deepG tutorial

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Introduction

The deepG library can be used for applying deep learning on genomic data. The library supports creating neural network architecture, automation of data preprocessing (data generator), network training, inference and visualizing feature importance (integrated gradients).

Create a model

deepG supports three functions to create a keras model.

create_model_lstm_cnn

The architecture of this model is k * LSTM, m * CNN and n * dense layers, where $k, m \ge 0$ and $n \ge 1$. LSTM (long short-term memory) layers are specifically designed to processes sequential data (order of data is important) by using feedback connections. CNN (convolutional neural network) layers are usually applied to images or audio data but can also be used for natural language processing or genomic sequences. Contrary to vanilla feedforward networks, they are able to process spatial relations in the data.

The user can choose the size of the individual LSTM, CNN and Dense layers and add additional features to each layer; for example the LSTM layer may be bidirectional (runs input in two ways) or stateful (considers dependencies between batches).

The last dense layer has a softmax activation and determines how many targets we want to predict. This output gives a vector of probabilities, i.e. the sum of the vector is 1 and each entry is a probability for one class.

The following implementation creates a model with 3 CNN layer (+ batch normalization and max pooling), 1 LSTM and 1 dense layer.

```
model <- create_model_lstm_cnn(
   maxlen = 500, # number of nucleotides processed in one sample
   layer_lstm = c(32), # number of LSTM cells
   layer_dense = c(4), # number of neurons in last layer (4 targets: A,C,G,T)
   vocabulary.size = 4, # input vocabulary has size 4 (A,C,G,T)
   kernel_size = c(12, 12, 12), # size of individual CNN windows for each layer
   filters = c(32, 64, 64), # number of CNN filters per layer
   pool_size = c(3, 3, 3) # size of max pooling per layer
)</pre>
```

	Model: "model"				
##	Layer (type)	Uutput Shape	Param #		
##	input_1 (InputLayer)	[(None, 500, 4)]	0		
##	conv1d (Conv1D)	(None, 500, 32)	1568		
##	max_pooling1d (MaxPooling1D)	(None, 166, 32)	0		
##	batch_normalization (BatchNormaliza	(None, 166, 32)	128		
##		(None, 166, 64)	24640		
## ## ##	batch_normalization_1 (BatchNormali	(None, 166, 64)	256		
	max_pooling1d_1 (MaxPooling1D)	(None, 55, 64)	0		
##	conv1d_2 (Conv1D)	(None, 55, 64)	49216		
	batch_normalization_2 (BatchNormali		256		
	max_pooling1d_2 (MaxPooling1D)	(None, 18, 64)	0		
##	lstm (LSTM)	(None, 32)	12416		
	dense (Dense)	(None, 4)	132		

```
## ------## Total params: 88,612
## Trainable params: 88,292
## Non-trainable params: 320
## _____
```

The model expects an input of dimensions (NULL (batch size), maxlen, vocabulary size) and a target of dimension (NULL (batch size), number of targets). Maxlen specifies the length of the input sequence.

[1] 3 4

```
colnames(pred) <- c("A", "C", "G", "T")
pred # prediction for initial random weights</pre>
```

create_model_lstm_cnn_target_middle

This architecture is closely related to create_model_lstm_cnn_target with the main difference that the model has two input layers (provided label_input = NULL).

```
model <- create_model_lstm_cnn_target_middle(
   maxlen = 500,
   layer_lstm = c(32),
   layer_dense = c(4),
   vocabulary.size = 4,
   kernel_size = c(12, 12, 12),
   filters = c(32, 64, 64),
   pool_size = c(3, 3, 3)
)</pre>
```

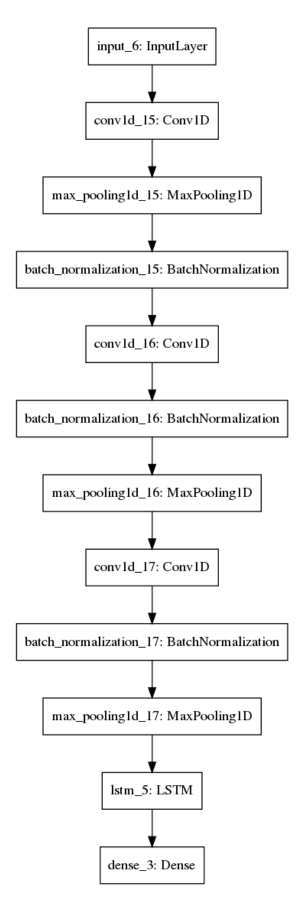


Figure 1: model with 3 CNN and 1 LSTM and 1 Dense layer ^{4}

##					
## ##	conv1d_6 (Conv1D)	(None,	250, 32)	1568	input_3[0][0]
	max_pooling1d_3 (MaxPooli	(None,	83, 32)	0	conv1d_3[0][0]
## ##	max_pooling1d_6 (MaxPooli	(None,	83, 32)	0	conv1d_6[0][0]
## ##	batch_normalization_3 (Ba	(None,	83, 32)	128	max_pooling1d_3[0][0]
	batch_normalization_6 (Ba	(None,	83, 32)	128	max_pooling1d_6[0][0]
	conv1d_4 (Conv1D)	(None,	83, 64)	24640	batch_normalization_3[0][0]
	conv1d_7 (Conv1D)	(None,	83, 64)	24640	batch_normalization_6[0][0]
	max_pooling1d_4 (MaxPooli	(None,	27, 64)	0	conv1d_4[0][0]
	max_pooling1d_7 (MaxPooli	(None,	27, 64)	0	conv1d_7[0][0]
	batch_normalization_4 (Ba	(None,	27, 64)	256	max_pooling1d_4[0][0]
## ##	batch_normalization_7 (Ba	(None,	27, 64)	256	max_pooling1d_7[0][0]
	conv1d_5 (Conv1D)	(None,	27, 64)	49216	batch_normalization_4[0][0]
	conv1d_8 (Conv1D)	(None,	27, 64)	49216	batch_normalization_7[0][0]
## ##	max_pooling1d_5 (MaxPooli	(None,	9, 64)	0	conv1d_5[0][0]
## ##	max_pooling1d_8 (MaxPooli	(None,	9, 64)	0	conv1d_8[0][0]
## ##	batch_normalization_5 (Ba	(None,	9, 64)	256	max_pooling1d_5[0][0]
## ##	batch_normalization_8 (Ba	(None,	9, 64)	256	max_pooling1d_8[0][0]
## ##	lstm_1 (LSTM)	(None,	32)	12416	batch_normalization_5[0][0]
##		(None,			batch_normalization_8[0][0]
## ## ##	concatenate (Concatenate)	(None,	64)		lstm_2[0][0]
	dense_1 (Dense)	(None,	4)	260	concatenate[0][0]
## ## ##	Total params: 177,220 Trainable params: 176,580 Non-trainable params: 640				
			·	·=== -	

