



Gigapixel Whole-Slide Images Classification Using Locally Supervised Learning

Jingwei Zhang¹(✉), Xin Zhang¹, Ke Ma², Rajarsi Gupta¹, Joel Saltz¹,
Maria Vakalopoulou³, and Dimitris Samaras¹

¹ Stony Brook University, New York, USA

{jingweizhang, xin.zhang.3, samaras}@cs.stonybrook.edu,
{Rajarsi.Gupta, Joel.Saltz}@stonybrookmedicine.edu

² Snap Inc., California, USA

kemma@cs.stonybrook.edu

³ CentraleSupélec, University of Paris-Saclay, Gif-sur-Yvette, France
maria.vakalopoulou@centralesupelec.fr

Abstract. Histopathology whole slide images (WSIs) play a very important role in clinical studies and serve as the gold standard for many cancer diagnoses. However, generating automatic tools for processing WSIs is challenging due to their enormous sizes. Currently, to deal with this issue, conventional methods rely on a multiple instance learning (MIL) strategy to process a WSI at patch level. Although effective, such methods are computationally expensive, because tiling a WSI into patches takes time and does not explore the spatial relations between these tiles. To tackle these limitations, we propose a locally supervised learning framework which processes the entire slide by exploring the entire local and global information that it contains. This framework divides a pre-trained network into several modules and optimizes each module locally using an auxiliary model. We also introduce a random feature reconstruction unit (RFR) to preserve distinguishing features during training and improve the performance of our method by 1% to 3%. Extensive experiments on three publicly available WSI datasets: TCGA-NSCLC, TCGA-RCC and LKS, highlight the superiority of our method on different classification tasks. Our method outperforms the state-of-the-art MIL methods by 2% to 5% in accuracy, while being 7 to 10 times faster. Additionally, when dividing it into eight modules, our method requires as little as 20% of the total gpu memory required by end-to-end training. Our code is available at https://github.com/cvlab-stonybrook/local_learning_wsi.

Keywords: Locally supervised learning · Whole slide image · Multiple instance learning · Classification

J. Zhang and X. Zhang—Contributed equally to this paper.

Supplementary Information The online version contains supplementary material available at https://doi.org/10.1007/978-3-031-16434-7_19.

© The Author(s), under exclusive license to Springer Nature Switzerland AG 2022
L. Wang et al. (Eds.): MICCAI 2022, LNCS 13432, pp. 192–201, 2022.
https://doi.org/10.1007/978-3-031-16434-7_19

1 Introduction

Computational pathology involving observation of tissue slides with a microscope, is the gold standard for cancer diagnosis. In recent years, digital pathology has emerged as a powerful technology for digitizing whole slide images (WSIs) for assessment, sharing and analysis [7]. This provides researchers a good opportunity to develop computer-aided analysis systems for various levels of applications, such as cell counting, gland segmentation, and WSI classification [6, 9, 13]. In particular, WSI-based cancer diagnosis faces unique challenges. The most typical characteristics of WSIs are their extremely large image size and high resolution. A WSI generally can be as large as $100,000 \times 100,000$ pixels at a 40X magnification, which makes it impractical to train deep neural networks in an end-to-end (E2E) manner. Consequently, the most popular methods nowadays follow a patch-based paradigm [4, 12], i.e. each WSI is first tiled into thousands of small patches. Then a model extracts and aggregates patch-level features to make the final prediction [14, 21, 23].

Such methods follow a Multiple Instance Learning (MIL) scheme which is currently the state-of-the-art for solving histopathology classification tasks [3, 10, 14, 21, 25]. Zhang et al. [25] proposed a spatial and magnification based attention sampling strategy to extract informative patches, and directly learned a WSI classification model on these patches. DSMIL [14] jointly trained a patch and an image classifier, where the patches are selected softly with instance-level attention. More recently, TransMIL [21] presented a transformer-based MIL framework to explore both morphological and spatial information among instances. However, such a technical paradigm has some intrinsic shortcomings; these methods do not explore the spatial relations of each tile, by failing to properly combine the local and global information of the tumor’s microenvironment. Moreover, these methods rely on pretrained features to represent the tiles since the current deep learning architecture cannot be trained in an end-to-end manner.

