User Manual

1 Computer requirements

System: Win10

GPU: Nvidia 1080Ti or better (at least 10G memory), with Nvidia corresponding driver

CUDA: cuda 10 **CPU**: Intel i7 or better

System Memory: 16G or better

2 Supported WSI formats

The software supports the WSI formats supported by the opensource *OpenSlide* library, including *x.svs*, *x.mrxs*, *x.tif*, etc.

WSI resolution: $20 \times$ or $40 \times (0.1 - 0.6 \,\mu\text{m/pixel}, 0.1 - 0.4 \,\mu\text{m/pixel})$ is better)

3 Functions of the software

Recommend the top 10 lesion cells and predict the positive probability of WSIs.

4 Steps of using the software

Step1: Edit the *slidePath* and *savepath* of the *config.ini* file according to your paths as the format of the example

Note: please use absolute path



Step2: Edit the paths of TfModel1, TfModel2 and TfRnn1-TfRnn6 as the format of the example

Note: please use absolute path

//// disconfig_test.ini - 记事本
文件(F) 编辑(E) 格式(O) 查看(V) 帮助(H)
[Config]
[Config]
slidePath=E:\\SoftWareCytologyAI\\TestSlides\\
savePath=E:\\SoftWareCytologyAl\\Results\\test\\
model1Path=E:\\SoftWareCytologyAI\\RNNPredictTool\\models\\m1.pb
model2Path=E:\\SoftWareCytologyAl\\RNNPredictTool\\models\\m2.pb
[TfModel2]
input=input 1:0
output=dense 2/Sigmoid:0,global max pooling2d 1/Max:0
path=E:\\SoftWareCytologyA \\RNNPredictTool\\models\\m2.pb
batti-E.\\SoftwareCytologyAf\\KivivrTedictToof\\fillodeis\\fill2.pb
[TfRnn1]
input=feature input:0
output=output/Sigmoid:0
path=E:\\SoftWareCytologyAI\\RNNPredictTool\\models\\rnn\\10_1.pb
[TfRnn2]
input=feature input:0
output=output/Sigmoid:0
path=E:\\SoftWareCytologyAI\\RNNPredictTool\\models\\rnn\\10_2.pb



Step3: Edit the paths of the *rnnPredict.exe* and the *config.int* in the *test.bat* file

```
■ testbat - 记事本
文件(F) 編輯(E) 格式(O) 章看(V) 報助(H)

Set CUDA_VISIBLE_DEVICES=1
E:\\SoftWareCytologyAl\\\RNNPredictTool\\rnnPredict_exe E:\\SoftWareCytologyAl\\\config_test.ini
E:\\SoftWareCytologyAl\\\RNNPredictTool\\rnnPredict_after.exe E:\\SoftWareCytologyAl\\\config_test.ini
pause
```

Step4: Run the test.bat file

```
C:\Windowe\system32>set CUBA_VISIBLE_DEVICES=1
C:\Windowe\system32>set CUBA_VISIB_DEVICES=1
C:\Win
```

Step5: View the results

For each slide, the results include the cropped image patches of recommended top 10 lesion cells and the predicted positive probability of the slide in *xx.rnnscore*.

In xx.txt below, the positive probability, (x, y) position of recommended cells are provided. Notably, the (x, y) position is relative to the rectangle bounding the non-empty region of the slide, if available by querying slide properties by OpenSlide.

```
文件(F) 编辑(E) 格式(O) 查看(V) 帮助(H)

1.000000,44767,67231

1.000000,47327,69983

1.000000,32671,70431

1.000000,32607,91039

1.000000,30047,70175

1.000000,28895,70559

1.000000,11679,70815

1.000000,79391,69727

1.000000,51231,14815

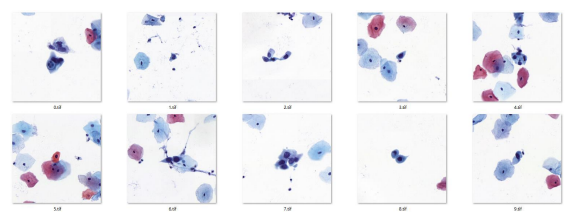
1.000000,15519,38751
```

5 Example

Slide 1: 1.mrxs

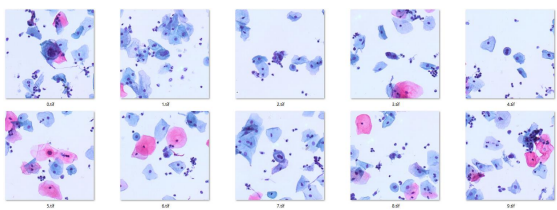
Predicted score of slide: 1.0

Recommended lesion cells: the recommended lesion cells refer to the cells in center region of the patch (about $243 \ \mu m^2$)

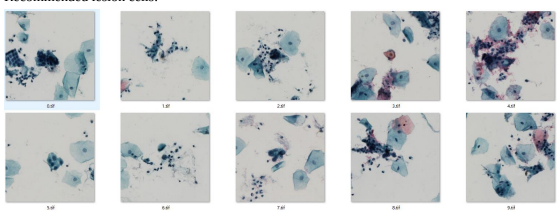


Slide2: 7.svs

Predicted score of slide: 0.995 Recommended lesion cells:



Slide3: 13.srp Predicted score of slide: 0.306 Recommended lesion cells:



Slide4: 19.sdpc

Predicted score of slide: 1.0 Recommended lesion cells:

