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1 Loading Data

All the working steps rely on the climate4R packages (loadeR, transformeR, downscaleR, visualizeR and climate4R.value), which is a set of libraries especifically developed to handle climate data. In particular to this study, climate4R functions are through the loading, post-processing, downscaling, validation and visualization of the data. Especific notebooks have been developed to illustrate the working of climate4R functions and we refer the reader to the github repository that allocates these notebooks for more explanation on the particularities of every package and function.

Therefore, we load the core libraries of climate4R: loadeR, transformeR, downscaleR and visualizeR. We also load climate4R.value of climate4R which would permit us to compute the validation indices as well as other auxiliary libraries mainly for plotting concerns.

```
library(loadeR)
library(transformeR)
library(downscaleR)
library(visualizeR)
library(climate4R.value)
library(magrittr)
library(gridExtra)
library(RColorBrewer)
library(sp)
```

To load the data we rely on the loadeR package who permits an easy access to the datasets listed in the User Data Getaway (UDG) which is maintained by the Santander Meteorology Group. To access the datasets we first have to log in to our UDG account. If you do not have an account follow the instructions in the UDG description page

```
loginUDG(username = "", password = "")
```

In order to avoid possible errors while running the notebook, you have to set the path to your desired working directory and create two files named "Data" and "models", that will contain the downscaled predictions and the trained deep models, respectively. Moreover, as we perform 2 distinct studies, one for precipitation and other for temperature, you should create 2 new directories named "precip" and "temperature" within the previous created directories (i.e., "Data" and "models"). An example of the latter would be "personalpath/Data/precip". The predictions and models infered will be automatically saved in these folders and therefore the not creating them will end into saving errors across the notebook.

```
path = ""
setwd(path)
dir.create("Data")
```

```
dir.create("Data/precip")
dir.create("models")
dir.create("models/precip")
```

We find the label associated to ERA-Interim via the UDG.datasets() function of loadeR: "ECMWF_ERA-Interim-ESD". Then we load the predictors by calling loadGridData of loadeR.

The E-OBS dataset version 14 is also accesible through UDG. Thus, we load the predictand dataset by calling again loadGridData.

We split into train (i.e., 1979-2002) and test (i.e., 2003-2008).

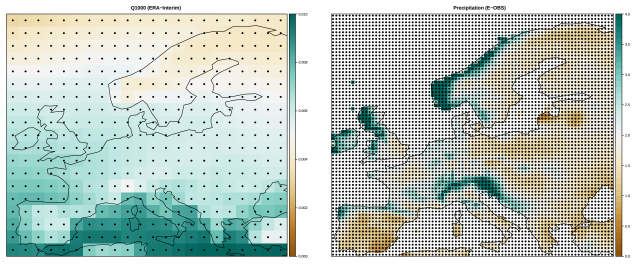
```
# Train
xT <- subsetGrid(x,years = 1979:2002)
yT <- subsetGrid(y,years = 1979:2002)
yT_bin <- binaryGrid(yT,threshold = 1,condition = "GT")
# Test
xt <- subsetGrid(x,years = 2003:2008)
yt <- subsetGrid(y,years = 2003:2008)
yt_bin <- binaryGrid(yt,threshold = 1,condition = "GT")

save(xT,file = "./Data/precip/xT.rda")
save(xt,file = "./Data/precip/xt.rda")
save(yT,file = "./Data/precip/yT.rda")
save(yt_bin,file = "./Data/precip/yt_bin.rda")
rm(x,y)</pre>
```

We can take a look at the grid resolutions of ERA-Interim and E-OBS in order to better visualize the gap we try to bridge with the downscaling.

```
colsindex <- brewer.pal(n = 9, "BrBG")
cb <- colorRampPalette(colsindex)
coords_x <- expand.grid(xt$xyCoords$x,xt$xyCoords$y) ; names(coords_x) <- c("x","y")
coords_y <- expand.grid(yt$xyCoords$x,yt$xyCoords$y) ; names(coords_y) <- c("x","y")
pplot <- list()</pre>
```

```
pplot[[1]] <- spatialPlot(climatology(subsetGrid(xt,var = "hus@1000")), backdrop.theme = "coastline",</pre>
                           main = "Q1000 (ERA-Interim)",
                           col.regions = cb,
                           at = seq(0,0.01, 0.0001),
                           set.min = 0, set.max = 0.01, colorkey = TRUE,
                           sp.layout = list(list(SpatialPoints(coords_x),
                                                 first = FALSE, col = "black",
                                                 pch = 20, cex = 1)))
pplot[[2]] <- spatialPlot(climatology(yt), backdrop.theme = "coastline",</pre>
                          main = "Precipitation (E-OBS)",
                           col.regions = cb,
                           at = seq(0, 4, 0.1),
                           set.min = 0, set.max = 4, colorkey = TRUE,
                           sp.layout = list(list(SpatialPoints(coords_y),
                                                 first = FALSE, col = "black",
                                                 pch = 20, cex = 1)))
lay = rbind(c(1,2))
grid.arrange(grobs = pplot, layout_matrix = lay)
```



