Deep Adversarial Training for Multi-Organ Nuclei Segmentation in Histopathology Images

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Problem

- 1. Generate synthetic H&E images
- Sthat model the distribution of cellular and extracellular spatial features represented in multiple organs.
- 2. Use both the synthetic and real histopathology data for training a context-aware CNN that can accurately **segment nuclei**.

Find learning mapping functions
G and S between two domains:
M (nuclei mask) and N (H&E images)



 $G: M \rightarrow N$

(for synthetic data generation)



 $S: N \to M$ (for nuclei segmentation)

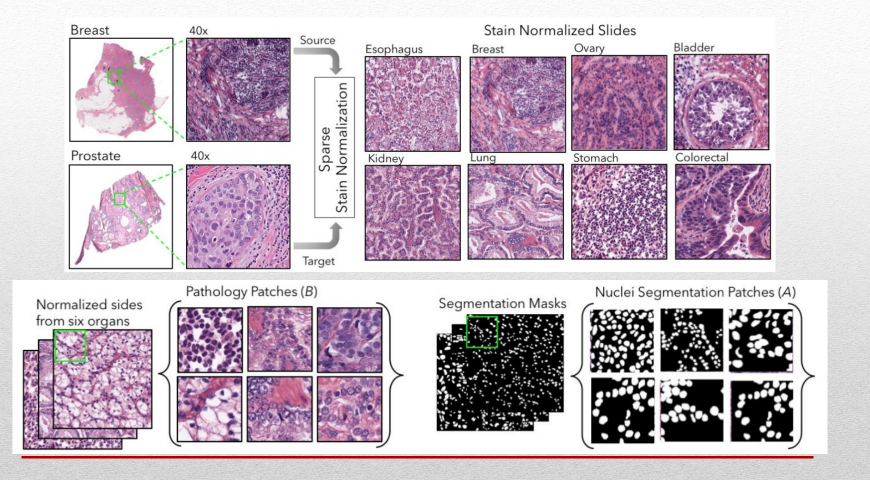
We denote m and n as training examples where $m \in M$ and $n \in N$.

Datasets

Multi organ pathology images (with annotations) of 34 1000 × 1000 pathology images from nine different organs (bladder, colon, stomach, breast, kidney, liver, and prostate, ovary, esophagus)



- Sparse stain normalization
- Decomposition into large patches of 256 x 256
- Training images: four slides from breast, liver, kidney and prostate
- Testing images: two slides from breast, liver, kidney, prostate, bladder, colon and stomach and one slide from esophagus and ovary
- Adding 4.650 synthetically generated images to training images (nuclei segmentation)



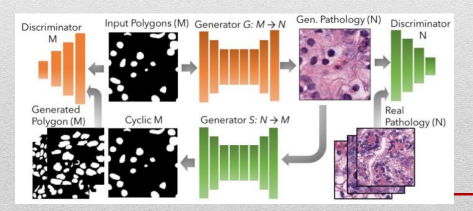
Problem 1: Synthetic data generation

Architectures: dual-GAN (with cycle consistency loss) → cycleGAN

Normalization: spectral normalization (to prevent mode colapse)

The cycleGAN framework learns a mapping between randomly generated polygon masks and unpaired pathology images. Since cycleGAN is based on consistency loss, the setup also learns a reverse mapping from pathology images to corresponding segmentation or polygon masks.

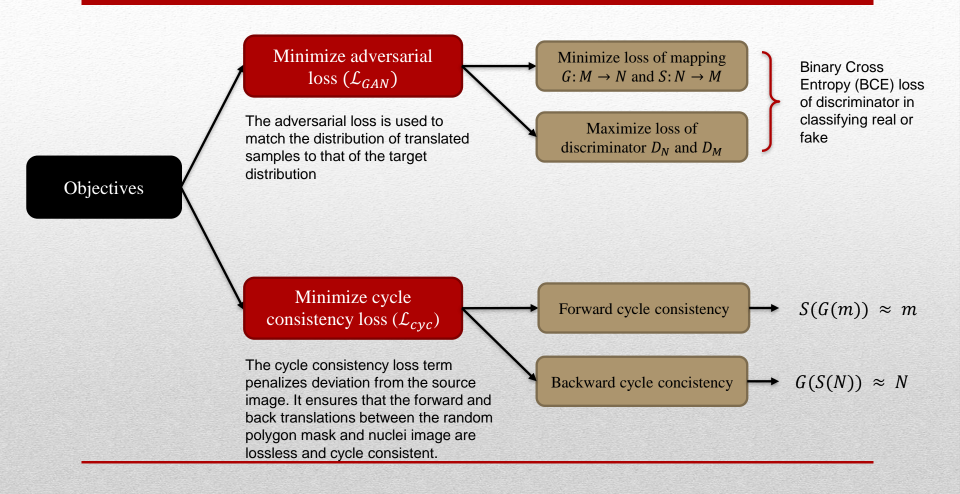
The reverse mapping is used only to train the forward mapping more effectively.



