

Deep Learning pipeline on histopathology images: detection of prostatic tumor

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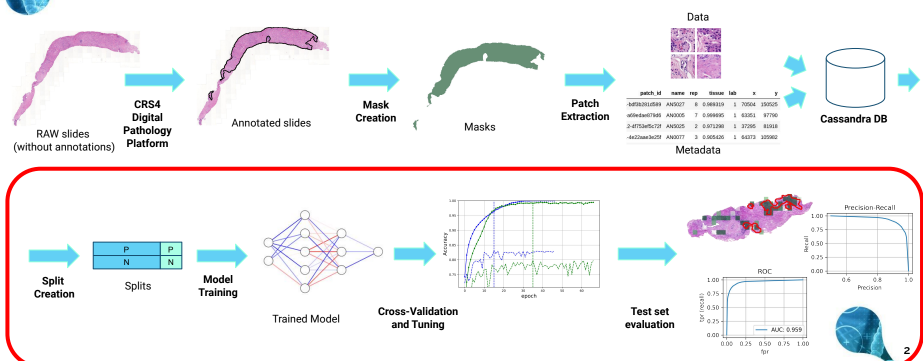






Pipeline: Predictive model









Creation of the splits



Р	Р
N	N
training split	validation split

All **splits** are **balanced**, meaning that the number of tumor patches is roughly the same of the normal ones.

We used the **80%** of the whole dataset for **training data**. The remaining **20%** is used for the **validation set**. The **split_ratios** parameter is set to **[8, 2]**. (**47782** and **11558** patches respectively)

The **test-set** data is organized as a **single split** obtained from different Cassandra tables (**44416** patches)

```
ap = PlainTextAuthProvider(username='xxxx', password='xxxx')
cd = CassandraDataset(ap, ['cassandra-db'])
cd.init listmanager(table= args.ids table,
                        metatable= args.metatable,
                        id col='patch id',
                        num classes=args.num classes,
                        label col=args.label,
                        grouping cols=['sample name']
cd.read rows from db()
min class = np.min(cd. clm. stats.sum(axis=0))
data size = min class*cd. clm.num classes
cd.init datatable(table=args.table)
cd.split_setup(split_ratios=args.split_ratios, max_patches=data_size, augs=[])
cd.save splits(args.out fn)
```





Conv 1-1 (64 x (3, 3)) Conv 1-2 (64 x (3, 3)) Max-pooling (2, 2)

VGG16



 VGG16 is a convolutional neural network with a top classifier made of dense layers

• The number of **trainable parameters** is about **166M** (images **256x256** pixel instead of 224x224)

 Pros: Classical CNN architecture, easy to implement from scratch and although not currently the state of the art, it was one of the best performing architectures in ImageNet Large Scale Visual Recognition Challenge (ILSVRC) challenge 2014

Cons: It is slow to train compared to more recent ANNs. The size of VGG16 weights is about 600MB which potentially takes a lot of disk space (e.g. if you store the model after each epoch) and hence a long time to synchronize in a distributed training setting

EDDL provides a VGG16 neural network pretrained on the Imagenet dataset

 We implemented a VGG16 by using PyEDDL to have more flexibility (e.g.: train from scratch with different initialization methods or pretrained net)

Max-pooling (2, 2)

Conv 3-1 (256 x (3, 3))

Conv 3-2 (256 x (3, 3))

Conv 2-1 (128 x (3, 3))

Conv 2-2 (128 x (3, 3))

Conv 3-2 (256 x (3, 3))

Max-pooling (2, 2)

Conv 4-1 (512 x (3, 3))

Conv 4-2 (512 x (3, 3))

Conv 4-3 (512 x (3, 3))

Max-pooling (2, 2)

Conv 5-1 (512 x (3, 3))

Conv 5-2 (512 x (3, 3))

Conv 5-3 (512 x (3, 3))

Max-pooling (2, 2)

Dense (4096) Dense (4096)

Dense (num classes)