We can visualize some statistics of the train and test distributions, such as the climatology, the frequency of rainy days and the percentile 98th in order to gain knowledge about the observed data. To compute the statistics we use the library climate 4R. value of climate 4R.

```
pplot <- at <- list()
n1 <- 0; n2 <- 3
indexNames <- c("Climatology", "Frequency of rain", "P98")
for (indexName in indexNames) {
   if (indexName == "Climatology") {
      indexTrain <- valueIndex(yT,index.code = "Mean")$Index %>% redim()
      indexTest <- valueIndex(yt,index.code = "Mean")$Index %>% redim()
      at[[1]] <- seq(0, 4, 0.1); at[[2]] <- seq(-1, 1, 0.1)
   }
   if (indexName == "Frequency of rain") {
      indexTrain <- valueIndex(yT_bin,index.code = "Mean")$Index %>% redim()
      indexTest <- valueIndex(yt_bin,index.code = "Mean")$Index %>% redim()
      at[[1]] <- seq(0, 0.5, 0.01); at[[2]] <- seq(-0.1, 0.1, 0.01)
}</pre>
```

```
if (indexName == "P98") {
    indexTrain <- valueIndex(yT,index.code = "P98")$Index %>% redim()
    indexTest <- valueIndex(yt,index.code = "P98")$Index %>% redim()
    at[[1]] \leftarrow seq(10, 20, 0.25); at[[2]] \leftarrow seq(-5, 5, 0.2)
  }
  for (i in 1:2) {
    if (i == 1) {
      dataset <- "(train)"; index <- indexTrain; n1 <- n1 + 1; n <- n1</pre>
    if (i == 2) {
      indexTest <- gridArithmetics(indexTest,indexTrain,operator = "-")</pre>
      dataset <- "(test bias)"; index <- indexTest; n2 <- n2 + 1; n <- n2
    pplot[[n]] <- spatialPlot(climatology(index), backdrop.theme = "coastline",</pre>
                                 main = paste(indexName,paste0(dataset,":"),
                                                round(mean(abs(index$Data), na.rm = TRUE), digits = 2)),
                                 col.regions = cb,
                                 at = at[[i]],
                                 set.min = at[[i]][1], set.max = at[[i]][length(at[[i]])],
                                 colorkey = TRUE)
lay = rbind(c(1,2,3),
             c(4,5,6)
grid.arrange(grobs = pplot, layout_matrix = lay)
                                                                                   P98 (train): 14.78
       Climatology (test bias): 0.16
                                           Frequency of rain (test bias): 0.02
```

Once the data is loaded we standardize the predictors by calling scaleGrid function of transformeR.

```
xt <- scaleGrid(xt,xT, type = "standardize", spatial.frame = "gridbox") %>% redim(drop = TRUE)
xT <- scaleGrid(xT, type = "standardize", spatial.frame = "gridbox") %>% redim(drop = TRUE)
```

2 Downscaling

2.1 Generalized Linear Models (GLM) - downscaleR

To downscale via generalized linear models (GLM) we rely on the downscaleR package of climate4R. In particular, we use the downscaleChunk function of downscaleR to downscale the precipitation. Below we define a new function, called trainPredictGLM, which encapsulates 4 calls to downscaleChunk that outputs the predictions for the occurrence (both deterministic and stochastic) and the quantity of precipitation (both deterministic and stochastic). Therefore, we input to the function the predictor (x), the number of local predictors to be used (neighbours), the predictand (y) and the test set where to apply the infered relationship. Moreover, the predictions are saved in a especific path indicated by the user through the function parameter 'filename'.

