Efficient 3D reconstruction of Whole Slide Images in 程序代码 4 编号

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Cutaneous melanoma is an invasive cancer with a worldwide annual death toll of 57,000 (Arnold et al., JAMA Dermatol 2022). In a metastatic state, surgical interventions are not curative and must be coupled with targeted therapy, or immunotherapy. However legitly cappears allows the action of letter tage books can remain poor. The complexity to eradicate melanoma stems from its plasticity; these cancer cells continually adapt to tumor microenvironments, which leads to resistance to treatment. Our primary assumption is that therapeutic resistance relies in part on a series of non-genetic transitions including changes in the metabolic states of these cancer cells Tie 3D patial distribution of proof of the sets that are squires of nutrition and oxygen that drive this metabolic status is an important variable for understanding zoning aspects of this adaptation process. Using Whole Slide Images (WSI) of melanoma tumors from Patient Derived Xenograft (PDX) mouse models, we build 3D vascular models to help predicting and understanding the metabolic states of cancer cells within the tumor. Our 30 reconstruction procline from ADX tumor samples sectioned over 2mm depth and stained with Hematoxylin and cosin (H&E), involves three primary steps, including 2D vessel segmentation using Deep Learning, intensity and affine-based image registration, and 3D reconstruction using interpolation and 3D rendering (allowing a better interaction with biologists, pathologists, and clinicians). The originality of our computer-assisted lineline is its capability to (a) deal with sparse data (i.e., not all tissue sections were readily available), and (b) all pt to a multitude of WH-clased challenges (e.g., epistemic uncertainty, extended processing times due to WSI scale, etc.). We posit both our 3D reconstruction pipeline, quantitative results of the major stages of the process and a detailed illustration of the challenges faced, presenting resolutions to improve the pipeline's efficiency.

Keywords: cutaneous melanoma, whole slide images, 3D reconstruction, vascular reconstruction, personalized medicine

1. INTRODUCTION

Historically a rare cancer, cutaneous melanoma has rapidly evolved to become one of the most fatal forms of cancers today, accounting for approximately 57,000 deaths worldwide annually as of 2020.^{1,2} Early stages of the disease have increased survival rates with surgical interventions. However, treatment of late-stage metastatic melanoma is much more perplexing, relying on targeted therapy, or immunotherapy for an improved line of defense.³ Despite these advanced therapeutic strategies, the clinical outcome of metastatic-affected patients remains poor because of major resistance that appears nearly systematically.

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The infiltration of resident host tissues by the cancer cells can stimulate molecular, cellular, and physical changes within the host tissue, creating a hicromyironner; that is consured to the substitule and proliferation of melanoma. Furthermore, melanoma cancer cells are plastic cells that have the ability to lewire their metabolism to adapt to changing environmental conditions (e.g., poorly vascularized and hypoxic tumor microenvironment, or as a result of the drug exposure), and thus thrive within their complex microenvironment. ⁵

In this study, our prin resistance to treatment stems at least in part from a series of tates. In a previous mathematical modeling work we showed ls, which are sources of nutrition and drug compounds for the non-genetic transitions ar that the spatial heteroger \Box distributed preferentially in highly vascularized regions.⁶ We tumor, may generate zon propose here a pipeline t chitecture of blood vessels distribution in a melanoma tumor using a artificial intelligen buld give here the different steps of the pipeline.... The proposed In to understanding the characteristics of the vascular network pipeline has a multi-funct pusly human-run and manual pathology grading systems that through reconstruction, The pipeline was designed for Whole Slide Images (WSIs) of Hematoxylin are time-consuming and arduous. and Eosin (H&E)-stained pathology section from melanoma tumors derived from Patient-Derived Xenograft (PDX) mouse models. In this study, we elucidate the steps of our pipeline, provide qualitative and quantitative assessment, and discuss WNArchard challengts (notations and objects or the size and depth of information in WSIs), and the solutions we have utilized to improve the efficiency and adaptability of our 3D reconstruction. This 3D vascular reconstruction can be used to explicate how characteristics such as vascular shape, size and bifurcations differ in alternate metabolic states, and thus can be used in predicting treatment efficacy.