The end-to-end training of deep neural networks requires storing in memory the entire computational graph as well as the layer activations during the forward pass. Then the loss backpropagates and updates the weights layer by layer based on the chain rule. Storage of the graph and the gradients occupy a large amount of GPU memory, limiting the input image size. Some researchers [20, 22] proposed to retain gradient information and train the model part by part to reduce memory consumption. Nevertheless, they still tiled images into patches, and trained networks on smaller regions with limited receptive field sizes.

Due to E2E training’s limited scalability to large input and large architectures, recent research attempts to seek alternatives to mitigate the memory constraints, among which locally supervised learning attracts increasing interest [2, 24]. Locally supervised learning aims to train each layer locally with a pre-defined objective function, without back-propagating the gradients end-to-end. The network training is free from storing *all* intermediate variables and the memory consumption is thus reduced. Belilovsky et al. [1] attached an auxiliary convolutional neural network classifier at each local module to predict the final target and evaluated it on ImageNet [5]. Nøkland et al. [18] proposed to use both classification loss and contrastive loss to supervise each local module and showed it was better than using a single loss.

In this paper, we introduce a locally supervised learning paradigm to train a classification network using the entire WSI. Our method splits a deep network into multiple

gradient-isolated modules and each part is trained separately with local supervision. Thus we can use the entire WSI as input and do not have a patch size limited receptive field. Moreover, we further propose the Random Feature Reconstruction (RFR) model to boost the performance and optimize the GPU usage. To the best of our knowledge, we are the first to propose a locally supervised learning scheme coupled with RFR for the classification of entire WSIs. Our method has been extensively evaluated on three public WSI datasets, and achieves state-of-the-art performance compared to MIL-based methods. Moreover, without tiling, our method is significantly faster during inference.

2 Method

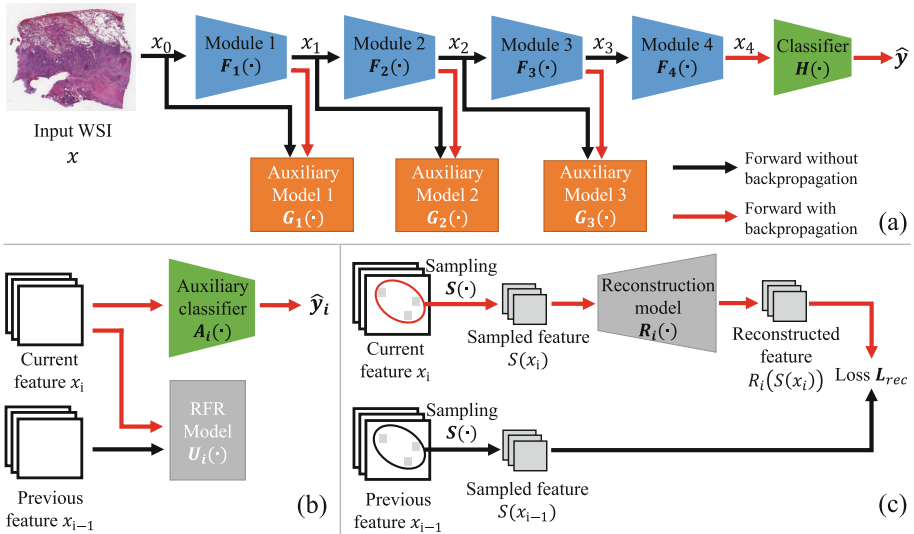


Fig. 1. Overview of the proposed method. (a) Overall structure of our locally supervised learning method. A network is divided into 4 modules and a classifier. The first 3 modules $F_i, i = 1, 2, 3$ are optimized using auxiliary models $G_i, i = 1, 2, 3$ respectively. The last module F_4 is optimized together with the classifier $H(\cdot)$. We assume $x_0 = x$. (b) Structure of the auxiliary model $G_i(\cdot)$. It has an auxiliary classifier $A_i(\cdot)$ and a Random Feature Reconstruction (RFR) model $U_i(\cdot)$. (c) Structure of the RFR model. It reconstructs randomly sampled regions in the previous feature map.

The key idea of our locally supervised learning is dividing a network layer by layer into several consecutive modules and optimizing them separately. Formally, let us assume without loss of generality, a network $F(\cdot)$ composed by K consecutive modules: $F(\cdot) = ((H \circ F_K) \circ F_{K-1} \circ \dots \circ F_1)(\cdot)$, where $F_i(\cdot)$ represents the i -th network module. $H(\cdot)$ is a gated attention multiple instance learning [11] based classifier and \circ is the function composition operation. Such a network is trained using pairs of (x, y) on which x denotes the entire WSI and y the corresponding label.