This architecture can be used to predict a character in the middle of a sequence. For example sequence: ACCGTGGAA

then the first input should correspond to ACCG, the second input to $\tt GGAA$ and T to the target. This can be used to combine the 2 tasks

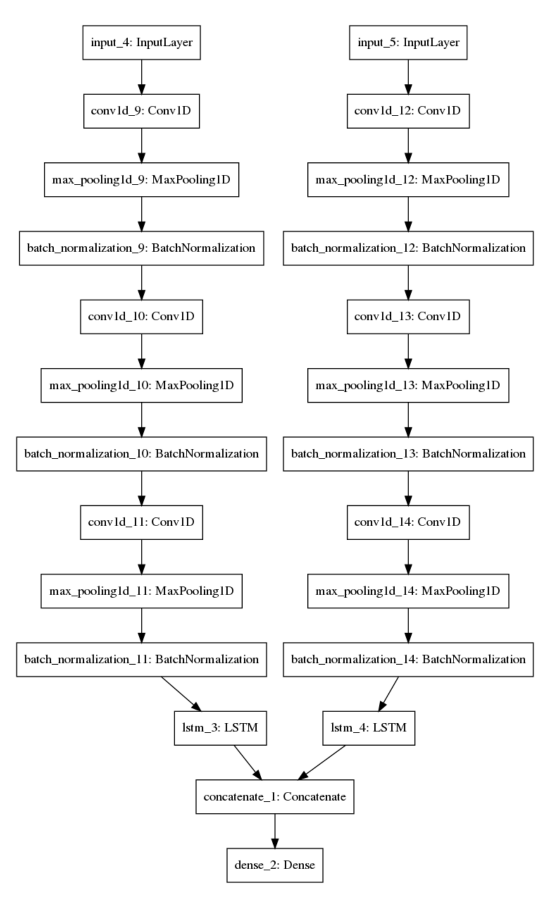


Figure 2: model with two input layers

- 1. predict T given ACCG
- 2. predict T given AAGG (note reversed order of input)

in one model.

create_model_wavenet

This model uses causal dilated convolution layers, which is suitable to handle long sequences. The original paper can be found here

```
## Model
## Model: "model_2"
## Layer (type) Output Shape Param # Connected to
## input_4 (InputLayer)
                 [(None, 500, 4)] 0
 _____
## conv1d_9 (Conv1D)
                  (None, 500, 32) 4096 input_4[0][0]
## r layer (RLayer) [(None, 500, 32), 0
                                 conv1d 9[0][0]
## r_layer_1 (RLayer) [(None, 500, 32), 0
                                  r_layer[0][0]
## r_layer_2 (RLayer) [(None, 500, 32), 0 r_layer_1[0][0]
## add (Add)
                  (None, 500, 32) 0
                                   r_layer[0][1]
##
                                    r_layer_1[0][1]
##
                                    r_layer_2[0][1]
## activation (Activation) (None, 500, 32) 0 add[0][0]
## ______
## conv1d_11 (Conv1D) (None, 500, 16) 512 activation[0][0]
## conv1d_10 (Conv1D) (None, 500, 4) 68 conv1d_11[0][0]
## -----
## Total params: 4,676
## Trainable params: 4,676
## Non-trainable params: 0
```

The model expects an input and output of dimension (batch size, maxlen, vocabulary.size). The target sequence should be equal to input sequence shifted by one position. For example, given a sequence ACCGGTC and maxlen = 6, the input should correspond to ACCGGT and target to CCGGTC.

Training

Preparing the data

Input data must be files in FASTA or FASTQ format and file names must have .fasta or .fastq ending; otherwise files will be ignored. All training and validation data should each be in one folder. deepG uses a data generator to iterate over files in train/validation folder.

Before we train our model, we have to decide what our training objective is. It can be either a language model or label classification.

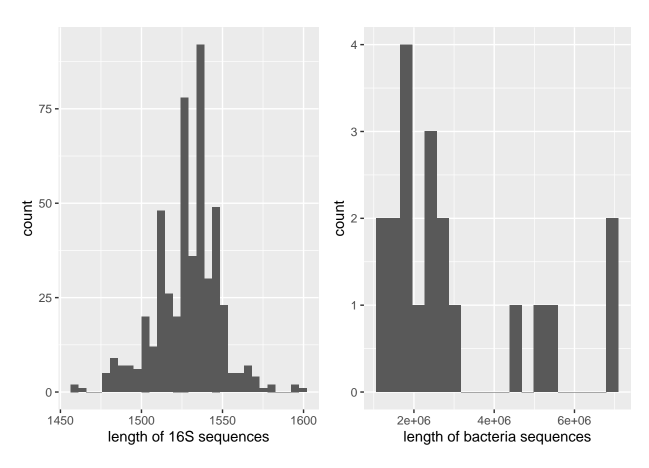
```
path <- "/home/rmreches/tutorial"
path_16S_train <- file.path(path, "16s/train")
path_16S_validation <- file.path(path, "16s/validation")
path_bacteria_train <- file.path(path, "bacteria/train")
path_bacteria_validation <- file.path(path, "bacteria/validation")

checkpoint_path <- file.path(path, "checkpoints")
tensorboard.log <- file.path(path, "tensorboard")
dir_path <- file.path(path, "outputs")
if (!dir.exists(checkpoint_path)) dir.create(checkpoint_path)
if (!dir.exists(tensorboard.log)) dir.create(tensorboard.log)
if (!dir.exists(dir_path)) dir.create(dir_path)</pre>
```

