely explored & prior work mainly focus on benign & malignant tissue characterization only, which forestalls further quantitative analysis of subordinate classes like adenosis, mucinous carcinoma & fibroadenoma etc, for diagnostic competence. In this work, a fully automated, near-real-time & computationally inexpensive robust multi-classification deep framework from HI is presented.

The proposed scheme employs deep neural network (DNN) aided discriminative ensemble of holistic class-specific manifold learning (CSML) for underlying HI sub-space embedding & HI hashing based local shallow signatures. The model achieves 95.8% accuracy pertinent to multi-classification, an 2.8% overall performance improvement & 38.2% enhancement for Lobular carcinoma (LC) sub-class recognition rate as compared to the existing state-of-the-art on well known BreakHis dataset is achieved. Also, 99.3% recognition rate at $200 \times \&$ a sensitivity of 100% for binary grading at all magnification validates its suitability for clinical deployment in hand-held smart devices.

Index Terms—breast cancer; image hash; sub-space learning; discriminative ensemble; deep learning; deep neural networks.

I. INTRODUCTION

Burgeoning cancer statistics in 2018 from World Cancer Report (WCR) estimate 2 million new cases of BC being registered worldwide [1]. World Health Organization (WHO) projects 627,000 women died from BC (which is approximately 15% of all cancer related deaths among women in 2018) & impacts 2.1 million women each year [2]. Significant research spanning imaging technique like computed tomography (CT), mammography, HI anatomization & magnetic resonance (MR) have been developed for early-stage BC prognosis through precision medicine initiative, but HI analysis is taken as the "gold standard" due to its rich encoded histological morphology & substantiating abnormal cellular activity. Additionally, deformation dynamics of these spatial tissue textures allows more specific characterizations from a diagnostic perspective [3] & aid pathologists to control growth & metastasis of tumor cells & devise therapeutic clinical schedules which are specific to particular sub-classes.

However, owing to ubiquitously abundant inhomogeneous morphology & complex intricate spatial correlations in the underlying inter-weaved biological tissue fabric of biopsy samples [4], automated machine vision for robust & accurate multi-class BC detection is still a challenging task & eludes researchers. Manual BC multi-classification by the pathologist

is arduous & requires domain expertise with interpretations being subjective in nature. An automated computer-aided diagnostic (CAD) system overcome these challenges & assist clinicians in reliable diagnosis by reducing their workload & avoid erroneous diagnosis [5-6]. Further, practical CAD systems face challenges of inefficient hand-crafted feature engineering, which is computationally intensive & time consuming. Secondly, optimal supervised feature selection for accurate BC identification via image segmentation & identification of primitives like nuclei deformation, tubule formation, lymphocytes presence etc..[7] & high-resolution HI analysis is computationally expensive requiring costly high-performance computation, which is generally not available in developing countries. Subtle inter-class & intra-class variability w.r.t., contrast & textures are evident in fig. 1. Fig. 1 shows representative Hematoxylin & Eosin (H&E) stained fine-grained multiclass biopsy tissue slides. (Taken from *BreaKHis* @400× magnification [8]). To resolve these challenges, a reliable &

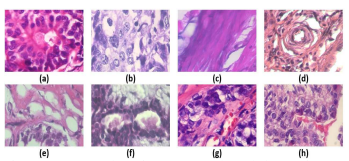


Figure 1: (a) Ductal carcinoma (DC), (b) Lobular carcinoma (LC), (c) Mucinous carcinoma (MC), (d) Papillary carcinoma (PC), (e) Adenosis, (f) Fibroadenoma, (g) Tubular adenoma (TA), (h) Phyllodes tumor (PT). (a) to (d) & (e) to (g) corresponds to malignant and benign class respectively.

more accurate practical method for BC multi-classification via DNN is developed. The proposed method discards feature engineering & employs end-to-end training of deep discriminative ensemble of holistic CSML of HI & local high-level to low-level semantic hierarchical hash signatures followed by softmax layer for classification. The model is validated with 7909 HI's to demonstrate its potency for deployment in clinical settings on generic processors & experimental result shows superior performance. The main contributions can be recapitulated as:

 A novel & scalable DNN for BC multi-classification framework is explored. The model, as shown in Fig. 2, yields the highest recognition rate with reduced computational time & complexity as compared to state-of-art.

