



# SpaCell: integrating tissue morphology and spatial gene expression to predict disease cells

Tan et al

Supplementary information







### Supplementary methods

### Cell Type Clustering

Data preprocessing includes tiling and normalisation to solve two inherent challenges, the small sample size and the technical variation between images. Since the number of images is often small, SpaCell implements a tiling strategy where each spatial spot in Slide-seq data is captured as an image tile with a corresponding column in the count matrix. While most histological images are viewed in isolation without taking into account other images, SpaCell can use spot image tiles from many tissue section images. Images are often different in contrast, colour, and baseline brightness due to various factors such as manufacturers, microscopy settings and slide preparation. The normalisation step in SpaCell removes this variation through colour cast removal and stain normalisation. As histological images are taken against a white background, any images with a coloured background can be assumed to have a colour cast. To remove this colour cast, SpaCell scales the R,G,B channels individually such that the background becomes white. Stain normalisation is implemented in StainTools as described below (Macenko et al., 2009):

$$OD_{flat} = C * S \tag{1}$$

where OD is the optical density image transformed from a H&E stained image in RGB format, S is a stain two colour matrix, and C is a pixel concentration matrix that is used to normalise a target image using the S matrix of that image. By default, the stain matrix is estimated by Vahadane et al. method (Vahadane et al., 2016), or alternatively by Macenko et al. method (Macenko et al., 2009). We found at least 3% improvement in accuracy for models with the normalisation step.

In the cell type identification model, SpaCell uses a pre-trained convolutional neural network (CNN), ResNet50, which makes use of network weights trained from the ImageNet database. SpaCell applies the ResNet50 model (He et al., 2015) to each spot image to find a latent variable vector representing informative features in each spot image. Due to the unbalanced number of features in the gene counts (13,000 genes) and tile feature vector (2048 features) for each spot, SpaCell has function to select 2048 top variable genes to equalise the dimensions of the spot gene counts and spot tile feature vector to 2048. SpaCell also performs min-max scaling method to scale spot gene counts and spot tile feature vector to range of 0 to 1 to minimize bias derived from variation in data ranges between image and count data. SpaCell implements two separate autoencoders for gene count and tile feature vector data to generate two latent spaces with the same dimensions. Those two latent spaces are able to output key features representative of both the original spot image and the spot gene expression, therefore, are concatenated as input for downstream k-means clustering (Fig. 1).

SpaCell implements three loss functions such as the Mean Square Error (MSE) (Equation (2)), Kullback-Leibler (KL) (Equation (3)) and Binary Cross Entropy (BCE) (Equation (4)) to measure the cost of the original input and the constructed output. Loss is minimised by the Adam (Kingma and Ba, 2014) optimiser during the training step (epoch).

$$MSE = \frac{\sum_{i=1}^{N} (y_i - \hat{y}_i)^2}{N}$$
 (2)

$$D_{KL}(p||q) = H_p(q) - H(p)$$

$$= \sum_{i=1}^{N} p(x_i)(\log p(x_i) - \log q(x_i))$$
(3)

$$= \sum_{i=1}^{N} p(x_i) \frac{\log p(x_i)}{\log q(x_i)}$$

$$H_p(q) = -\frac{1}{N} \sum_{i=1}^{N} (x_i \log(p) + (1 - x_i) \log(1 - p))$$
(4)

where i is the index for spot i.  $X_i$  represents 2048 feature vectors from ResNet50 model or a vector of counts for the top 2048 most variable genes measured for  $Spot_i$ . p and q denote the probability distributions for input and constructed output of all N spots.

To quantify performance of the cell type identification model, SpaCell implements a validation tool that utilises pathologist annotations. Pathologist annotations are often low-resolution so SpaCell registers this annotation image to the whole slide image. To achieve this, SpaCell uses a sliding window approach to find the best location and scale for the annotation image, as indicated by a maximum normalised correlation coefficient (Yoo and Han, 2009). In combination with a user-specified annotation colour, SpaCell extracts the annotation contours. These annotations may be open or closed contours. SpaCell preprocesses open contours with a convex hull approach (Barber et al., 1996) to close the contours. Closed contours are filled in to create an annotation mask. By referencing spot coordinates against this pathological annotation mask, SpaCell generates a pathologist label for each spot. SpaCell compares the labels predicted by the cell type identification model to the pathologist labels to generate performance metrics such as accuracy, F-score and ROC curves.