Our data set for this tutorial consists of 16S sequences and bacterial genomes.

```
cat("number of files in 16S train:", length(list.files(path_16S_train)), "\n")
## number of files in 16S train: 498
cat("number of files in 16S validation:", length(list.files(path_16S_validation)), "\n")
## number of files in 16S validation: 100
cat("number of files in bacteria train:", length(list.files(path_bacteria_train)), "\n")
## number of files in bacteria train: 20
cat("number of files in bacteria validation:", length(list.files(path_bacteria_validation)), "\n")
## number of files in bacteria validation: 5
# print first 16S file
print(microseq::readFasta(list.files(path_16S_train, full.names = TRUE)[1]))
## # A tibble: 1 x 2
##
    Header
                                Sequence
                                <chr>
## 1 16S_rRNA::AM295250.1:1217~ TATGGAGAGTTTGATCCTGGCTCAGGATGAACGCTGGCGGCGTGCCTAAT~
```

```
seq_length_16_train <- vector("integer", length(path_16S_train))</pre>
i <- 1
for (file_name in list.files(path_16S_train, full.names = TRUE)) {
  fasta.file <- microseq::readFasta(file_name)</pre>
  seq_length_16_train[i] <- nchar(fasta.file$Sequence)</pre>
  i <- i + 1
}
df_16S <- as.data.frame(seq_length_16_train)</pre>
seq_length_bacteria_train <- vector("integer", length(path_bacteria_train))</pre>
i <- 1
for (file_name in list.files(path_bacteria_train, full.names = TRUE)) {
  fasta.file <- microseg::readFasta(file name)</pre>
  seq_length_bacteria_train[i] <- nchar(fasta.file$Sequence)</pre>
  i <- i + 1
}
df_bact <- as.data.frame(seq_length_bacteria_train)</pre>
p1 <- ggplot(df_16S, aes(x=seq_length_16_train)) +
  xlab("length of 16S sequences") +
  geom_histogram(bins=30)
p2 <- ggplot(df_bact, aes(x=seq_length_bacteria_train)) +</pre>
  xlab("length of bacteria sequences") +
  geom_histogram(bins=20)
ggpubr::ggarrange(p1, p2, ncol = 2, nrow = 1)
```



Language model

With language model, we mean a model that predicts a character in a sequence. The target can be at the end of the sequence, for example

ACGTCAG

or in the middle

ACGTCAG

Language model for 16S (predict next character)

Say we want to predict the next character in a sequence given the last 500 characters and our text consists of the letters A,C,G,T. First we have to create a model. We may use a model with 1 LSTM, 3 CNN and 1 dense layer for predictions.

```
model <- create_model_lstm_cnn(
  maxlen = 500,
  layer_lstm = c(32),
  layer_dense = c(4),
  vocabulary.size = 4,
  kernel_size = c(12, 12, 12),
  filters = c(32, 64, 128),
  pool_size = c(3, 3, 3),
  learning.rate = 0.001
)</pre>
```

Next we have to specify the location of our training and validation data and the output format of the data generator. We randomly select 95% from each file so the generator does not take samples from the same position repeatedly.

```
trainNetwork(train_type = "lm", # train a language model
             model = model.
             path = path_16S_train, # location of trainig data
             path.val = path_16S_validation, # location of validation data
             checkpoint_path = checkpoint_path,
             tensorboard.log = tensorboard.log,
             validation.split = 0.2, # use 20% of train size for validation
             run.name = "lm_16S_target_right",
             batch.size = 256,
             epochs = 5,
             steps.per.epoch = 15, # 1 epoch = 15 batches
             step = 100, # take a sample every 100 steps
             output = list(none = FALSE,
                           checkpoints = TRUE,
                           tensorboard = TRUE,
                           log = FALSE,
                           serialize_model = FALSE,
                           full model = FALSE
             ),
             tb images = TRUE,
             output_format = "target_right", # predict target at end of sequence
             proportion_per_file = c(0.95)
```

```
# randomly select 95% of file
## Trained on 15 samples (batch_size=NULL, epochs=5)
## Final epoch (plot to see history):
##
       loss: 1.118
        acc: 0.5359
##
##
         f1: Inf
## val_loss: 1.12
   val_acc: 0.5182
     val_f1: Inf
##
##
         lr: 0.001
# tensorflow::tensorboard(tensorboard.log)
```

Predict character in middle of sequence

If we want to predict a character in the middle of a sequence and use LSTM layers, we should split our input into two layers. One layer handles the sequence before and one the input after the target. If, for example

sequence: ACCGTGGAA

then first input corresponds to ACCG and second to AAGG. We may create a model with two input layers using the create_model_cnn_lstm_target_middle

```
model <- create_model_lstm_cnn_target_middle(
    maxlen = 500,
    layer_lstm = c(32),
    layer_dense = c(4),
    vocabulary.size = 4,
    kernel_size = c(12, 12, 12),
    filters = c(32, 64, 128),
    pool_size = c(3, 3, 3),
    learning.rate = 0.001
)</pre>
```