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2.1 3D Reconstruction Pipeline

2.1.1 Overall Pipeline

Fig. 1 provides an overall outline of the 3D reconstruction pipeline. Images first underwent preliminary preprocessing steps, including being (A) exported in Tagged Image File (TIF) Format, (B) split into top and bottom tissues, (C) crop-centered, inpainted, and cleaned via the removal of the noisy background. Once images were pre-processed they were (D) registered in a parties of shirt sequentially using an optimized intensity- and affine-based registration method. This was followed by patch-level segmentation (E & F) to produce complete, segmented WSIs (G). All segmented WSIs were rendered and interpolated to produce a final 3D vascular model (H & I).

2.1.2 Image Pre-procesittos://tutorcs.com

The pre-processing stage is designed to normalize the WSIs in order to ease subsequent processes. In essence, the following steps are taken to create WSIs that are more uniform and almost mimic the standardized radiological images, such as those of computed tomography (CT) scans. To create this uniformity, several artifacts and variabilities in WSIs need to be accounted for (Fig. 2), with the most predominant concern being tissue shift. WSI acquisition is littered with epistemic uncertainty (i.e., uncertainty due to incomplete knowledge of a disease or process, or reducible errors, such as subjective, measurement, or human-related errors). This is mostly due to the process being heavily reliant on humans manually carrying out each task of WSI acquisition (i.e., the slicing of the tissue, its addition to a glass slide, etc). To minimize tissue shift, images were crop-centered. Crop-centering in this pipeline includes: establishing an image size with a width of 6000 and height of 5000 (choice of the size based on the size of the baseline and largest tissue section); the identification of the central point of the WSI; and from the identified center, cutting a perimeter based on the pre-set width and height. This process ensures all tissue sections are centrally located while retaining their original tissue size. The maintenance of the original tissue size takes priority. Cutting directly around the perimeter of each tissue would have resulted in small tissue sections appearing erroneously large that would not reflect their true size relative to the whole tissue. Thus, there would have been an inconsistency in the image size that could have impacted rendering and interpolation techniques later in the pipeline (refer to Section 2.1.6). An additional step was required prior to crop-centering. This included the addition of a border of a width of 2000 pixels around each tissue section following its split into top and bottom images. The reasoning for this addition was to account for the disparity in the location of



some tissue sections. Some tissue samples were close to the edge of the slide as shown in Fig. 3. Thus, when crop-centering was attempted the images would reconstruct the images would reconstruct the correct and central placement of 6000×5000 . The addition of a 2000-pixel border granted additional and sufficient space to ensure the correct and central placement of our tissue samples.

Noisy backgrounds were also removed (Fig. 4). Noisy backgrounds in WSIs consist of straining artifacts and pixelation caused by the characteristic for straining artifacts and pixelation caused by the characteristic for straining artifacts and pixelation caused by the characteristic for straining and pixelation their original background and pixelation and pixelation. This was achieved using foreground extraction. The original images were binarized using a simple Otsu thresholding technique, creating a foreground mask. To remove small, erroneous regions, such as extraneous stains around the tissue, the largest contour (i.e., the tissue section) was selected, eliminating any small and unnecessary regions from the binary mask. Using the Bitwise Operation, the white portions of our masks were used to identify the region of interest within the original image, and therefore only foreground information was extracted. The final step was to simply add a white color to the background, resulting in a nice, cleaned tissue section for processing.