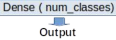
VGG16

in_ = eddl.Input([3, patch_size[0], patch_size[1]])



```
Input
Conv 1-1 (64 x (3, 3))
Conv 1-2 (64 x (3, 3))
  Max-pooling (2, 2)
Conv 2-1 (128 x (3, 3))
Conv 2-2 (128 x (3, 3))
  Max-pooling (2, 2)
Conv 3-1 (256 x (3, 3))
Conv 3-2 (256 x (3, 3))
Conv 3-3 (256 x (3, 3))
  Max-pooling (2, 2)
Conv 4-1 (512 x (3, 3))
Conv 4-2 (512 x (3, 3))
Conv 4-3 (512 x (3, 3))
  Max-pooling (2, 2)
Conv 5-1 (512 x (3, 3))
Conv 5-2 (512 x (3, 3))
Conv 5-3 (512 x (3, 3))
  Max-pooling (2, 2)
    Dense (4096)
```

```
def VGG16(in_, num_classes, seed=1234, init=eddl.HeNormal, 12 reg=None, dropout=None):
    x = in
    x = eddl.ReLu(init(eddl.Conv(x, 64, [3, 3]), seed))
    x = eddl.MaxPool(eddl.ReLu(init(eddl.Conv(x, 64, [3, 3]), seed)), [2, 2], [2, 2])
    x = eddl.ReLu(init(eddl.Conv(x, 128, [3, 3]), seed))
    x = eddl.MaxPool(eddl.ReLu(init(eddl.Conv(x, 128, [3, 3]), seed)), [2, 2], [2, 2])
    x = eddl.ReLu(init(eddl.Conv(x, 256, [3, 3]), seed))
    x = eddl.ReLu(init(eddl.Conv(x, 256, [3, 3]), seed))
    x = eddl.MaxPool(eddl.ReLu(init(eddl.Conv(x, 256, [3, 3]), seed)), [2, 2], [2, 2])
    x = eddl.ReLu(init(eddl.Conv(x, 512, [3, 3]), seed))
    x = eddl.ReLu(init(eddl.Conv(x, 512, [3, 3]), seed))
    x = eddl.MaxPool(eddl.ReLu(init(eddl.Conv(x, 512, [3, 3]), seed)), [2, 2], [2, 2])
    x = eddl.ReLu(init(eddl.Conv(x, 512, [3, 3]), seed))
    x = eddl.ReLu(init(eddl.Conv(x, 512, [3, 3]), seed))
    x = eddl.MaxPool(eddl.ReLu(init(eddl.Conv(x, 512, [3, 3]), seed)), [2, 2], [2, 2])
```



Dense (4096)



Input Conv 1-1 (64 x (3, 3))

VGG16



```
Conv 1-2 (64 x (3, 3))
  Max-pooling (2, 2)
Conv 2-1 (128 x (3, 3))
Conv 2-2 (128 x (3, 3))
  Max-pooling (2, 2)
Conv 3-1 (256 x (3, 3))
Conv 3-2 (256 x (3, 3))
Conv 3-3 (256 x (3, 3))
  Max-pooling (2, 2)
Conv 4-1 (512 x (3, 3))
Conv 4-2 (512 x (3, 3))
Conv 4-3 (512 x (3, 3))
  Max-pooling (2, 2)
Conv 5-1 (512 x (3, 3))
Conv 5-2 (512 x (3, 3))
Conv 5-3 (512 x (3, 3))
```

```
Max-pooling (2, 2)
   Dense (4096)
   Dense (4096)
Dense (num classes)
      Output
```

```
x = eddl.Reshape(x, [-1])
   x = eddl.Dense(x, 4096)
    if dropout:
        x = eddl.Dropout(x, dropout, iw=False)
    if 12 reg:
        x = edd1.L2(x, 12 reg)
    x = init(x, seed)
    x = eddl.ReLu(x)
    x = eddl.Dense(x, 4096)
    if dropout:
        x = eddl.Dropout(x, dropout, iw=False)
    if 12 reg:
        x = eddl.L2(x, 12 reg)
    x = init(x, seed)
    x = eddl.ReLu(x)
    x = eddl.Softmax(eddl.Dense(x,
num classes))
return x
```