Clustering method were tested on a prostate cancer dataset (Berglund et al., 2018) containing 12 tissue slides from one patient but taken from different prostate locations. Two slides with pathologist annotation were used to test the model performance; P3.3 which represents cancer and non-cancer regions with an open annotation contour and P4.4 which represents inflamed stromal and non-inflammed stromal regions with a closed annotation contour.

### Disease Stage Classification

In the disease stage prediction model, SpaCell uses a two-input deep neural framework to integrate spot image data and spot gene count data (Fig. 1). Spot images feed into an followed by a hidden layer for image feature input and spot gene counts feed into a hidden layer for gene expression input. A merged layer connects these two hidden layers and is followed by a fully connected neural network classifier. This architecture enables the model to learn effective information from both spot image data and spot count data that is relevant to the disease stage. This model uses a Softmax activation function in the last layer to calculate the probability over the C classes for each input Z, defined as:

$$Softmax(Z_i) = \frac{e^{Z_i}}{\sum_{C=1}^{C} e^{Z_C}}$$
 (5)

As this model is designed for multi-class classification problem, a Categorical Cross-Entropy Loss (CCE) (Equation (6)) is implemented where  $t_i$  and  $Softmax(Z_i)$  denote the ground truth and predicted score for class  $C_i$ .

$$CCE = -\sum_{i=1}^{n} t_i log(Softmax(Z_i))$$
 (6)

Dropout layer and 12 regularization for dense layer are used to avoid overfitting. To improve the robustness of the model and assess the risk of overfitting, SpaCell implements stratified 5 fold cross-validation on the training dataset.

Classification method were tested on ALS dataset (Maniatis *et al.*, 2019) consisting of 348 Spinal cord spatial transcriptomics tissue slides in the raw dataset, of which 146 were classified into different ALS disease stages. After removing low quality spots, 31771 spots within 143 WSIs were kept for classification model. Each WSI was labeled as one of the four disease progression categories, including p30, p70, p100 and









p120 representing pre-symptomatic, onset, symptomatic and end-stage respectively. In total, 9267 tiles from 46 WSIs with p30 labelled, 6370 tiles from 33 WSIs with p70 labelled, 6337 tiles from 27 WSIs with p100 labelled and 9797 tiles from 37 WSIs with p120 labelled were used in classification model. 143 ALS mice were randomly split to 100 mice for training and the remaining 43 mice as unseen data for testing such that each disease stage was represented in similar proportions in the test set. Five fold cross validation was applied for the training dataset. After tilling without overlapping, 22179 tiles from 100 mice were used for model training and 9592 tiles from 43 unseen mouse tissue images were used for testing the model performance.

### Supplementary results

### Compare models for cell type classification

For cell type clustering model, we assessed SpaCell's performance in two applications different in biological contexts. In one case, we distinguished cancer cells from non-cancer cells (image P4.2) and in another case we identified stromal cells from the whole tissue (image P3.3), (Supp. Fig 1). Two images were selected because they had pathological annotation, which can be used as a reference for assessing spot predicted values. We successfully mapped the contours from low-resolution pathological annotation images, available as pdf files in the original paper (Berglund et al., 2018), to the original images that are 1000 times larger. The mapping enables us to assess the model performance by accuracy, precision, Fscore, and ROC curve. In both cases, the combined model shows higher performance than using one data type alone, with up to 25% in precision, 14% in accuracy and 38% for F-score. Furthermore, we compared 33 models, with different options for data inputs (combined with two latent spaces, combined with one latent space, single gene-count, single tiled image), data-prepossessing (scaled, top variable genes, and no-scaled), and loss functions (BCE, KL, and MSE ) (Supp. Fig 2). The four heatmap  $\,$ blocks demonstrate the comprehensive comparisons and the superior performance of the combined pixel and gene-count models, with the performance ranked in descending order as: combined with two latent spaces, combined with one latent space, single gene-count model, and single tiled images (Supp. Fig 2). We also found the optimal architecture for the cell type classification model with two latent spaces, BCE loss, and scaled preprocessing. With this optimal model, the performance of the combined image and gene-count data is 8-14 % higher in accuracy, precision, F-score and Area Under the Curve than the models with only one data type (Supp. Fig 2).