Finally, the epidermis was inpainted using a whole-image level trained U-Net model (Fig. 5). Initial registration attempts showed the epidermis hindered the performance of the registration and its presence provided little value for the final vascular reconstruction. The epidermis U-Net model had mostly similar hyperparameters to that of our patch-level blood vessel segmentation model (refer to Section 2.1.4) and included: batch size=32, the ADAM optimizer, learning rate= 3×10^{-5} , epoch=300, and the Binary Cross Entropy Dice loss function (BCE DICE). Given the (almost) consistent nature of the epidermis (i.e., shape, location, patterns, etc.), a smaller epoch size proved to be more than adequate for inpainting. The BCE DICE was chosen based on its ability to better discern segmentation boundaries as compared to other loss functions.



Figure 2: Sample image artifices in acting liberal to. The fattly manual actustion of issue sections in WSI processing, these imaging modalities are highly prone to several artifacts, such as erroneous staining and tissue shifts. There are also pixelations (shown as horizontal bars) that appear around the perimeter of tissue sections.

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Figure 3: Demonstration of proper and improper crop-centering. In some cases, tissue sections would appear at the edge of a slide, resulting in a lack of space to cut the pre-set perimeter of 6000×5000 . This often resulted in images that were not centered, and had smaller image sizes as compared to other tissue sections. To correct this anomaly, an additional step was taken to first add a border size of 2000 pixels before the crop-centering.

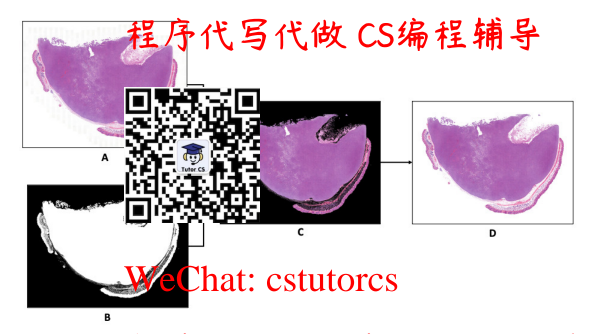


Figure 4: Noisy background Subjact Professional extraction WSI artificial Very removed by using a foreground extraction process. This involved the creation of a binary mask using the Otsu thresholding, selecting the largest contour (i.e., the centralized WSI, eliminating all smaller artifacts that may be surrounding the tissue), then using this binary mask as a reference to which regions of the original image we wish to extract. Finally, a clean white background was re-added, leading to Science tissue lide.



Figure 5: Whole-image level epidermis model trained on U-Net architecture. To remove the epidermis, which provided little value in terms of vascular reconstruction but caused a hindrance in the registration process, a U-Net model was trained to detect and inpaint the epidermis.

2.1.3 Image Registration

Histopathological image registration poses several haller with testian as registration methods limited by their ability to identify distinct points throughout time-series images, while intensity-based methods alone result in a high number of local optima when there is noisy data. In this study, for the registration of the WSI stack, we propose of the symmetric, intensity-based affine registration framework based on a family of symr introduced by Öfverstedt et al.⁸ More specifically, the authors combined intensity-based fuzzy point-to-set bidirectional or fuzzy point-to-set inwards distances into an asymme stance. The limitation of this method is that, when compared \blacksquare uared Differences, Pearson Correlation Coefficient, and Mutual to traditional similarity m Information), this method emory to store auxiliary data structures (e.g., a single, pairedbrking memory). This results in extensive computation time. image registration may re took approximately one week to finalize the registration of our For example, the initial v entire WSI dataset, even h-performance cluster (HPC).

In order to optimize the execution time, we profiled this approach and deduced the following bottlenecks/flaws:

- Sequential image registration takes longer to finish registering all slides for a given tumor because a given $slide_{s+1}$ requires the veristered $slide_s$ (which is lie registration. Sults of $slide_{s-1}$ and $slide_s$) to initiate the subsequent registration. Thus, we can not take advantage of the parallelization offered by HPC. This dependency limits the scalability of the algorithm.
- Registration using the native resolution is unjustifiably costly for such a registration task III used a global approach that estimates one affine transformation applicable to all side pixels. For instance, given a pair of RGB slides with 6000x5000 pixels each, when re-scaling to 90% of the original size, the registration takes 1961.59 sec while yielding a mean absolute error (MAE) of 45.72. However, re-scaling to 10% of the original scale takes only 16.30 sec with a 45.53 error.
- The tumors analyzed in this study are ball-like. Thus, the mid-section slides contain more tissue when compared to the top/bottom slides. This registration approach does not account for this specification since it stretches the slides tissue (affine transformation) to minimize the global error.