A network module contains several network layers of the original network, for example, the first 6 layers in a ResNet34. The input to a network module $F_i, i = 1, \dots, K$

is x_{i-1} and the output is x_i , assuming $x_0 = x$. An overview of our approach is presented in Fig. 1 on which the forward and backward passes are indicated.

In such a setup, each module is trained locally. More specifically, given a network module F_i and its input x_{i-1} , we use an auxiliary model G_i , and compute the loss as $\mathcal{L}_i = G_i(F_i(x_{i-1}), x_{i-1}, y) = G_i(x_i, x_{i-1}, y)$. We train F_i by minimizing the \mathcal{L}_i . Then, the trained module F_i is frozen and in an iterative process, the same technique is applied to F_{i+1} by minimizing $\mathcal{L}_{i+1} = G_{i+1}(x_{i+1}, x_i, y)$. The same process is applied to each of the $K - 1$ modules. Finally, the final module $F_K(\cdot)$ is optimized together with the classifier $H(\cdot)$ without an additional auxiliary model since the label y serves as the final supervision.

2.1 Auxiliary Model

The training of each model is performed using an auxiliary model with a greedy strategy [1]. As shown in Fig. 1(b), the auxiliary model has two parts: an auxiliary classifier $A_i(\cdot)$ and an RFR model $U_i(\cdot)$. The auxiliary classifier has a similar structure of classifier $H(\cdot)$ and computes a classification loss $\mathcal{L}_{cls}(\hat{y}_i, y)$ between the prediction $\hat{y}_i = A_i(x_i)$ and ground truth y . Such a design enables the training of the module $F_i(\cdot)$ locally. However, as discussed in [24], the shallower layers in a network have limited ability to extract discriminative features, making the training difficult.

To overcome this problem, the authors proposed to reconstruct the input image x from the feature map x_i and applied a reconstruction loss as a regularization to preserve the discriminative features. However, this strategy cannot be applied to WSIs as reconstructing an entire WSI is too costly.

To deal with this issue, we propose to use a RFR model instead of the reconstruction module. More specifically, a RFR model reconstructs randomly sampled regions from the previous feature map.

As shown in Fig. 1(c), the first step of a RFR model $U_i(\cdot)$ is to randomly sample $(S(\cdot))$, 10 corresponding spatial locations on the latent representations from the i -th module x_i as well as from the previous module x_{i-1} . Feature patches $S(x_i)$ and $S(x_{i-1})$ are cropped according to the sampled spatial locations. Then, a reconstruction network $R_i(\cdot)$ is applied to the cropped features from i -th module $S(x_i)$, aiming to reconstruct the target $S(x_{i-1})$. A reconstruction loss \mathcal{L}_{rec} is used to minimize the distance between the reconstructed feature patches $R_i(S(x_i))$ and feature patches from its previous module in the corresponding spatial locations $S(x_{i-1})$. During training, this random sampling process eventually iterates over most locations and thus encourages the network to preserve discriminative features with limited GPU memory cost.

2.2 Optimization

In our framework, the first $K - 1$ modules are optimized locally with the following setting:

$$\mathcal{L}_i = \mathcal{L}_{cls}(A_i(F_i(x_{i-1})), y) + \alpha \mathcal{L}_{rec}(R_i(S(F_i(x_{i-1}))), S(x_{i-1})), \quad (1)$$

$$i = 1, \dots, K - 1, \quad (2)$$

where x_{i-1} is the input feature of the i -th module $F_i(\cdot)$ and hyperparameter α is a regularization term. Since the last module $F_K(\cdot)$ is jointly optimized with the classifier $H(\cdot)$ the training scheme is changed to:

$$\mathcal{L}_K = \mathcal{L}_{cls}(H(F_K(x_{K-1})), y), \quad (3)$$

where x_{K-1} is the input feature of the last module. We used L1 loss as the reconstruction loss \mathcal{L}_{rec} and cross entropy loss as the classification loss \mathcal{L}_{cls} . We set α to be 1 after grid search on the validation dataset. However, our methodological design is independent of these losses and different reconstruction and classification losses can be applied.

3 Experiments and Discussion

3.1 Datasets

TCGA-NSCLC. The TCGA-NSCLC (The Cancer Genome Atlas-Non-Small Cell Lung Cancer) dataset includes two sub-types of lung cancer, Lung Adenocarcinoma (LUAD) and Lung Squamous Cell Carcinoma (LUSC). The dataset contains a total of 1053 diagnostic WSIs. We randomly split them into 663 training slides, 166 validation slides and 214 testing slides (10 slides without magnification labels are discarded). We benchmarked the performance of our model on this dataset for the lung cancer sub-type classification task. The WSIs were on 5X magnification and the size of the slides ranges from 1581×1445 to 23362×11345 .

TCGA-RCC. The TCGA-RCC (Renal Cell Carcinoma) dataset includes three sub-types of kidney cancer, Kidney Chromophobe Renal Cell Carcinoma (KICH), Kidney Renal Clear Cell Carcinoma (KIRC) and Kidney Renal Papillary Cell Carcinoma (KIRP). The dataset contains a total of 939 diagnostic digital slides. We randomly split them into 603 training slides, 150 validation slides and 186 testing slides. We benchmarked the performance of our model on this dataset for the classification of these three different kidney cancer types. The WSIs were on 5X magnification and the size of the slides ranges from 2610×1351 to 23849×10257 .

LKS. The Liver-Kidney-Stomach(LKS) [17] dataset is a multi-tissue indirect immunofluorescence slides dataset. It includes four classes: Negative, Anti-Mitochondrial Antibodies (AMA), Vessel-Type Anti-Smooth Muscle Antibodies (SMA-V) and Tubule-Type Anti-Smooth Muscle Antibodies (SMA-T). The dataset contains a total of 684 slides, including 205 testing slides. To perform our experiments, we further split the rest into 383 training slides and 96 validation slides. Each slide in this dataset has the original size of 40000×40000 . We further resized the images to 10000×10000 at 5X magnification.

3.2 Implementation Details

For all our experiments, we used ResNet34 [8] pretrained on ImageNet [5] as our backbone network ($F(\cdot)$). We froze the first 4 layers of ResNet34 and increased the stride of the first convolution from 2 to 3 to enlarge the receptive field. We set the batch size

to be 1, since each WSI has a different size. To mitigate potential instability, we used the common optimization practice of accumulating the gradients of 8 batches before updating the parameters. All the batch normalization layers were frozen as well. In the RFR model, we sampled 10 patches of the size 128×128 for the first auxiliary model $G_1(\cdot)$ and the spatial dimensions sampled in the following modules depended on the size of the feature map. The number and size of patches are determined by the validation dataset. We used a gated attention multiple instance learning (GABMIL) [11] based network in the auxiliary classifier $A_i(\cdot)$ and classifier $H(\cdot)$.

We used AdamW [15] with weight decay 10^{-6} as the optimizer. For the two TCGA datasets, the learning rate was initially set to 1×10^{-5} for the pre-trained backbone modules $F_i(\cdot)$ and 2×10^{-5} for the randomly initialized auxiliary models $G_i(\cdot)$ and classifier $H(\cdot)$. Learning rates were decreased by a factor of 0.1 when the loss and validation accuracy plateaued. For the LKS dataset, the initial learning rates were doubled to 2×10^{-5} for pre-trained backbone modules and 4×10^{-5} for randomly initialized auxiliary models.

We used the PyTorch library [19] and trained our models on a NVIDIA Tesla V100 or a Nvidia Quadro RTX 8000 GPU.

3.3 Results

Evaluation of Overall Performance. We chose overall accuracy and area under Receiver Operating Characteristic curve (AUROC) as the main metrics to evaluate our method. Our baselines included ImageNet pre-trained ResNet34 with two different pooling methods: average pooling and max pooling. We also included the current state-of-the-art deep MIL models: the attention based multiple instance learning (ABMIL) [11] and its gated variant GABMIL [11], dual stream attention based model

Table 1. Comparison of accuracy and AUROC on three datasets. Our method, of both $K = 4$ and $K = 8$, outperforms existing state-of-art MIL models