# Compare models for disease stage classification

For multiclass classification of the four ALS disease stages, we implemented a stringent design to create a test set completely unseen from the training dataset at both the tile and image levels and performed cross-validation to assess overfitting and model robustness. The design allowed us to assess model performance based on ground-truth labels from known phenotype for each of the above 31,000 tiles representing 143 mice and four disease states. Supp Fig. 3 shows higher performance for the combined model especially for distinguishing the to very similar class, presymtomatic (p30) and onset (p70). The confusion matrix in the supplementary Figure 3 B show that the gene count only model was unable to separate these two classes (P30 and P70). The combined model also performed markedly better compared to the model using image only input.

### Implementation steps

```
# Step 1. Installation
```

```
# 1.1 Build from sources
# Download SapCell from GitHub and install all required packages:
git clone https://github.com/BiomedicalMachineLearning/Spacell.git
cd Spacell
conda env create -f requirements.yml
# To activate environment:
conda activate SpaCell
# To exit environment:
conda deactivate
# 2.2 Build Docker container:
# Download the SpaCell Docker image
docker pull biomedicalmachinelearning/spacell:latest
# Run Docker container
docker run \
-v /path/to/your/data:/home/Spacell/dataset/ \ # mount your local data data
biomedicalmachinelearning/spacell:latest
# Step 2. Setup configurations in config.pv
\ensuremath{\text{\#}} Path to metadata which contains at least sample name column and
# corresponding label column
META_PATH = '../dataset/metadata/mouse_sample_names_sra.tsv'
```

# # Alignment transform matrix # If ST imaging data were aligned, leave it

# Path to spatial transcriptomics imaging data

IMG\_PATH = '../dataset/image/'

CM\_PATH = '../dataset/cm/'

# If ST imaging data were aligned, leave it to None.
# Otherwise, give the path to affine transformation
# matrix generated by st\_spot\_detector
ATM\_PATH = None

# Path to spatial transcriptomics gene counts data

# Path to folder that save the tiles
TILE\_PATH = '../dataset/tile/'

# Path to save intermediate output and final result
DATASET\_PATH = '../dataset/'

# Path to an image which will be used as a template
# for stain normalization
TEMPLATE IMG = '../dataset/image/CN94 D2 HE.ipg'

# Tile size (DO NOT CHANGE)
SIZE = 299, 299

# Color channel (RGB)
N\_CHANNEL = 3









4

```
# Image stain normalization method
                                                        cross validation = False
NORM_METHOD = 'vahadane'
                                                        # Number of splits for cross validation
# Threshold for removing low abundant gene, genes
                                                        k_fold = 2
# expressed in less than THRESHOLD_GENE of total
# number spots will be removed
                                                        # Step 3. Image Preprocessing
THRESHOLD GENE = 0.01
                                                        python image_normalization.py
# Threshold for removing low quality spots, spots
                                                        # Step 4. Count Matrix PreProcessing
# with less than THRESHOLD_SPOT of total genes
                                                        python count_matrix_normalization.py
# expressed will be removed
THRESHOLD\_SPOT = 0.01
                                                        # Step 5. Generate paired image and gene count
                                                        # training dataset
                                                        python dataset_management.py
# Minimum gene count value for counting whether
# expressed or not
MIN\_EXP = 1
                                                        # Step 6. Classification
                                                        python spacell_classification.py
# Specify column name of sample name column,
# label column and condition column (used
                                                        # Step 7. Clustering
# for subset if provided otherwise leave it to None) python spacell_clustering.py -i /path/to/one/image.jpg -l
                                                         /path/to/iamge/tiles/ -c /path/to/count/matrix/
# in metadata file
SAMPLE_COLUMN = 'sample_name'
                                                         -e 100 -k 2 -o /path/to/output/
LABEL_COLUMN = 'age'
CONDITION_COLUMN = 'breed'
                                                        # -e is number of training epochs
                                                        # -k is number of expected clusters
# Subset dataset that all samples have certain
# CONDITION in CONDITION_COLUMN
                                                        # Step 8. Clustering Validation and Quantification
CONDITION = 'B6SJLSOD1-G93A'
                                                        python spacell_validation.py -d /path/to/data
ADDITIONAL_COLUMN = 2 if CONDITION_COLUMN else 1
                                                         -a annotation.png -w wsi.jpeg
                                                         -m affine_tranformation_matrix.txt
# Set random seed for reproducibility
seed = 37
                                                        # -o output_folder
                                                        # -k clustering_predictions.tsv
# Color map for spots color in final clustering plot # -c annotation_colour_range
color_map = ['#ff8aff', '#6fc23f', '#af63ff',
                                                        # -c is annotation colour range thresholds
 '#eaed00', '#f02449', '#00dbeb', '#d19158',
                                                        # -blue_low green_low red_low blue_upper green_upper red_low
 '#9eaada', '#89af7c', '#514036']
                                                        # -t indicates that annotations are not closed paths,
                                                        # so spacell with try to close the paths
# Options for running different models:
                                                        # -f downscale factor if the input whole slide image has
# "combine" : uses both image and gene count data
                                                        # already been downscaled
# "gene_only" : takes gene count data only as input # -s spot size, optional, usually set automatically
# "tile_only" : takes images data only as input
model = ["combine", "gene_only", "tile_only"]
# Number of tiles that will be propagated through
                                                        References
# the model at each step
                                                        Barber, C. B. \it et\,al. (1996). The quickhull algorithm for convex hulls. \it ACM
batch_size = 32
                                                         Trans. Math. Softw., 22(4), 469-483.
                                                        Berglund, E. et al. (2018).
                                                                                 Spatial maps of prostate cancer
# Number of times that all training dataset will be
                                                          transcriptomes reveal an unexplored landscape of heterogeneity. Nature
# passed forward and backward through model
                                                          Communications, 9(1), 2419.
                                                        He, K. et al. (2015). Deep residual learning for image recognition. CoRR,
                                                          abs/1512.03385.
# The ratio for splitting training and test datasets
                                                        Kingma, D. P. and Ba, J. (2014). Adam: A method for stochastic
train_ratio = 0.5
                                                          optimization. cite arxiv:1412.6980Comment: Published as a conference
                                                          paper at the 3rd International Conference for Learning Representations,
# Number of categories in label
                                                          San Diego, 2015.
n_classes = 4
                                                        Macenko, M. et al. (2009). A method for normalizing histology slides
```

for quantitative analysis. In 2009 IEEE International Symposium on

Maniatis, S. et al. (2019). Spatiotemporal dynamics of molecular

pathology in amyotrophic lateral sclerosis. Science, 364(6435), 89–93.

Biomedical Imaging: From Nano to Macro, pages 1107-1110.



# Option for stratified K-fold

# True : run stratified K-fold cross validation

# False : run model without cross validation









Vahadane, A. et al. (2016). Structure-preserving color normalization and sparse stain separation for histological images. IEEE Transactions on Medical Imaging, **35**(8), 1962–1971.

"output" — 2019/11/28 — 2:32 — page 5 — #5

Yoo, J. C. and Han, T. (2009). Fast normalized cross-correlation.  $\it Circuits$ , Systems and Signal Processing, 28, 819–843.