^{*}This paper is accepted for presentation at 44^{th} International Conference on Acoustics, Speech, & Signal Processing (IEEE ICASSP), UK, 2019

The scheme strengthen the intra-class morphological similarities & inter-class dissimilarities of the hierarchical feature space in a discriminative manner via deep ensemble of CSML & heterogeneous local hash signatures.

II. MATERIALS & PRIOR WORK

A. BreakHis Dataset

The proposed BC multi-classification method is examined on publicly available large-scale BreaKHis dataset [8] containing 7,909 H&E stained microscopic images from surgical biopsy (SOB) breast tumors, taken from 82 patients & collected at multiple magnification factor of: $40 \times ,100 \times ,200 \times \&400 \times$. The images are of 700×460 pixels dimension with 24-bit color depth in 3-channel RGB (Red-Green-Blue) format. Table 1 summarizes the distribution of different histological sub-types & a detailed description can be traced from [8].

Table I: Distribution of classes, sub-classes in *BreaKHis* @ different magnification factors.

		Magnification Factor					
Class	SC	40 ×	100×	200×	400×	Total	Patient
	DC	864	903	896	788	3451	
M	LC	156	170	163	137	626	
	MC	205	222	196	169	792	58
	PC	145	142	135	138	560	
	A	114	113	111	106	444	
В	F	253	260	264	237	1014	
	TA	109	121	108	115	453	24
]	PT	149	150	140	130	569	
Tot	al	1995	2081	2013	1820	7909	82

B. Prior Art using BreaKHis

Significant research is concentrated around binary classification of benign & malignant classes. Over the past few years, researches have broadly investigated optimal feature engineering & deep learning based architectures. These can be found from the works of Spanhol et al., [8], ensemble classifier (EC) of shallow features by Gupta et al., [9], multiple feature vector (MFV) & transfer learning [10], graph-manifold & BI-LSTM models by Pratiher et al., [11], Grassmann manifold (GM) aided vector of locally aggregated descriptors (VLAD) by Dimitropoulos et al., [12] and convolution neural network (CNN) model with fusion rule (FR) [13]. Efficacy of ConvNet based fisher vector (CFV) & Gaussian mixture model (GMM) by Song et al., [14], deep CNN by Wei et al., [15] and BC classification using incremental boosting convolution networks in [16]. An exhaustive comparative study can be traced from Table III. Studies concerning multi-classification of sub-classes for clinical diagnosis or prognosis is done by Han Zhongyi et al., using class structured deep CNN (CSD-CNN) model [15] & Bardou et al. using CNN based approach [17]. Multi-classification BC histological research comparison can be found in Table IV & V.

III. METHODOLOGY & RELATED THEORY

A. Experiment Design: **Deep Discriminative Ensemble of Histological Hashing & Class-Specific Manifold Learning**

Recently, DNN has shown efficacy in achieving state-ofthe-art performance in diverse research problems spanning

medical imaging [4], natural language processing (NLP) & speech processing. Here, we propose to use stacked sparse autoencoder (SSAE) based DNN [21] for robust BC multiclassification. Fig. 2 highlights the workflow. Our model pipeline consists of three main stages: Pre-processing stage for stain normalization, & overlapping & optimal patch segments of a specific size are generated for subsequent tissue index profile abstraction. Thereafter, class-specific manifold learning (CSML) of different histological sub-types are comprehended for nonlinear dimensionality reduction (NLDR). NLDR distillates discriminative low-dimensional structures pertinent to particular sub-class hidden in the high-dimensional HI. CSML preserves the intrinsic quasi-isometric geometry & local contour connectivity of HI point-cloud within tolerable limits via feature-space geometry constraints & is very much crucial for diagnosis. Thereafter, different hash signature obtained via discrete wavelet transform (DWT), singular-value decomposition (SVD) & perceptual feature-points augments the local shallow statistical HI descriptors. The holistic CSML & Hash vectors are fused in a discriminative fashion, which contemplates class structure based feature fusion. These ensemble discriminative super feature vectors are fed to SSAE for learning deep features & classification thereof.