Dataset	TCGA-NSCLC		TCGA-RCC		LKS	
Metric	Accuracy	AUROC	Accuracy	AUROC	Accuracy	AUROC
Max-pooling	0.8318	0.9036	0.8495	0.9306	0.8049	0.9366
Avg-pooling	0.7944	0.8669	0.8172	0.9309	0.6000	0.9086
ABMIL [11]	0.8037	0.8816	0.8495	0.9423	0.8341	0.9392
GABMIL [11]	0.8364	0.8762	0.8602	0.9535	0.8146	0.9399
MIL-RNN [3]	0.8178	0.9011	/	/	/	/
DSMIL [14]	0.8271	0.8909	0.8710	0.9590	0.8390	0.9328
CLAM-SB [16]	0.8224	0.9185	0.8763	0.9701	0.8293	0.9446
CLAM-MB [16]	0.8598	0.9131	0.8763	0.9716	0.8439	0.9448
StreamingCNN [20]	0.8692	0.9260	0.8817	0.9660	0.8927	0.9652
Ours (K=4)	0.8785	0.9377	0.9140	0.9740	0.8976	0.9562
Ours (K=8)	0.8785	0.9322	0.9032	0.9760	0.8829	0.9633

Table 2. Comparison of GPU memory consumption. Our method required around 20% ($K = 8$) to 30% ($K = 4$) GPU memory compared to E2E training. * is measured on CPU because of GPU memory limitation

Image size	8698×7496	12223×10057	23849×10257
E2E	17.89G	33.63G	78.14G*
Ours ($K = 4$)	5.35G	9.64G	18.32G
Ours ($K = 8$)	3.71G	6.47G	11.85G

DSMIL [14], single-attention-branch CLAM-SB [16], multi-attention-branch CLAM-MB [16], and also two-stage recurrent neural network based aggregation MIL-RNN [3], which considers binary classifications only. All these baselines are trained on 5X resolution and using ResNet34 for fair comparison. Our method was able to fine tune the ImageNet pretrained weights to adapt to the medical image domain, while other methods directly used the ImageNet pretrained features.

As shown in Table 1, our method ($K = 4$) outperformed all the compared methods in the overall accuracy and AUROC metrics. To the best of our knowledge the SOTA for the TCGA-NSCLC dataset is 96.3% AUROC (95% confidence interval: 93.7%-99.0%) reported by CLAM [16]. For experimental uniformity, we used the exact same splits on all comparisons and reported accuracy and AUROC, reporting a 91.9% AUROC for CLAM for the same resolution which is also higher than the rest of the compared methods. Our method achieved 1.87% higher accuracy than the best compared method CLAM-MB, and 2.46% higher AUROC. On the TCGA-RCC dataset, our method achieved 3.77% higher accuracy and 0.24% higher AUROC. On the LKS dataset, our method had 5.37% higher accuracy and 1.14% higher AUROC compared with the best performing method CLAM-MB. Also, comparing with the SOTA on this dataset, SOS [17], our method achieved comparable performance to it (90.73% accuracy).

Moreover, Table 1 highlights the robustness of our method with respect to the different modules. In particular, our model was divided into 8 modules and each of them trained locally using the proposed strategy performs as well as the $K = 4$ and outperformed the compared methods. Also, we compared our method with a non-MIL approach StreamingCNN [20], as shown in Table 1, our method ($K = 4$) outperformed it on TCGA-NSCLC and TCGA-RCC. On LKS dataset, our method achieved better accuracy and comparable AUROC.

Evaluation of GPU Memory Consumption. Another major advantage of our method is that our method significantly reduced the GPU memory required during training and thus enables training on the entire WSI. We compared the GPU memory consumption of our method (for $K = 4$ and $K = 8$) and that of the end-to-end (E2E) training. As the sizes of images in a WSI dataset usually vary a lot, instead of evaluating the memory consumption on three datasets, we evaluated it on three different sized images: a 23849×10257 image, the largest image in the TCGA-RCC dataset, a 12223×10057 image, and an 8698×7496 image. Note that for the 23848×10257 image, we performed the E2E measurement on the CPU since the GPU memory was not enough to perform

Table 3. Comparison of inference speed on three different sized inputs. Our method ran 7 to 10 times faster than GABMIL since we do not have the tiling and feature extraction step.

Image size	8698 \times 7496			12223 \times 10057			23849 \times 10257		
Method	GABMIL	Ours	Speed gain	GABMIL	Ours	Speed gain	GABMIL	Ours	Speed gain
Tiling	0.3 s	/	/	0.7 s	/	/	1.8 s	/	/
Features	2.6s	/	/	3.7s	/	/	9.6s	/	/
Prediction	<0.1 s	0.3 s	/	<0.1 s	0.6 s	/	<0.1 s	1.2 s	/
Total	2.9 s	0.3s	9.7x	4.4 s	0.6 s	7.3x	11.4 s	1.2s	9.5x

this task. As shown in Table 2, when the input image size was 8698×7496 , our 4 divided network required only 29.9% of the GPU memory that the E2E training needs. This number further dropped to 20.7% if we divided the network into $K = 8$ modules.

The same memory usage held for the other two input image sizes. In general, our 4 divided network required only around 30% memory and our 8 divided network required only around 20% memory compared to E2E training.

Evaluation of Time Efficiency. Besides the higher accuracy and the lower GPU memory cost, our method is faster in inference than the standard MIL approaches. We measured the total time (in seconds) that our method requires to infer an entire WSI and compared it with GABMIL [11], a high performance MIL model. We timed the whole inference pipeline including patch tiling, feature extraction, and final prediction. We reported the total inference time on WSIs in 3 different sizes. Table 3 highlights the time efficiency of our method. Our method took only 1.2 s to classify a 23849×10257 WSI, while GABMIL needed more than 11.4 s due to the time consuming step of feature extraction on the large amount of patches.

Ablation Study on RFR. We conducted an ablation study on the Random Feature Reconstruction (RFR) model. Table 4 shows the comparison on the accuracy of our method with and without RFR. On the TCGA-NSCLC dataset, using the RFR model improved the accuracy by 0.47% for $K = 4$ and 3.27% for $K = 8$. On the TCGA-RCC dataset, using the RFR model improved the accuracy by 2.15% for $K = 4$ and 3.22% for $K = 8$. On the LKS dataset, using the RFR model improved the accuracy by around 1% for both $K = 4$ and $K = 8$.

Table 4. Comparison of accuracy of our method with and without RFR. The RFR model improved the accuracy of our method by 1% to 3%.

Dataset	TCGA-NSCLC	TCGA-RCC	LKS
K = 4, w/o RFR	0.8738	0.8925	0.8829
K = 4, w. RFR	0.8785	0.9140	0.8976
K = 8, w/o RFR	0.8458	0.8710	0.8780
K = 8, w. RFR	0.8785	0.9032	0.8829

4 Conclusion

In this paper, we introduced a locally supervised learning framework to train using entire whole slide images. We evaluated it on three WSI datasets and achieved a 2% to 5% accuracy improvement compared to existing MIL methods. This significant performance gain was achieved by reducing GPU memory consumption and enabling fine-tuning of the feature extractor. Compared with end-to-end training, our method required only 20% to 30% of the memory. Moreover, our method did not require tiling as existing MIL methods do, thus it was 7 to 10 times faster during inference. We also demonstrated that the proposed random feature reconstruction (RFR) model improved the performance of our locally supervised learning framework by 1% to 3%. Our proposed approach showed the greater potential of locally supervised learning on classifying whole slide images and we will explore its applications on other tasks including segmentation.

Acknowledgements. This work was partially supported by the ANR Hagnodice ANR-21-CE45-0007, the NSF IIS-2123920 award, Stony Brook Cancer Center donors Bob Beals and Betsy Barton as well as the Partner University Fund 4D Vision award.

References

1. Belilovsky, E., Eickenberg, M., Oyallon, E.: Greedy layerwise learning can scale to imagenet. In: International Conference on Machine Learning, pp. 583–593. PMLR (2019)
2. Belilovsky, E., Eickenberg, M., Oyallon, E.: Decoupled greedy learning of cnns. In: International Conference on Machine Learning, pp. 736–745. PMLR (2020)
3. Campanella, G., et al.: Clinical-grade computational pathology using weakly supervised deep learning on whole slide images. *Nat. Med.* **25**(8), 1301–1309 (2019)
4. Coudray, N., et al.: Classification and mutation prediction from non-small cell lung cancer histopathology images using deep learning. *Nat. Med.* **24**(10), 1559–1567 (2018)
5. Deng, J., Dong, W., Socher, R., Li, L.J., Li, K., Fei-Fei, L.: Imagenet: a large-scale hierarchical image database. In: 2009 IEEE Conference on Computer Vision and Pattern Recognition, pp. 248–255. IEEE (2009)
6. Deng, S., et al.: Deep learning in digital pathology image analysis: a survey. *Front. Med.* **14**(4), 470–487 (2020). <https://doi.org/10.1007/s11684-020-0782-9>
7. Dimitriou, N., Arandjelović, O., Caie, P.D.: Deep learning for whole slide image analysis: an overview. *Front. Med.* **6**, 264 (2019)
8. He, K., Zhang, X., Ren, S., Sun, J.: Deep residual learning for image recognition. In: Proceedings of the IEEE Conference on Computer Vision and Pattern Recognition, pp. 770–778 (2016)
9. Hou, L., Agarwal, A., Samaras, D., Kurc, T.M., Gupta, R.R., Saltz, J.H.: Robust histopathology image analysis: to label or to synthesize? In: Proceedings of the IEEE/CVF Conference on Computer Vision and Pattern Recognition, pp. 8533–8542 (2019)
10. Hou, L., Samaras, D., Kurc, T.M., Gao, Y., Davis, J.E., Saltz, J.H.: Patch-based convolutional neural network for whole slide tissue image classification. In: Proceedings of the IEEE Conference on Computer Vision and Pattern Recognition (CVPR), June 2016
11. Ilse, M., Tomczak, J., Welling, M.: Attention-based deep multiple instance learning. In: International Conference on Machine Learning, pp. 2127–2136. PMLR (2018)