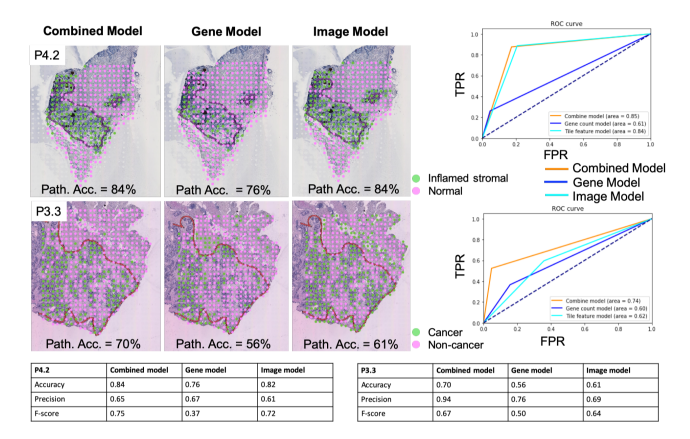






6

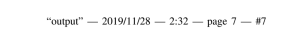
# Supplementary materials



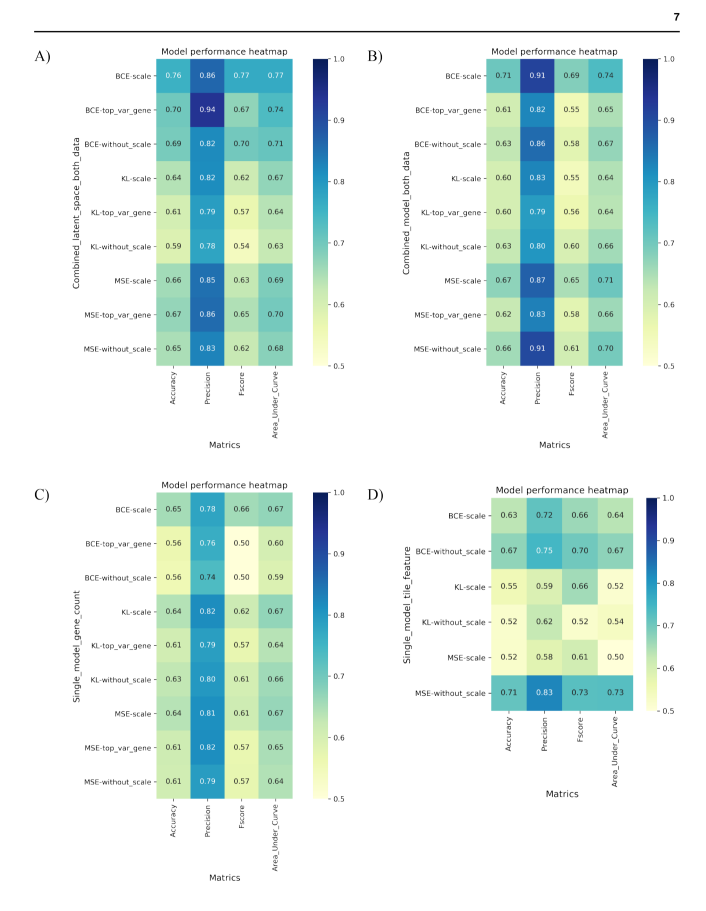
**Supplementary Figure 1.** Cancer and inflamed cell type classification. Comparisons between three models that used either only gene count data or image data or the combination of both as the model training dataset. P4.2, inflamed stromal cell. P3.3, cancer cell types. The red and purple contours denote pathological annotation. All three models implemented same AE architecture with 100 training epochs where losses were calculated by BCE. Top 2048 variable genes were selected in gene model and combined model to balance the weight of gene expression data and tile feature data.











**Supplementary Figure 2.** Comparing cell type classification models. Four data input categories are compared. A) is model using the combination of gene counts and images with two separate autoencoder streams, followed by the concatanation of the two latent spaces. B) is model using the combination of gene counts and images with a single autoencoder used for both pixel and gene count. C) and D) are gene count only model and image only model, respectively.



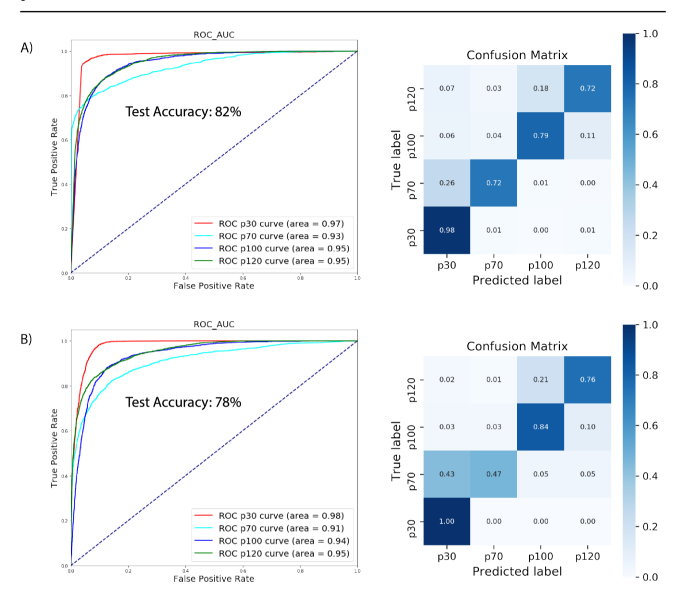












Supplementary Figure 3. Disease stage classification model. Three models are compared. A) and B) are the model using the combination of gene counts and images or the gene count only, respectively.