B. Class-Specific Manifold Learning (CSML)

CSML is envisaged via Landmark Isomap (L-ISOMAP) aide Eigen sub-space estimation of a particular histological sub-type. For a vectorized HI point-cloud of Y data-points, arbitrarily $m(m \ll M)$ points are selected as Landmarks. Euclidean distance based embedding of the input data using m landmark points are selected from Y & followed by multidimensional scaling (MDS) using the $m \times m$ matrix $G_{m,M}$ of geodesic distances for each landmark pair to compute the low dimensional feature space. Mathematically, it is given by:

$$m^{T}(p,q) = -\frac{1}{2} \left(F_{pq}^{2} - e_{p} \frac{1}{m} \sum_{l} H_{pl}^{2} \right),$$
 (1)

where, H^2 is the means geodesic distance matrix H is element-wise squared & e_p is the Eigen vector with zero Eigen value. Details about L-ISOMAP can be traced from [22].

C. Histological Hashing for Local Signatures

Rudimentary image hashing details can be found in [23-26]. Histological image hashing encodes locality reference & inherent neighborhood connectivity of the underlying HI. Here, we have used the following hash signatures:

- 1) Discrete Wavelet Transform (DWT) Based Image Hash: Computes robust & compact hash via 2D DWT on HI, which decomposes into four sub-bands. Both edge or high frequency information & coarse stable low-frequent coefficients are perceived via DWT coefficients [23].
- 2) Hashing Via Singular Value Decomposition (SVD): SVD based image hasing has robust tolerance to small rotational changes until 10° & is translation in-variance & incorporates low rank approximation of original normalized sub-image of HI & non-correlated directional feature space encoding [24].
- 3) Feature Point Based Image Hashing: Statistical image features like Hessian affine, maximally stable extremal region (MSER) detectors, Harris corner detector & feature points

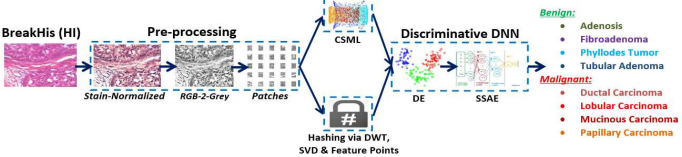


Figure 2: Stages of integrated pipeline: pre-processing, CSML & HI hash extraction, discriminative ensemble & DNN learning.

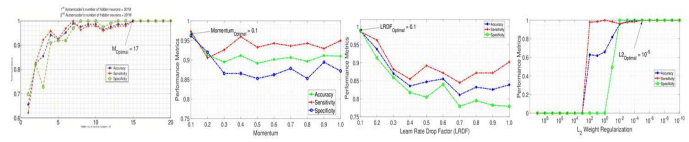


Figure 3: Hyper-parameter tuning in DNN. (a) hidden size, (b) momentum, (c) learn rate drop factor, (d) L_2 weight regularization

based on end-stopping behavior like end-stopped Wavelets such as Morlets preserves the momentous image geometry feature-space constraints of 2-D HI pixels & mapped to 1-D feature vector which is compressed to generate the hash vector [25].

D. SSAE aided DNN Training & Hyper-parameters Tuning

Discriminative feature ensemble of holistic CSML & shallow hash signatures is fused via discrimination correlation analysis [27]. Stacked Sparse Autoencoder (SSAE) [28] involves optimal parameter $\theta = (V, a_k, a_y)$ computation by minimizing the error between the model input and output. Rudimentary details can be traced from [21]. The model is trained on a generic system woth mode configuration as 3.50 GHz processor, 16GB RAM & AMD FX-8320 Octacore. Fig 3 shows hyper-tuned optimal SSAE parameters. A maximum epoch of 500 & number hidden layer neurons in the first and second autoencoders (AE) is kept at 500 & 300 respectively. Further, L_2 weight regularization, sparsity regularization & sparsity proportion in the two AE are set to 0.001, 4 & 0.15. 'tanh' activation function (AF) in the hidden units is connected to by fully connected layer & classification done via a softmax layer. 'tanh' AF is used due to its robust tolerance to approximate intrinsic manifold non-linearity & extraction of mutual dependence for further segregation thereof. Piece-wise back propagation learning via stochastic gradient descent (SGD) algorithm is envisaged with a learning rate of 10^{-4} & the initial random weights drawn uniformly from [-0:1;0:1], gradient decay factor of 0.2, momentum of 0.6, learning rate drop period (LRDP) of 5 & L_2 weight regularization of 10^{-4} is used during the training phase. The weights (W) & bias (b) are updated as