12. Le, H.: Utilizing automated breast cancer detection to identify spatial distributions of tumor-infiltrating lymphocytes in invasive breast cancer. *Am. J. Pathol.* **190**(7), 1491–1504 (2020)
13. Lerousseau, M., et al.: Weakly supervised multiple instance learning histopathological tumor segmentation. In: Martel, A.L., et al. (eds.) *MICCAI 2020*. LNCS, vol. 12265, pp. 470–479. Springer, Cham (2020). https://doi.org/10.1007/978-3-030-59722-1_45
14. Li, B., Li, Y., Eliceiri, K.W.: Dual-stream multiple instance learning network for whole slide image classification with self-supervised contrastive learning. In: *Proceedings of the IEEE/CVF Conference on Computer Vision and Pattern Recognition*, pp. 14318–14328 (2021)
15. Loshchilov, I., Hutter, F.: Decoupled weight decay regularization. In: *International Conference on Learning Representations* (2018)
16. Lu, M.Y., Williamson, D.F., Chen, T.Y., Chen, R.J., Barbieri, M., Mahmood, F.: Data-efficient and weakly supervised computational pathology on whole-slide images. *Nat. Biomed. Eng.* **5**(6), 555–570 (2021)
17. Maksoud, S., Zhao, K., Hobson, P., Jennings, A., Lovell, B.C.: SOS: selective objective switch for rapid immunofluorescence whole slide image classification. In: *Proceedings of the IEEE/CVF Conference on Computer Vision and Pattern Recognition*, pp. 3862–3871 (2020)
18. Nøkland, A., Eidnes, L.H.: Training neural networks with local error signals. In: Chaudhuri, K., Salakhutdinov, R. (eds.) *Proceedings of the 36th International Conference on Machine Learning*. *Proceedings of Machine Learning Research*, 09–15 June, vol. 97, pp. 4839–4850. PMLR (2019)
19. Paszke, A., et al.: Pytorch: an imperative style, high-performance deep learning library. In: *Advances in Neural Information Processing Systems*, vol. 32 (2019)
20. Pinckaers, H., van Ginneken, B., Litjens, G.: Streaming convolutional neural networks for end-to-end learning with multi-megapixel images. *IEEE Trans. Pattern Anal. Mach. Intell.* **44**, 1581–1590 (2020)
21. Shao, Z., et al.: Transmil: transformer based correlated multiple instance learning for whole slide image classification. In: *Advances in Neural Information Processing Systems*, vol. 34 (2021)
22. Takahama, S., et al.: Multi-stage pathological image classification using semantic segmentation. In: *Proceedings of the IEEE/CVF International Conference on Computer Vision*, pp. 10702–10711 (2019)
23. Tellez, D., Litjens, G., van der Laak, J., Ciompi, F.: Neural image compression for gigapixel histopathology image analysis. *IEEE Trans. Pattern Anal. Mach. Intell.* **43**, 567–578 (2019)
24. Wang, Y., Ni, Z., Song, S., Yang, L., Huang, G.: Revisiting locally supervised learning: an alternative to end-to-end training. In: *International Conference on Learning Representations* (2020)
25. Zhang, J., et al.: A joint spatial and magnification based attention framework for large scale histopathology classification. In: *Proceedings of the IEEE/CVF Conference on Computer Vision and Pattern Recognition*, pp. 3776–3784 (2021)