The trainNetwork call is identical to the previous model, except we have to change the output format of the generator by setting output_format = "target_middle_lstm". This reverses the order of the sequence after the target.

```
trainNetwork(train_type = "lm", # train a language model
    model = model,
    path = path_16S_train, # location of trainig data
    path.val = path_16S_validation, # location of validation data
    checkpoint_path = checkpoint_path,
    tensorboard.log = tensorboard.log,
    validation.split = 0.2, # use 20% of train size for validation
    run.name = "lm_16S_target_middle_lstm",
    batch.size = 256,
    epochs = 5,
    steps.per.epoch = 15, # 1 epoch = 15 batches
    step = 100, # take a sample every 100 steps
```

```
## loss: 0.8906

## acc: 0.6727

## f1: Inf

## val_loss: 0.9077

## val_acc: 0.6406

## val_f1: Inf

## lr: 0.001
```

Label classification

With label classification, we describe the task of mapping a label to a sequence. For example: given the sequence ACGACCG, does the sequence belong to a viral or bacterial genome?

deepG offers two options to map a label to a sequence

- 1. the label gets read from the fasta header
- 2. files from every class are in separate folders

```
>165

| Telectititide of the content of the content
```

Figure 3: Example: read class label from header

Label by folder

We put all data from each class into separate folders. In the following example, we want to classify if a sequence belongs to 16s or bacterial genome. We have to put all 16S/bacteria files into their own folder. In this case the path and path.val arguments should be vectors, where each entry is the path to one class.

```
model <- create_model_lstm_cnn(
    maxlen = 500,
    layer_lstm = NULL,
    layer_dense = c(2), # predict 2 classes
    vocabulary.size = 4,
    kernel_size = c(12, 12, 12),
    filters = c(32, 64, 128),
    pool_size = c(3, 3, 3),
    learning.rate = 0.001
)</pre>
```

```
trainNetwork(train_type = "label_folder", # reading label from folder
             model = model,
             path = c(path_16S_train, # note that path has two entries
                      path_bacteria_train),
             path.val = c(path_16S_validation,
                          path_bacteria_validation),
             checkpoint_path = checkpoint_path,
             tensorboard.log = tensorboard.log,
             validation.split = 0.2,
             run.name = "16S_vs_bacteria",
             batch.size = 256, # half of batch is 16S and other half bacteria data
             epochs = 5,
             save_best_only = FALSE,
             steps.per.epoch = 15,
             step = c(100, 500), # smaller step size for 16S
             labelVocabulary = c("16s", "bacteria"), # label names
             output = list(none = FALSE,
                           checkpoints = TRUE,
                           tensorboard = TRUE,
                           log = FALSE,
                           serialize_model = FALSE,
                           full_model = FALSE
             ),
             tb_images = TRUE,
             proportion_per_file = c(0.95, 0.05)
             # randomly select 95% of 16S and 5% of bacteria files,
             # since bacteria files are much larger
```

```
## Trained on 15 samples (batch_size=NULL, epochs=5)
## Final epoch (plot to see history):
## loss: 0.01451
## acc: 0.9958
## f1: 0.9958
## val_loss: 0.001222
## val_acc: 1
## val_f1: 1
## lr: 0.001
```

Checkpoints

We can save the architecture and weights of a model after every epoch using checkpoints. The checkpoints get stored in h5 format. The file names contain the corresponding epoch, loss and accuracy. For example, we can display the checkpoints from binary classification model for 16S/bacteria.

```
[1] "Ep.001-val loss0.06-val acc0.984.hdf5"
##
    [2] "Ep.001-val_loss0.09-val_acc0.962.hdf5"
##
       "Ep.001-val loss0.10-val acc0.966.hdf5"
##
    [4]
        "Ep.001-val_loss0.42-val_acc0.854.hdf5"
##
        "Ep.002-val loss0.03-val acc0.991.hdf5"
        "Ep.002-val_loss0.04-val_acc0.987.hdf5"
##
    [6]
##
    [7]
        "Ep.002-val_loss0.06-val_acc0.982.hdf5"
##
    [8]
        "Ep.002-val_loss0.17-val_acc0.947.hdf5"
##
    [9]
        "Ep.003-val_loss0.00-val_acc0.999.hdf5"
        "Ep.003-val_loss0.00-val_acc1.000.hdf5"
##
   [10]
##
   [11]
        "Ep.003-val_loss0.07-val_acc0.986.hdf5"
##
   [12]
        "Ep.004-val_loss0.00-val_acc0.999.hdf5"
##
  [13]
        "Ep.004-val_loss0.01-val_acc0.992.hdf5"
##
   [14]
        "Ep.004-val_loss0.01-val_acc0.996.hdf5"
   [15]
        "Ep.004-val_loss0.03-val_acc0.988.hdf5"
##
   [16]
        "Ep.005-val loss0.00-val acc1.000.hdf5"
##
   [17]
        "Ep.005-val_loss0.01-val_acc0.994.hdf5"
##
   [18]
        "Ep.005-val_loss0.20-val_acc0.940.hdf5"
  [19]
        "Ep.006-val_loss0.00-val_acc1.000.hdf5"
##
  [20]
##
        "Ep.006-val loss0.01-val acc0.996.hdf5"
  [21]
        "Ep.007-val loss0.00-val acc0.999.hdf5"
##
##
   [22]
        "Ep.007-val loss0.02-val acc0.991.hdf5"
##
   [23]
        "Ep.008-val_loss0.00-val_acc1.000.hdf5"
   [24]
        "Ep.008-val_loss0.04-val_acc0.992.hdf5"
   [25]
        "Ep.009-val_loss0.00-val_acc0.999.hdf5"
##
##
   [26]
        "Ep.009-val_loss0.00-val_acc1.000.hdf5"
   [27]
        "Ep.010-val_loss0.00-val_acc0.998.hdf5"
##
##
   [28]
        "Ep.010-val_loss0.00-val_acc1.000.hdf5"
##
   [29]
        "Ep.011-val_loss0.00-val_acc1.000.hdf5"
   [30]
        "Ep.011-val_loss0.01-val_acc0.998.hdf5"
##
   [31]
       "Ep.012-val_loss0.00-val_acc0.997.hdf5"
   [32] "Ep.012-val_loss0.00-val_acc0.999.hdf5"
##
   [33] "Ep.013-val loss0.00-val acc0.999.hdf5"
##
   [34]
       "Ep.014-val_loss0.01-val_acc0.998.hdf5"
   [35] "Ep.015-val_loss0.00-val_acc1.000.hdf5"
```

After training, we can load a trained model and continue training or use the model for predictions/inference. Let's create a model with random weights identical to our 16S/bacteria classifier and make some predictions.

```
model <- create_model_lstm_cnn(
  maxlen = 500,
  layer_lstm = NULL,
  layer_dense = c(2),</pre>
```

```
vocabulary.size = 4,
  kernel_size = c(12, 12, 12),
  filters = c(32, 64, 128),
 pool_size = c(3, 3, 3),
 learning.rate = 0.001
# evaluate 1000 samples, 500 from each class
eval_model <- evaluateFasta(fasta.path = c(path_16S_validation,</pre>
                                             path_bacteria_validation),
                             model = model,
                             batch.size = 100,
                             step = 100,
                             label_vocabulary = c("16s", "bacteria"),
                             numberOfBatches = 10,
                             mode = "label_folder",
                             verbose = FALSE)
eval_model[["accuracy"]]
## [1] 0.5
eval_model[["confusion_matrix"]]
##
             Truth
## Prediction 16s bacteria
              500
                        500
##
     16s
##
                          0
     bacteria
As expected, the performance is not better than random guessing. Let's repeat evaluation but load the
weights of our pretrained model
weight_path <- cp[length(cp)]</pre>
model <- keras::load_model_weights_hdf5(model, weight_path)</pre>
eval_model <- evaluateFasta(fasta.path = c(path_16S_validation,</pre>
                                             path_bacteria_validation),
                             model = model,
                             batch.size = 100,
                             step = 100,
                             label vocabulary = c("16s", "bacteria"),
                             numberOfBatches = 10,
                             mode = "label_folder",
                             verbose = FALSE)
eval_model[["accuracy"]]
## [1] 1
eval_model[["confusion_matrix"]]
##
             Truth
## Prediction 16s bacteria
              500
                      500
##
     bacteria 0
```

Inference

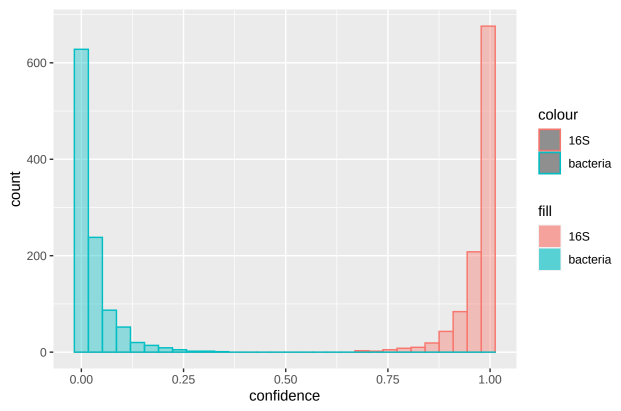
Once we have trained a model, we may use the model to get the activations of a certain layer and write the states to an h5 file. First, we apply our model to a file from our 16S validation set.

```
model <- keras::load_model_hdf5(weight_path, compile = FALSE)</pre>
model <- keras::load_model_weights_hdf5(model, weight_path)</pre>
maxlen <- model$input$shape[[2]]</pre>
num_layers <- length(model$get_config()$layers)</pre>
layer_name <- model$get_config()$layers[[num_layers]]$name</pre>
cat("get output at layer", layer_name)
## get output at layer dense_4
fasta.path <- list.files(path_16S_validation, # make predictions for 16S file
                           full.names = TRUE)[1]
fasta.file <- microseq::readFasta(fasta.path)</pre>
head(fasta.file)
## # A tibble: 1 x 2
##
    Header
                                   Sequence
     <chr>
                                   <chr>
## 1 16S rRNA::CP015410.2:15033~ TATGAGAGTTTGATCCTGGCTCAGGACGAACGCTGGCGGCGTGCCTAAT~
sequence <- fasta.file$Sequence[1]</pre>
filename <- file.path(dir_path, "states.h5")</pre>
if (!file.exists(filename)) {
  writeStates(
    model = model,
    layer_name = layer_name,
    sequence = sequence,
    round_digits = 4,
    filename = filename,
    batch.size = 10,
    mode = "lm")
}
We can access the h5 file as follows
states <- readRowsFromH5(h5_path = filename, complete = TRUE)</pre>
## states matrix has 1058 rows and 2 columns
colnames(states) <- c("conf_16S", "conf_bacteria")</pre>
rownames(states) <- paste0("sample_", 1:nrow(states))</pre>
head(states)
```

```
##
            conf_16S conf_bacteria
## sample_1
               0.9960
                              0.0040
## sample 2
               0.9919
                              0.0081
               0.9951
                              0.0049
## sample_3
## sample_4
               0.9956
                              0.0044
## sample 5
                              0.0047
               0.9953
## sample 6
               0.9961
                              0.0039
```

```
ggplot(as.data.frame(states)) +
  geom_histogram(aes(conf_16S, color = "16S", fill = "16S"), alpha = 0.4) +
  geom_histogram(aes(conf_bacteria, color = "bacteria", fill = "bacteria"), alpha = 0.4) +
  xlab("confidence") + ggtitle("model confidence for 16S file")
```

model confidence for 16S file



The matrix shows the models confidence in its predictions. Every row corresponds to one sample. If the value in the 16s column is > 0.500, the model will classify the sample as 16s.