G: random polygon mask to pathology image generator

S: pathology image to polygon mask generator

 D_N : discriminator for G D_M :discriminator for S



Loss of mapping $G: M \to N$

$$\mathcal{L}_{GAN}(G, D_N) = \mathbb{E}_{n \sim p_{\text{data}}(n)}[\log D_N(n)] + \mathbb{E}_{m \sim p_{\text{data}}(m)}[\log(1 - D_N(G(m)))],$$

Loss of mapping $S: N \to M$

$$\mathcal{L}_{cyc}(G, S) = \lambda_n \mathbb{E}_{n \sim p_{\text{data}}(n)}[||G(S(n)) - n||_1]$$
$$+ \lambda_m \mathbb{E}_{m \sim p_{\text{data}}(m)}[||S(G(m)) - m||_1]$$

 λ controls the importance of the forward and backward cycle constraints. For synthetic data generation, the λ_m term is relaxed.

Loss of mapping $S: N \to M$

$$\mathcal{L}_{\text{GAN}}(S, D_M) = \mathbb{E}_{m \sim p_{\text{data}}(m)}[\log D_M(m)] + \mathbb{E}_{n \sim p_{\text{data}}(n)}[\log(1 - D_M(S(n)))].$$

full objective for synthetic data generation

$$\underset{G,S}{\operatorname{arg\,min}} \underset{D_N,D_M}{\operatorname{arg\,max}} \mathcal{L}_{\operatorname{GAN}}(G,D_N) + \mathcal{L}_{\operatorname{GAN}}(S,D_M)$$

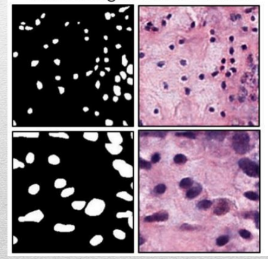
$$+ \mathcal{L}_{cyc}(G,S)$$

Training Details

- Pytorch 4.0
- $\lambda_n = 70, \lambda_m = 10$
- Optimizer: Adam solver
- Number of epoch: 300
- Normalization: spectral normalization
- Learning rate: 0.0002 (for first 150 epochs) → linearly decayed to 0 (for remaining epochs)
- Weight initialized from a Gaussian distribution with a mean 0 and standard deviation 0.02

Output

Synthetic Pathology Images with Nuclei Segmentation Masks

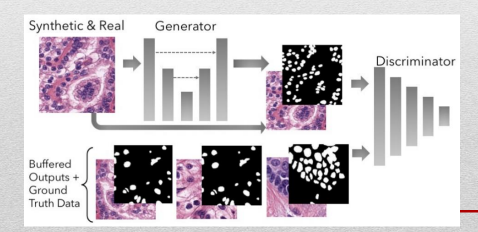


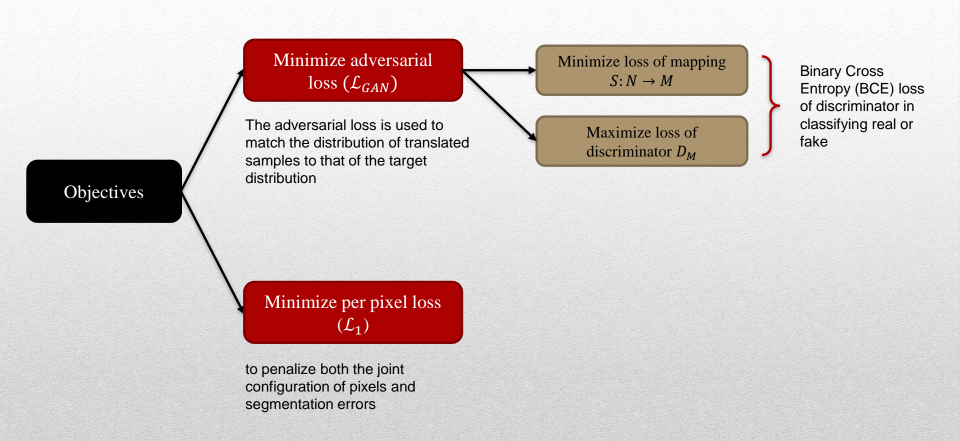
Problem 2: Nuclei segmentation

Architectures: cGAN (conditional GAN)

Normalization: spectral normalization (to prevent mode colapse)

Conditional GANs can learn a **structured context-aware loss** considering a larger receptive field. The cGAN framework learns a mapping S for nuclei segmentation, in which S can adapt H&E nuclei images to their segmentation masks.