We propose a generic optimization scheme for WSI registration (Fig. 6): (i) Chose the down-scaled percentage that does not influence the registration performance, (ii) Register each two consecutive WSI in a separate CPU (job array in case of HPC), (iii) Apply a correction ratio based on the stretched area and original tissue. In order to quantify the added value from the optimization scheme, we registered 220 consecutive H&E WSI. It takes less than 10 minutes compared to one week previously (without the optimization). Note that the same HPC resources were used on both tests.

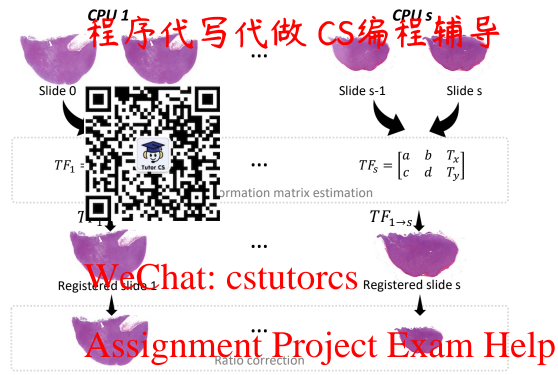


Figure 6: Optimization scheme for whole slide images registration.

2.1.4 2D Blood Vessel Egmand: tutores@163.com

Similar to the epidermis inpainting discussed in Section 2.1.2, our blood segmentation model was trained using the U-Net architecture, with the primary differences being (i) the blood vessel model was trained at a patch-level, with patch sizes of 612×112 , and if the 62×112 finds ances and the variety of shape and locations (even within patches), the training epoch was extended to 1000×122 . Furthermore, data were trained and tested using a 5-fold cross-validation approach, with each fold split into 80% training and 20% test. The final hyper-parameters included: batch size=32, the ADAM optimizer, learning rate= 3×10^{-5} , epoch=1000, and the BCE DICE. As stated earlier, the BCE/DICE was used for its ability to better discern boundaries. This is particularly imperative in the dase of vascular network segmentation. The hypothesis is based on the evaluation of non-genetic transitions that may impact chemotherapeutic response. Thus, capturing the branching and size of blood vessel networks can impact our understanding of nutrient and oxygen diffusion, so we need the most accurate representation of the network.

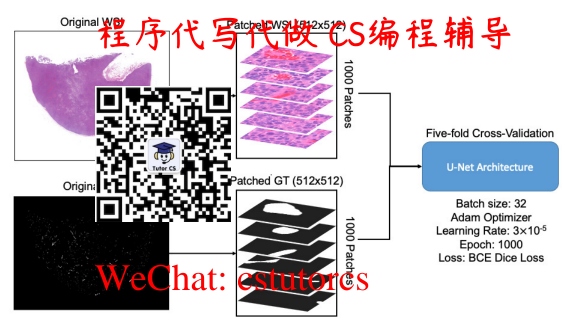


Figure 7: Patch Sestion In the latter of the Control of the Contro

2.1.5 Pipeline Evaluation

The evaluation metrics used to a sess 2D segmentation of the lift of vessels included 176E DICE, MAE, accuracy (ACC; $\frac{TP+TN}{TP+TN+FP+FN}$), Dice coefficient (DICE; $\frac{TP}{2TP+FP+FN}$), Dice Loss (i.e., 1 - DICE; DICE LOSS), the F_1 -score (2 × $\frac{PREC \times REC}{PREC+REC}$), Precision (PREC; $\frac{TP}{TP+FP}$), recall (REC; $\frac{TP}{TP+FN}$), specificity (SPEC; $\frac{TN}{TN+FP}$) and Matthews Correlation Coefficient (MCOR; $\frac{(TP \times TN) - (FP \times FN)}{2(Q \cdot P) \cdot TP+FN}$). Image registration performance was validated using the Structural Similarity index (SIM).

2.1.6 3D Rendering and Interpolation

For a complete 3D reconstruction, a hybrid solution was created using rendering and interpolation, where rendering stacks existing data to create a pyelinibal vascular volume will the interpolation imputes the gaps between the H&E rendered tissues sections. Rendering was achieved with marching cubes. The shape- and distance-based method proposed by Schenk *et al.* was used for interpolation. Their proposed interpolation involves the following:

- First, binary scenes for the *i* and *i+t* images are established. While in their publication these are user-defined contours, in our application the binary scenes involve the segmented and sequentially stacked WSIs.
- Gray-level distance maps are generated for the *i* and *i+t* binary scenes, with distance being relative to the binary scene boundaries. Distances within the boundaries are given positive values, while those outside of the boundaries are negative.
- Gray-level distance maps are interpolated using conventional gray-scale interpolation techniques, such as linear interpolation.
- The interpolated gray-level distance maps are converted back into interpolated binary scenes.

2D-level blood vessel model training was based on the tiles extracted at the highest resolution. For the purposes of 3D rendering and interpolation, the whole slides at their highest resolution would be too large to

export (requiring inordinate amounts of memory) and execute in a timely manner. Therefore in the final phase of the pipeline, whole tissue section, were exported with a townsampling figure of 3. The highest resolution has a magnification of $\times 40$, which translates to 0.25 µm per pixel. At downsampling of 8, we have a magnification level of $\times 5$ and a resolution of 2 µm per pixel. Given the training of the U-Net (described in Section 2.1.4) was based on patch sizes of 512×512 at a resolution of 0.25 µm per pixel, the equivalent patch size and resolution at a downsampling of 8 w sizes. As illustrated in Fig. 8, 512×512 patches with an $\times 40$ magnification (A) and 64 with a size and resolution (C) capture the same contextual and spatial information within WSIs. Sizes and resolution at 512×512 patches with an $\times 5$ magnification (B), reger spatial zone, which would lead to inefficient blood vessel segmentation, as our modern odd over the same context and spatial odd vessels 'close up' within these patches.

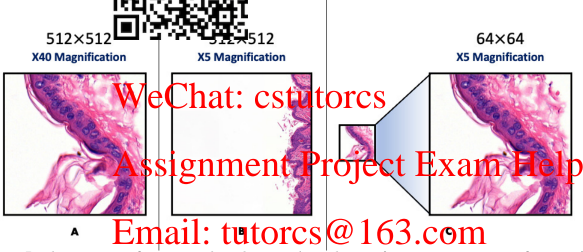


Figure 8: Patch sizes, magnifications, and resolutions. A patch size of 512×512 at a magnification of $\times 40$ captures the same spatial and contextual information as a patch size of 64×64 at a magnification of $\times 5$.

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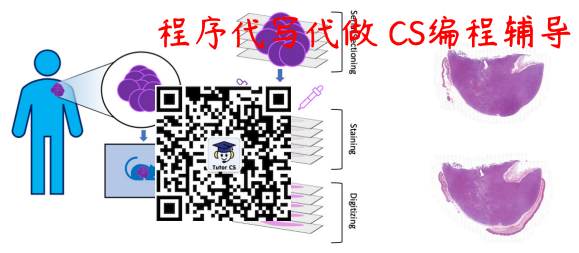
3.1 Data preparation