Detect 16S region

In the following example we use the binary model trained to classify 16S/bacteria to predict the location of 16S sequences (similar to https://github.com/tseemann/barrnap). We use the same model architecture as before, but load the weights of a model that was trained on more data and for a longer time. We iterate over a new bacteria file (not present in train or validation data) and make predictions every 100 steps

```
model = model,
    layer_name = layer_name,
    sequence = sequence,
    round_digits = 4,
    filename = filename,
    batch.size = 500,
    step = 100)
}
states <- readRowsFromH5(h5_path = filename, complete = TRUE, getTargetPositions = TRUE)</pre>
## states matrix has 30252 rows and 2 columns
pred <- states[[1]]</pre>
position <- states[[2]] - 1</pre>
df <- cbind(pred, position) %>% as.data.frame()
colnames(df) <- c("conf_16S", "conf_bacteria", "seq_end")</pre>
head(df)
##
     conf_16S conf_bacteria seq_end
## 1
      0.0018
                      0.9982
## 2 0.0027
                                 600
                      0.9973
## 3 0.0029
                      0.9971
                                 700
## 4 0.0028
                      0.9972
                                 800
## 5
                      0.9988
      0.0012
                                 900
## 6 0.0014
                      0.9986
                                1000
index 16S pred \leftarrow df[ , 1] > 0.5
df_16S <- df[index_16S_pred, ]</pre>
head(df_16S)
##
         conf_16S conf_bacteria seq_end
## 20241
           0.6726
                          0.3274 2024500
                          0.3767 2037200
## 20368
           0.6233
## 21125 0.6095
                         0.3905 2112900
## 21897
           0.9963
                          0.0037 2190100
## 21898
           0.9946
                          0.0054 2190200
## 21899
           0.9627
                          0.0373 2190300
Next, we search for the true rRNA region in the corresponding gff file to validate our models predictions.
fasta.path <- file.path(path, "E_faecalis.fasta")</pre>
gff.file <- file.path(path, "E_faecalis.gff")</pre>
gff.data <- rtracklayer::readGFF(gff.file, version = 0,</pre>
                                   columns = NULL, tags = NULL, filter = NULL, nrows = -1,
                                  raw_data = FALSE)
rRNA_index <- stringr::str_detect(gff.data$product, "^16S ribosomal") &
```

if (!file.exists(filename)) {

(gff.data\$strand == "+")

start <- gff.data[rRNA_index, "start"]
end <- gff.data[rRNA index, "end"]</pre>

cat("start positions of 16S sequences:", start, "\n")

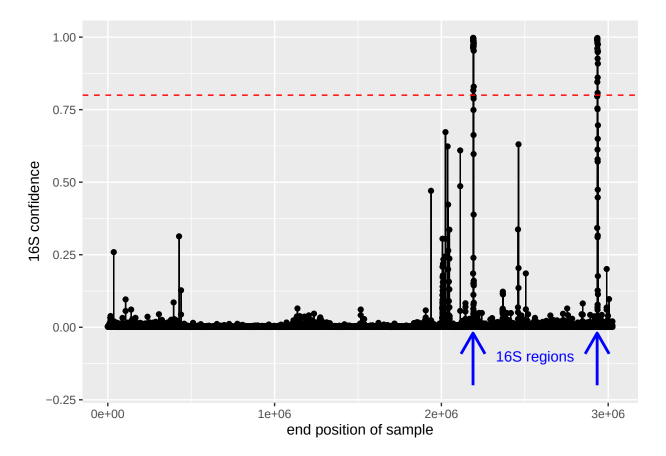
writeStates(

```
## start positions of 16S sequences: 2189670 2933745
```

```
cat("end positions of 16S sequences:", end, "\n")
```

end positions of 16S sequences: 2191227 2935302

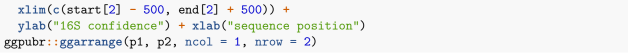
We can now plot the models confidence in 16S over the whole genome.

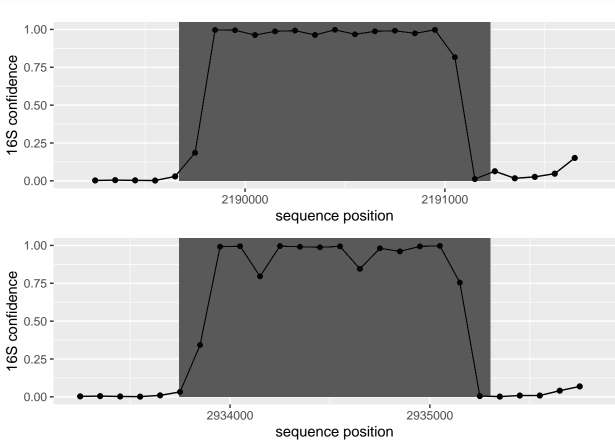


Next we may zoom into areas with high 16S confidence, the true 16S regions are shaded grey.

```
p1 <- ggplot(df, aes(x = seq_end - 250, y = conf_16S)) + geom_point() + geom_line() +
    geom_rect(aes(xmin=start[1], xmax=end[1], ymin=0, ymax=Inf), alpha = 0.01) +
    xlim(c(start[1] - 500, end[1] + 500)) +
    ylab("16S confidence") + xlab("sequence position")

p2 <- ggplot(df, aes(x = seq_end - 250, y = conf_16S)) + geom_point() + geom_line() +
    geom_rect(aes(xmin=start[2], xmax=end[2], ymin=0, ymax=Inf), alpha = 0.01) +</pre>
```