Loss of mapping $S: N \to M$

S is penalized if the group configuration of the pixels in the predicted mask is unrealistic, i.e. masks that look like salt-and-pepper noise.

$$\mathcal{L}_{\text{GAN}}(S, D_M) = \mathbb{E}_{m, n \sim p_{\text{data}}(m, n)}[\log D_M(m, n)]$$
$$+ \mathbb{E}_{n \sim p_{\text{data}}(n)}[\log (1 - D_M(m, S(n)))]$$

Pixel loss

$$\mathcal{L}_1(S) = \mathbb{E}_{m,n \sim p_{\text{data}}}(m,n)[||m - S(n)||_1].$$

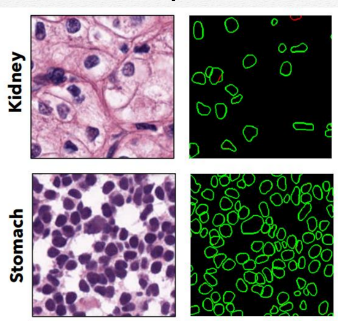
full objective for conditional GAN segmentation

 $\operatorname{arg\ min}_{S} \operatorname{max}_{D_{M}} \mathcal{L}_{\operatorname{GAN}}(S, D_{M}) + \mathcal{L}_{1}(S).$

Training Details

- Pytorch 4.0
- Random jitter (resize image to 286 x 286 → crop back to 256 x 256)
- Optimizer: Adam optimizer
- Number of epoch: 400
- Normalization: spectral normalization
- Learning rate: 0.0002 (for first 200 epochs) → linearly decayed to 0 (for remaining epochs)
- Weight initialized from a Gaussian distribution with a mean 0 and standard deviation 0.02
- Randomly select 64 patch pairs of segmented output and ground truth to be used in the discriminator

Output



Results

Organ	Aggregated Jaccard Index (AJI) ↑				Average Hausdorff distance ↓				F1-Score ↑			
	FCN	U-Net	Mask R-CNN	Proposed	FCN	U-Net	Mask R-CNN	Proposed	FCN	U-Net	Mask R-CNN	Proposed
Breast	0.339	0.485	0.483	0.686	8.17	7.927	6.286	4.761	0.794	0.774	0.749	0.881
Ovary	0.322	0.471	0.513	0.728	8.48	8.142	7.236	4.221	0.675	0.702	0.725	0.876
Esophagus	0.293	0.392	0.433	0.759	8.71	8.349	7.394	3.963	0.667	0.698	0.733	0.891
Liver	0.364	0.422	0.469	0.692	7.93	7.463	6.927	4.173	0.683	0.697	0.741	0.794
Kidney	0.472	0.532	0.542	0.724	7.34	6.858	6.538	4.492	0.775	0.783	0.779	0.829
Prostate	0.281	0.276	0.536	0.731	8.94	8.347	7.912	4.013	0.794	0.802	0.784	0.857
Bladder	0.461	0.491	0.477	0.768	7.11	7.423	7.612	4.273	0.746	0.791	0.769	0.904
Colorectal	0.247	0.218	0.391	0.686	9.27	8.729	7.743	4.712	0.719	0.733	0.691	0.836
Stomach	0.383	0.437	0.629	0.721	7.52	7.314	7.518	4.017	0.866	0.874	0.843	0.922
Overall	0.351	0.414	0.497	0.721	8.163	7.839	7.136	4.291	0.746	0.761	0.757	0.866

The segmentation networks demonstrates a 29.19% improvement in AJI as compared to DIST and 42.98% as compared to CNN-3C. In terms of standard architectures there is a 44.27% improvement over standard Mask R-CNN and 73.19% over a U-Net.

Resources

Mahmood, Faisal dkk. (2018). Deep Adversarial Training for Multi-Organ Nuclei Segmentation in Histopathology Images.

Thank You