3.1.1 Raw data https://tutorcs.com

The PDX model is illustrated in Fig. 9a. Cutaneous melanoma tumors were extracted from humans during biopsy, then implanted onto immunodeficient mice for amplification. PDX samples underwent serial sectioning at every 12 μ m over a depth of 2mm and were stained with H&E. Each tissue section had a thickness of 4 μ m. This sectioning protocol resulted in gaps between each tissue section, which were imputed using interpolation methods (refer to Section 2.1.6). Slides were digitized into MIRAX format. Each slide contained two tissue sections (i.e., top and bottom) as shown in Fig. 9b; this method was used to expedite data acquisition time and reduce production costs. A total of 120 slides (240 tissues sections) were available. Blood vessels were annotated by an expert pathologist (co-author CP) using the open-source software QuPath. ¹⁰

3.1.2 Sampling

In preparation for training using deep learning methods, WSIs along with their annotated counterparts were exported as tiles/patches from QuPath. Patches were sized 512x512 (with size selection based on preliminary experimentation with various patch sizes) and were exported at their highest resolution (0.25 µm per pixel). A total of 1000 patches with positive (i.e., presence of blood vessels) and negative (i.e., background/other molecular markers) cases were captured.



(a) Complete PDX model illustration.

(b) Sample H&E slide with two tissue sections.

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Figure 9: Human tumors are engrafted onto immunodeficient mice and grown. Mice-harvested samples then undergo serial sectioning. Slides stained with hematoxylin and eosin (H&E) are digitized into whole slide images (WSIs).

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5. NOVELTY OF WORK

We present an end-to-end approach for reconstructing a vascular 3D model based on WSIs. An original feature of our pipeline is its ability to handle sparse data, given that not all tissue sections were made readily available. This is achieved through rendering and interpolation. Furthermore, the pipeline has been built to accommodate the multitude of challenges faced with processing WSIs. These include: conducting patch-level training and segmentation; creating uniformity in the available data; and improving registration by using inpainting, crop centering, and artefact removal techniques to minimize registration-related errors. Other breakthrough components of our project include the utilization of epistemic uncertainty. This involves understanding human-related or reducible errors, and using statistical techniques to detect and minimize the impact of uncertainty on the pipeline. Our work further includes optimization via parallel processing to improve the efficiency and speed of our reconstruction process.

Table 1: Average training and test results for 2D-level blood vessel segmentation based on U-Net architecture across 5-fold cross-validation

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	Metrics	🔰 Iran	mig -	resung		
		Mean	SD	Mean	SD	
	BCE DICE	0.081	0.019	0.183	0.083	
		.009	0.001	0.027	0.017	
7 2		.992	0.001	0.973	0.016	
3		.886	0.037	0.813	0.086	
W	Tutor CS	-1 .114	0.037	0.188	0.085	
ſŧŧ		.836	0.052	0.767	0.129	
i		.842	0.049	0.847	0.082	
بعا	. z wiane	.832	0.054	0.795	0.112	
	SPEC	0.996	0.001	0.985	0.015	
**	MCOR	0.830	0.053	0.784	0.105	
W	SIP Standard	Deviation	f11f <i>(</i>	rcs		

Table 2: Quantitative assessment of registered images using SSIM.

		~-	Median					
P	0-864 SE = Star	0.0032 Grd Err	nent	Pre	jæc	0.964	kam	Help

SD = Standard Deviation



(a) Overlap between large tissue sections following registration. (b) Overlap between small tissue sections following registration.

Figure 10: Qualitative assessment of registered images. While visually near-perfect, variability in SSIM between larger and smaller tissues are attributable to greater heterogeneity of blood vessels in larger tissues.

6. CONCLUSIONS

In this study, we have proven that AI can be used to develop automated and clinically operational pipelines for WSIs. Using a combination of deep learning, image processing, registration, statistical and parallel processing techniques, we can develop 3D models that are adaptable to WSI challenges and produce results which elucidate vascular networks that could predict chemotherapy efficacy in melanoma patients. This body of work can be readily extended to other biomarkers and cancer types and is not limited in its scope or application.

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