Building the model:

- optimizer and its params
- loss function
- metric
- device

```
net = eddl.Model([in ], [out])
eddl.build(
    net,
    eddl.rmsprop(lr),
    ["soft cross entropy"],
    ["categorical accuracy"],
    eddl.CS GPU(gpus, mem=mem) if
gpus else eddl.CS CPU()
```







Training and validation



Training

```
for e in range(args.epochs):
    eddl.set mode(net, 1) # TRMODE = 1, TSMODE = 0
    cd.current split = ∅
    cd.rewind splits(shuffle=True)
    pbar = tqdm(range(num batches tr))
    for b index, b in enumerate(pbar):
        x, y = cd.load batch()
        rescale tensor(x) # to range [-1,1] if tf net
        tx, ty = [x], [y]
        eddl.train batch(net, tx, ty)
    pbar.close()
```

For each epoch:

- Set the network mode to Training. This makes the dropout layers work (if indicated)
- Set the training split by selecting the split with index 0
- Randomly reshuffle samples in each split
- Load samples along with their labels by means of the *load_batch* Cassandra
 Data loader method
- Train the network by using the current batch and labels







Training and validation



Validation

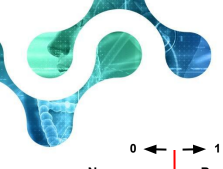
```
eddl.set mode(net, 0) # Set test mode.
cd.current split = 1 ## Set validation split
pbar = tqdm(range(num batches val))
for b index, b in enumerate(pbar):
    x, y = cd.load batch()
    rescale tensor(x) # to range [-1,1] if tf net
    eddl.forward(net, [x])
    output = eddl.getOutput(out)
   result = output.getdata()
   target = v.getdata()
pbar.close()
```

- Set the network mode to Test. This deactivates the dropout layers (if indicated)
- Set the validation split by selecting the split with index 1
- Load samples along with their labels by means of the *load_batch* Cassandra
 Data loader method
- Perform a forward pass and get results from the output of the network





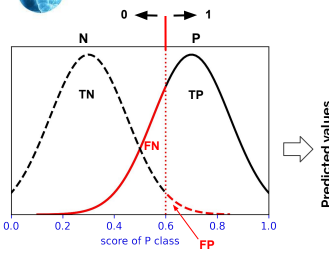




Metrics



Performance Measure: Accuracy, Recall and Precision



Binarization of predictions (th=0.6)

Positive Sample - Pred(P) = $0.75 \rightarrow [1] \rightarrow TP$ Positive Sample - Pred(P) = $0.58 \rightarrow [0] \rightarrow FN$ Negative Sample - Pred(P) = $0.62 \rightarrow [1] \rightarrow FP$ Negative Sample - Pred(P) = $0.22 \rightarrow [0] \rightarrow TN$

	Actual values				
		N	Р		,
Predicted values	N	TN	FN		F
Predicte	Р	FP	TP		F

Actual values

Confusion Matrix

Accuracy =	Correct predictions _	TP+TN
Accuracy -	Total	TD + TN + FD + FN

Recall = Correct pos preds = TP

Positive samples TP + FN

Precision = Correct pos preds = TP

Positive preds TP + FP







Metrics

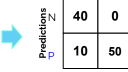


Performance Measure: Accuracy

We use Accuracy to evaluate the validation set during the training process (Recall and Precision are used for cross-validation and test set evaluation)

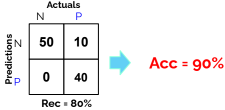
 Pros: Intuitive measure, a model with and accuracy of 90% means that it correctly classifies 90% of observations

- Cons:
 - No information on single class performance.



Actuals



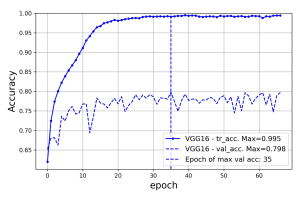


It does not work with imbalanced datasets.