$$W_{l} = W_{l} - \eta \frac{\partial}{\partial W_{l}}(W, b; X, t) = B_{l}W_{l} - \eta \frac{\partial}{\partial B_{l}}(W, B; X, t)$$
(2)

where, W_l represents the weights, B_l represents the l^{th} layer bias, η is learning rate, (X,t) is the mini-batch comprising of 'm' training samples. In order to eschew skewed model overfitting & bias minimization, the data is randomly partitioned in the ratio 60:20:20 for to training, validation & testing phase.

IV. RESULTS AND DISCUSSION

The adequacy of the proposed method is evaluated in terms of standard evaluation metrics i.e., classification accuracy (AC), sensitivity (SN) & specificity (SP). Experimental results for both binary & multi-classification are compared with state-of-the-art techniques on benchmark *BreaKHis* dataset.

Table II: Results for Binary Classification. PM = Performance Metrics

Class	PM (%)	Magnification Factor					
		40X	100X	200X 400X			
	AC	99.1	98.7	99.3 98.4			
Binary	SN	100	100	100 100			
	SP	98.3	97.9	98.5 96.8			

A. Grading Tumor Malignancy: Binary Classification

Initially, binary classification of BC into benign & malignant classes is done to ensure its competence in coarse HI characterization. Table II highlights the performance measure of the experimental results for different magnification factors. 100% sensitivity for all magnification factor ensures that all malignant classes are recognized correctly. As such, pathologist may invest more time for identifying benign cases with our system & demonstrates the effectiveness of the DNN model to learn deep malignancy biomarkers for discriminative HI grading. A comparative study of the proposed mode with the existing state-of-the-art methods [8-20] is given in Table III. It may be noted that the proposed framework outperforms

all the previously used methods in terms of classification accuracy with a significant enhancement at all magnification factors. Further, our proposed discriminative DNN framework outperforms the conventional & visual feature descriptor based approach such LBP [8], VLAD [12] and KAZE [20] & also surpasses recent CNN & BiLSTM based methods [10-11][14-16], which is proved to be optimal in analyzing visual imagery. This indicates that our proposed system which is robust in terms of performance & computationally much more efficient as it runs on generic laptops as compared to CNN models which requires graphics processing unit (GPU).

Table III: State-of-the-art comparison for binary classification of benign and malignant classes

Ref, Years	Feature + Method	Performance (%)			
[40X	100X	200X	400X
[13], 2016	CNN, FR	85.6	83.5	82.7	80.7
[8] 2016,	CLBP, SVM	77.40	76.40	70.20	72.80
[9], 2017	C-TID, EC	87.2	88.22	88.89	85.82
[10], 2017	CNN, DeCAF, MFV	84.6	84.8	84.2	81.6
[11], 2017	GML, BI-LSTM	96.2	97.2	97.1	95.4
[12], 2017	VLAD, GM	91.8	92.1	91.4	90.2
[14], 2017	I-EM, CFV, CNN, GMM	87.7	87.6	86.5	83.9
[15], 2017	CSDCNN	95.8	96.9	96.7	94.9
[16], 2018	DCNN	95.1	96.3	96.9	93.8
[17], 2018	Dense SIFT, SURF, BOW, LCLC	98.33	97.12	97.85	96.15
[18], 2018	FV, CSE	87.5	88.6	85.5	85.0
	TL, DenseNet	84.72	89.44	95.65	82.65
[19], 2018	CNN, DenseNet	91.90	93.64	95.84	90.15
[MV, XGboost	94.71	95.9	96.76	89.11
[20]	KAZE, BOF, binary SVM	85.9	80.4	78.1	71.1
This Work	CSML, Hashing DNN	99.1	98.7	99.3	98.4

B. Multi-classification Performance

It's evident from literature survey that there are very few literature available in the multi-class classification of BC from HI & as such, we further examine the proposed framework towards the multi-class classification of BC to demonstrate its efficacy towards practical use from a clinical perspective. State-of-the-art comparative evaluation is given in Table IV, whereas Table V gives class specific performance accuracy. It is evident that the proposed discriminative DNN framework not only surpasses the state-of-art on the basis of magnification factor but also on the basis of HI sub-classes in terms of recognition rate. Earlier in the state-of-art, CSDCNN [15] based approach exhibited best recognition rate for multi-class classification but our proposed method surpasses it in all the magnification factor by 3.3%, 1.8%, 2.6%, 3.5%, for $40\times$, $100\times$, $200\times$ & $400\times$ magnification. Table V shows that our method surpasses the class-specific recognition rate with high margin & in some case as high as 38.2% enhancement for Lobular carcinoma (LC) sub-class.

Table IV: State-of-the-art comparison for multi-Classification histological sub-types

Ref	Feature + Method	Performance (%)				
1		40X	100X	200X	400X	
[15]	CSDCNN	92.8	93.9	93.7	92.9	
	DSIFT + BoW	41.80	38.56	49.75	38.67	
	SURF + BoW	53.07	60.80	70.00	51.01	
	DSIFT + LLC	60.58	57.44	70.00	49.96	
	SURF + LLC	80.37	63.84	74.54	54.70	
[17]	DSIFT, BoW +SVM	18.77	17.28	20.16	17.49	
	SURF, BoW+ SVM	49.65	47.00	38.84	29.50	
	DSIFT, LLC+SVM	48.46	49.44	43.97	32.60	
	SURF, LLC+ SVM	55.80	54.24	40.83	37.20	
	CNN, SVM-RBF	75.43	71.20	67.27	65.12	
This Work	CSML, Hashing, DNN	95.1	95.7	95.8	95.2	

Table V: Class-specific performance comparison with [17]

Tuote V. Class specific performance comparison with [17]						
		Magnification Factor				
Class	Ref, years	SC	40 ×	100×	200×	400×
	[17], 2018	DC	91.51	90.77	91.14	92.74
	This Work	DC	96.7	97	97.6	96.9
Malignant	[17], 2018	LC	78.72	54.90	63.27	56.10
	This Work	LC	93.8	94.7	92.8	93.1
	[17], 2018	MC	70.49	82.09	61.02	70.59
	This Work	MC	94.4	95.8	96.6	94.9
	[17], 2018	PC	67.44	83.72	57.50	68.29
	This Work	PC	93.1	95.2	93	93.1
	[17], 2018	A	85.29	79.41	84.85	90.63
	This Work	A	93.5	93.9	97	94.1
Benign	[17], 2018	F	86.84	91.03	91.14	77.46
	This Work	F	95	95.6	94.9	95.1
	[17], 2018	TA	75.56	93.33	76.19	82.05
	This Work	TA	94.5	94.1	94.3	94.4
	[17], 2018	PT	76.19	63.89	62.50	58.82
	This Work	PT	94.4	95.1	95.2	94.7

V. CONCLUSION

A novel deep discriminative ensemble learning CAD for multi-class BC characterization is introduced in this work. The method implements deep contextual grading of hybrid holistic-level CSML representations & local hash signatures of HI, thereby, effectively discriminating between benign & malignant sub-classes. The proposed approach was validated using *BreakHis* dataset & experimental results exemplify superior discriminating performance as compared to the existing state-of-the-art. In particular, it shows high specificity towards malignant sub-classes, which can assist pathologists by reducing their heavy workload & arrange optimal therapeutic schedules for further diagnosis or prognosis of benign tissues.

Currently, the method is being escalated to include deeper structures using graph CNN & sequential contextual learning with other tissue images to investigate the diagnostic modality.

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