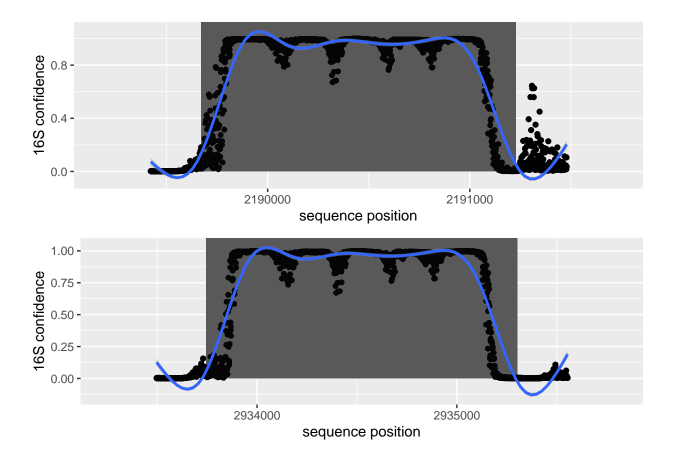




Finally, let's decrease our step size parameter to 1 and make a predictions for every possible sample around the true 16S regions (add 500 nucleotides to start/end)

```
fasta.file <- microseq::readFasta(fasta.path)</pre>
buffer <- 500
sequence_16S_1 <- fasta.file$Sequence[1] %>%
  substr(start = start[1] - buffer, stop = end[1] + buffer)
sequence_16S_2 <- fasta.file$Sequence[1] %>%
  substr(start = start[2] - buffer, stop = end[2] + buffer)
filename_16S_1 <- file.path(dir_path, "states_16S_1.h5")</pre>
filename_16S_2 <- file.path(dir_path, "states_16S_2.h5")
if (!file.exists(filename_16S_1)) {
  writeStates(
    model = model,
    layer_name = layer_name,
    sequence = sequence_16S_1,
    round_digits = 4,
    filename = filename_16S_1,
    batch.size = 500,
    step = 1)
```

```
}
if (!file.exists(filename_16S_2)) {
  writeStates(
    model = model,
    layer_name = layer_name,
    sequence = sequence_16S_2,
    round_digits = 4,
    filename = filename_16S_2,
    batch.size = 500,
    step = 1)
}
states_1 <- readRowsFromH5(h5_path = filename_16S_1, complete = TRUE, getTargetPositions = TRUE)
## states matrix has 2058 rows and 2 columns
pred_1 <- states_1[[1]]</pre>
position_1 <- start[1] - buffer + 1 + states_1[[2]] - floor(maxlen/2)</pre>
df_1 <- cbind(pred_1, position_1) %>% as.data.frame()
colnames(df_1) <- c("conf_16S", "conf_bacteria", "seq_end")</pre>
states_2 <- readRowsFromH5(h5_path = filename_16S_2, complete = TRUE, getTargetPositions = TRUE)
## states matrix has 2058 rows and 2 columns
pred_2 <- states_2[[1]]</pre>
position_2 <- start[2] - buffer + 1 + states_2[[2]] - floor(maxlen/2)</pre>
df_2 <- cbind(pred_2, position_2) %>% as.data.frame()
colnames(df_2) <- c("conf_16S", "conf_bacteria", "seq_end")</pre>
and plot the results again
p1 \leftarrow ggplot(df_1, aes(x = seq_end, y = conf_16S)) + geom_point() +
  geom_rect(aes(xmin=start[1], xmax=end[1], ymin=0, ymax=Inf), alpha = 0.01) +
  xlim(c(start[1] - buffer, end[1] + buffer)) +
 ylab("16S confidence") + xlab("sequence position") +
  geom_smooth()
p2 <- ggplot(df_2, aes(x = seq_end, y = conf_16S)) + geom_point() +
  geom_rect(aes(xmin=start[2], xmax=end[2], ymin=0, ymax=Inf), alpha = 0.01) +
  xlim(c(start[2] - buffer, end[2] + buffer)) +
  ylab("16S confidence") + xlab("sequence position") +
  geom_smooth()
ggpubr::ggarrange(p1, p2, ncol = 1, nrow = 2)
```



Tensorboard

We can use tensorboard to monitor our training runs. To track the runs, we have to specify a path for tensorboard files and give the run a unique name.

```
# trainNetwork(run.name = "unique_run_name",
# tensorboard.log = "tensorboard_path",
# ...
# )
```

We can inspect our previous training runs in tensorboard

```
## open tensorboard in browser
# tensorflow::tensorboard(tensorboard.log)
```

The "SCALARS" tab displays accuracy, loss and percentage of files seen for each epoch

In the "IMAGES" tab, we implemented a display of train and validation confusion matrices after every epoch. We can see for our binary classification of bacteria/16S sequences, that the model misclassified more bacteria sequences as 16S than vice versa in one of the epochs.

The "TEXT" tab shows the trainNetwork call as text.

The "HPARAM" tab tracks the hyper parameters of the different runs (maxlen, batch size etc.). This can helpful to find the optimal hyper parameter settings for a given task

Further tensorboard documentation can be found here.

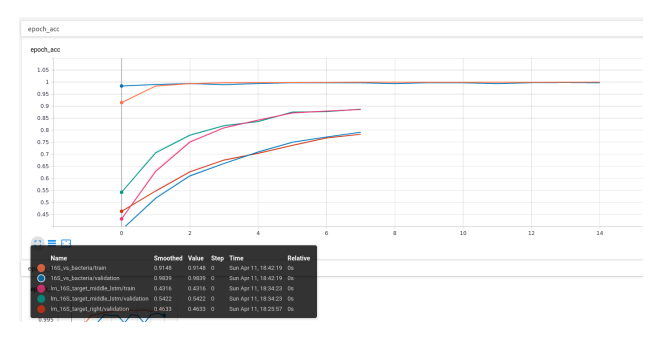


Figure 4: accuracy

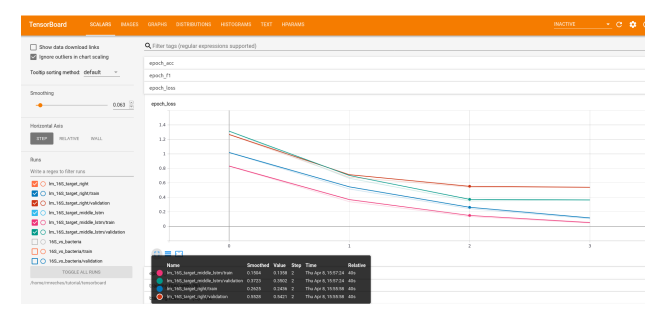


Figure 5: loss

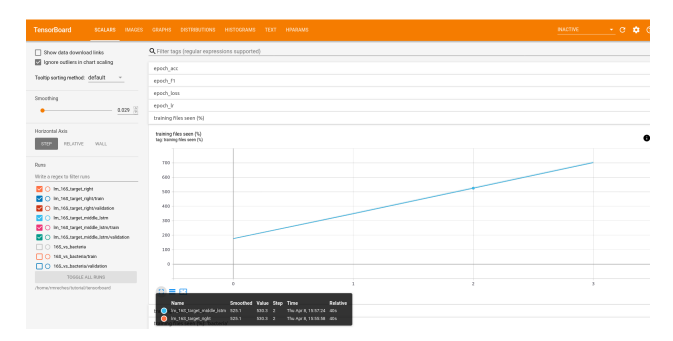


Figure 6: percentage of seen training files

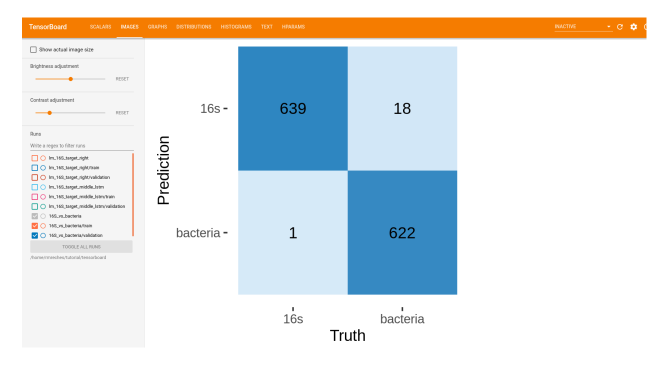


Figure 7: confusion matrix

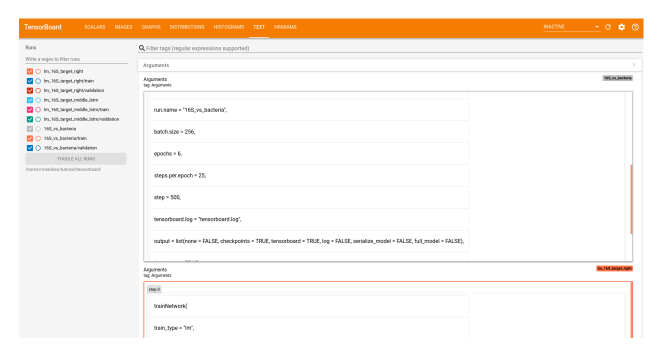


Figure 8: trainNetwork call



 $Figure \ 9: \ hyperparameters$

Integrated gradient

To visualize which parts of an input sequence is important for the models decision, we may use a method called Integrated Gradient (paper). This compares the predictions for a given sequence to a baseline sequence (usually a zero tensor or shuffling of original sequence) to determine which features were important for the models decision.

```
# load trained model
model_path <- "pretrained_models/bact_16S_model.hdf5"
model <- keras::load_model_hdf5(model_path, compile = FALSE)
model <- keras::load_model_weights_hdf5(model, model_path)</pre>
```

```
fasta_path <- file.path(path_16S_validation,</pre>
                          "GCF_000427035.1_09mas018883_genomic.16s.fasta.fasta")
fasta_file <- microseq::readFasta(fasta_path)</pre>
# extract input tensors with data generator
gen <- labelByFolderGenerator(corpus.dir = fasta path,
                                batch.size = 1,
                                maxlen = 500,
                                reverseComplements = FALSE,
                                numTargets = 2,
                                onesColumn = 1,
                                step = 1,
                                padding = FALSE)
ig_list <- list()</pre>
seq_len <- nchar(fasta_file$Sequence)</pre>
for (i in 1:(seq_len - 500)) {
  z <- gen()
  input <- z[[1]]
  ig <- integrated_gradients(m_steps = 50,</pre>
                               baseline type = "shuffle",
                               input_seq = input,
                               target class idx = 1,
                               model = model,
                               num_baseline_repeats = 2)
  py$integrated_grads <- ig</pre>
  py_run_string("attribution_mask = tf.reduce_sum(tf.math.abs(integrated_grads), axis=-1)")
  attribution_mask <- py$attribution_mask</pre>
  attribution_mask <- as.matrix(attribution_mask, ncol = 1)</pre>
  df <- data.frame(position = (1:(nrow(attribution_mask))) + i - 1,</pre>
                    ig_sum = attribution_mask[ , 1])
  ig_list[[i]] <- df</pre>
#saveRDS(ig_list, dir_path)
ig_df <- data.table::rbindlist(ig_list) %>% as.data.frame()
ig_df <- aggregate(x = ig_df$ig_sum,</pre>
                            by = list(ig_df$position),
                            FUN = mean)
names(ig_df) <- c("position", "ig_sum")</pre>
```

```
ggplot(ig_df, aes(x = position, y = ig_sum, alpha = 0.001)) +
  #geom_point() +
geom_smooth() + ylab("feature importance") +
theme(legend.position = "none") +
geom_line()
```