For example, consider we have 98 dogs and 2 cats, and our model provides only dogs as output. The accuracy is 98/100 = 98%. **The model has high accuracy but** is obviously not a good model and it **fails to generalize**.



The Accuracy of the validation set is generally lower than the training set. This is due to overfitting. The model is able to learn the relationship between the training set images and their labels but lacks generalization capabilities.



- Get More (or better) training data: Time consuming and demanding task because a pathologist has to annotate new slides
- Early stopping: Don't wait for the training loss to converge, just get the model with the best validation accuracy.
- L1, L2 Regularization: Add a penalty term to the loss function
- **Dropout:** Some neurons of the selected layers are switched off at training time (remember to reactivate them at evaluation time)
- Data augmentation: Random image transformations (e.g.: flip, rotation)







Stain Normalization

What is H&E stain:

- **Hematoxylin** and **eosin** (**H&E**) stain is used to emphasize cell and tissue structure details to help pathologists in analyzing biopsies of suspected cancer.
- It is the most widely used stain.
- The hematoxylin stains cell nuclei a purplish blue, and eosin stains the extracellular matrix and cytoplasm pink. The other structures take on different shades, hues, and combinations of these colors.

Why Normalize:

- Histological slides from different labs (and even across batches within the same lab) show heterogeneous
 appearance as a result of the different preparation and staining procedures.
- This variability can heavily affect the result of automatic image analysis algorithms.

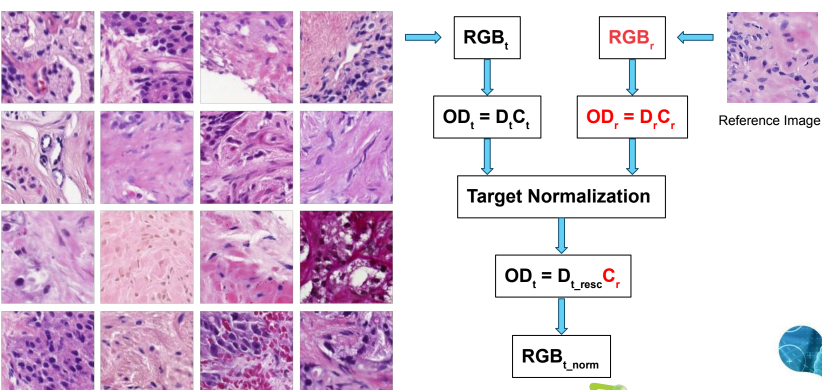
How we normalize:

- We used the Stain separation method proposed in the paper "Structure-preserving color normalization and Sparse Stain Separation for Histological Images" (1).
- We normalized patches within a Cassandra data-table storing the normalized version to a new data-table.
 This allowed us to make a direct comparison by using the same splits but taking patches either with or without normalization



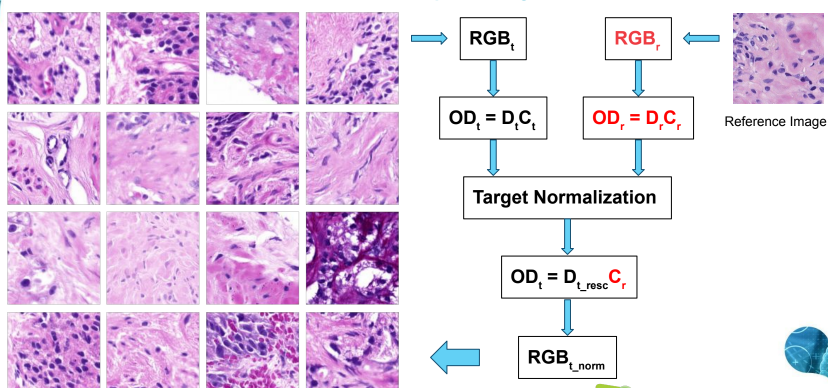


Stain Normalization: Structure-preserving color normalization





Stain Normalization: Structure-preserving color normalization

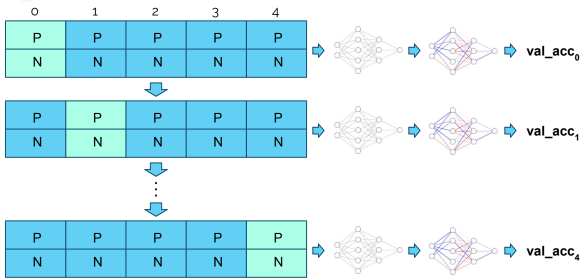






Cross-validation





cross-validation with 5 splits:

- Five splits are created with a split ratio of [1, 1, 1, 1, 1] to have almost equal sized splits.
- The validation set is selected with a round robin policy
- The training set is composed of the remaining splits

Model Accuracy:

It is the average of **val_acc**; with i in [0,4]

training splits

validation split







Cross-validation data loading



Training

for b_index, b in enumerate(pbar): x, y = cd.load_batch_cross(not_splits=val_splits) rescale_tensor(x) ### [0,1] or [-1,1] tx, ty = [x], [y] eddl.train_batch(net, tx, ty) #... pbar.close()

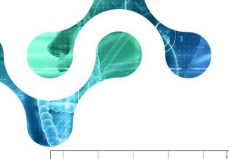
Validation

```
for b_index, b in enumerate(pbar):
    x, y = cd.load_batch_cross(not_splits=train_splits)
    rescale_tensor(x)

eddl.forward(net, [x])
    output = eddl.getOutput(out)
#...
pbar.close()
```

- To make **cross-validation** work and load the right data during the training and validation steps, the **load batch** function must be replaced with the **load batch cross** function.
- **load_batch_cross** takes a list of split indexes as the input argument and load batches from all the splits except those specified in the list
- For example, we have 5 splits and we want to use split 2 for validation. We will provide the
 list [2] to load_batch_cross during the training phase and the list [0,1,3,4] during the
 validation step

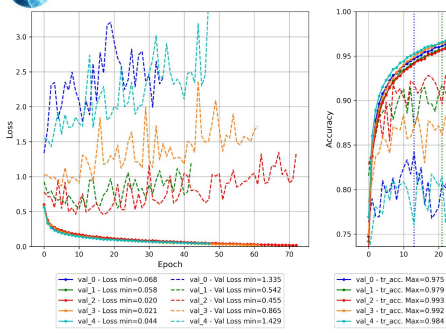


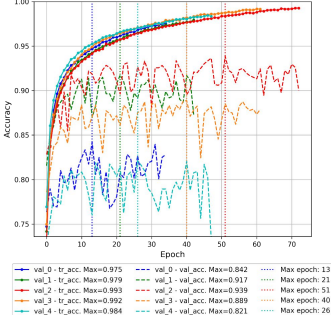


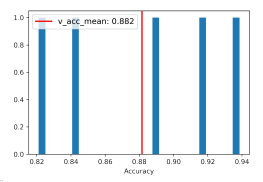
Model evaluation



Cross validation results





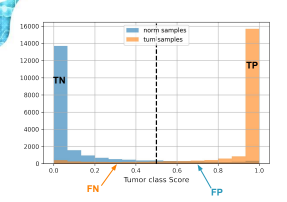


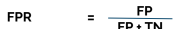


Model evaluation



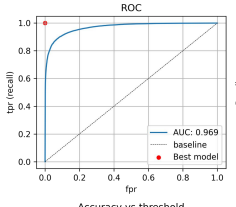
Test Set Results

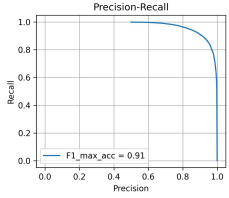


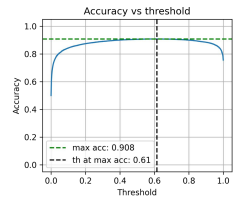


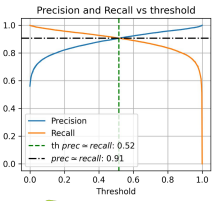
Precision =
$$\frac{TP}{TD + FD}$$

$$Recall(TPR) = \frac{TP}{TP + FN}$$





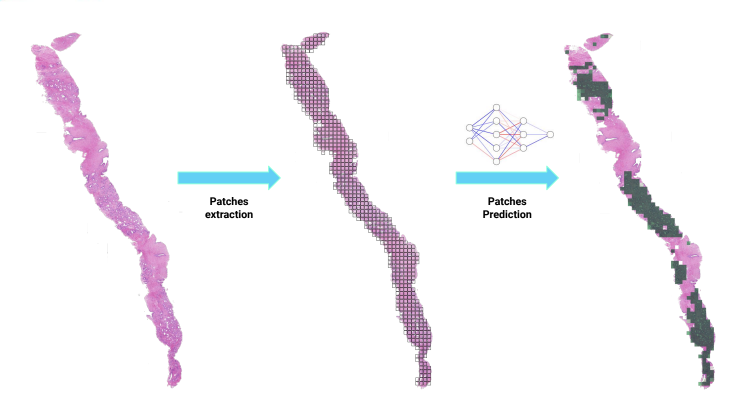








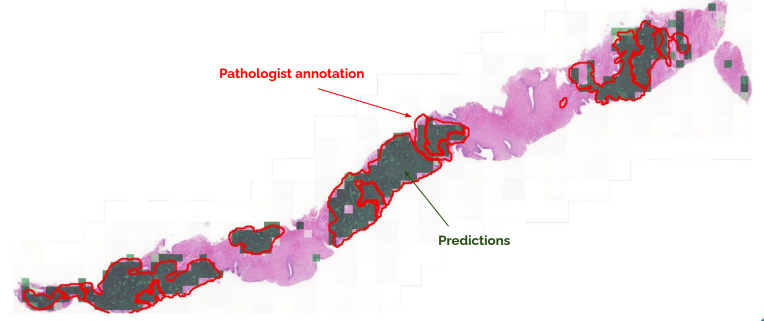








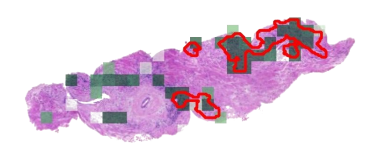




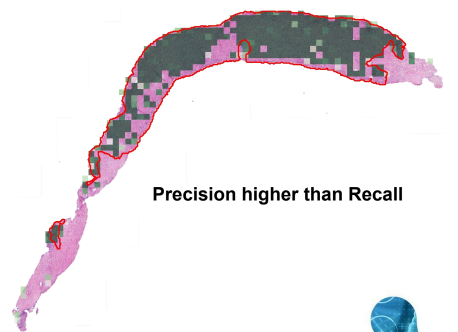








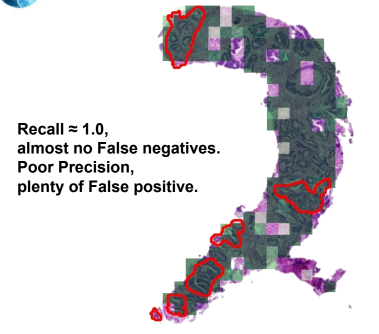
Precision lower than Recall

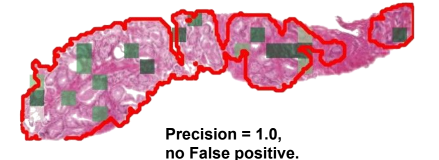












Poor recall,



plenty of False negative.







How to fight mispredictions



- Data Preprocessing:
 - Remove noisy patches
 - More sophisticated normalization algorithms (e.g.: GAN based)
- Try more recent neural network architectures (e.g.: ResNet, Xception, EfficientNet)
- Use different types of data augmentations
- Increase the dataset size by using more uncorrelated images (better to have a lot of cases with a small number of slides each rather than having a small number of cases each with many slides)
- **Increase the number of patches** of the class for which the classifier shows worse performance







Thank you! Q&A