Note that downscaleChunk temporarily creates .rda files in your working directory, containing the predictions per chunk. The final binded predictions are saved in your previously created "Data/precip/" directory.

```
trainPredictGLM <- function(x,y,newdata,neighbours=1,filename) {</pre>
  y.ocu <- binaryGrid(y,condition = "GT",threshold = 1)</pre>
  y.rest <- gridArithmetics(y,1,operator = "-")</pre>
  # Logistic Regression (DETERMINISTIC)
  pred <- downscaleChunk(x = x, y = y.ocu, newdata = newdata,</pre>
                 method = "GLM", family = binomial(link = "logit"),
                 prepareData.args = list(local.predictors = list(n=neighbours, vars = getVarNames(x)))
  predDET_ocu_train <- pred[[1]] %>% redim(drop = TRUE)
  predDET_ocu <- pred[[2]] %>% redim(drop = TRUE)
  # Logistic Regression (STOCHASTIC)
  pred <- downscaleChunk(x = x, y = y.ocu, newdata = newdata,</pre>
                 method = "GLM", family = binomial(link = "logit"), simulate = TRUE,
                 prepareData.args = list(local.predictors = list(n=neighbours, vars = getVarNames(x)))
  )
  predSTO_ocu <- pred[[2]] %>% redim(drop = TRUE)
  # Gamma Regression with link logarithmic (DETERMINISTIC)
  pred <- downscaleChunk(x = x, y = y.rest, newdata = newdata,</pre>
                 method = "GLM", family = Gamma(link = "log"), condition = "GT", threshold = 0,
                 prepareData.args = list(local.predictors = list(n=neighbours, vars = getVarNames(x)))
  predDET_reg <- pred[[2]] %>% redim(drop = TRUE)
  # Gamma Regression with link logarithmic (STOCHASTIC)
  pred <- downscaleChunk(x = x, y = y.rest, newdata = newdata,</pre>
                 method = "GLM", family = Gamma(link = "log"), simulate = TRUE,
                 condition = "GT", threshold = 0,
                 prepareData.args = list(local.predictors = list(n=neighbours, vars = getVarNames(x)))
  predSTO_reg <- pred[[2]] %>% redim(drop = TRUE)
  ###
  save(predDET_ocu,predDET_reg,
       predSTO_ocu,predSTO_reg,
```

```
file = paste0("./Data/precip/predictions_test_",filename,".rda"))
predDET_ocu <- predDET_ocu_train
save(predDET_ocu,
    file = paste0("./Data/precip/predictions_train_",filename,".rda"))
}</pre>
```

Now, we downscale by calling the function defined above.

2.2 Downscaling - Deep Neural Networks

2.2.1 Training

To infer deep learning models we rely on Keras.

```
library(keras)
```

In this work, all deep learning functions have optimized the negative log-likelihood of a Bernouilli-Gamma distribution. We implement the custom loss in Keras.

```
discreteGamma <- custom_metric("custom_loss", function(true, pred){
   K <- backend()
   D <- K$int_shape(pred)[[2]]/3
   ocurrence = pred[,1:D]
   shape_parameter = K$exp(pred[,(D+1):(D*2)])
   scale_parameter = K$exp(pred[,(D*2+1):(D*3)])
   bool_rain = K$cast(K$greater(true,0),K$tf$float32)
   epsilon = 0.000001
   return (- K$mean((1-bool_rain)*K$tf$log(1-ocurrence+epsilon) + bool_rain*(K$tf$log(ocurrence+epsilon))
})</pre>
```

In the following we provide the code related to the deep learning architectures.

```
architectures <- function(architecture,input_shape,output_shape) {
  if (architecture == "CNN-LM") {
    inputs <- layer_input(shape = input_shape)
    x = inputs
    11 = layer_conv_2d(x ,filters = 50, kernel_size = c(3,3), activation = 'linear', padding = "same")
    12 = layer_conv_2d(l1,filters = 25, kernel_size = c(3,3), activation = 'linear', padding = "same")
    13 = layer_conv_2d(l2,filters = 1, kernel_size = c(3,3), activation = 'linear', padding = "same")
    14 = layer_flatten(l3)
    parameter1 = layer_dense(l4,units = output_shape, activation = "sigmoid")
    parameter2 = layer_dense(l4,units = output_shape)
    parameter3 = layer_dense(l4,units = output_shape)
    outputs = layer_concatenate(list(parameter1,parameter2,parameter3))
    model <- keras_model(inputs = inputs, outputs = outputs)
}</pre>
```

```
if (architecture == "CNN1") {
  inputs <- layer_input(shape = input_shape)</pre>
 x = inputs
 11 = layer_conv_2d(x ,filters = 50, kernel_size = c(3,3), activation = 'relu', padding = "same")
 12 = layer_conv_2d(11,filters = 25, kernel_size = c(3,3), activation = 'relu', padding = "same")
 13 = layer_conv_2d(12, filters = 1, kernel_size = c(3,3), activation = 'relu', padding = "same")
 14 = layer_flatten(13)
 parameter1 = layer dense(14,units = output shape, activation = "sigmoid")
 parameter2 = layer dense(14,units = output shape)
 parameter3 = layer_dense(14,units = output_shape)
 outputs = layer_concatenate(list(parameter1,parameter2,parameter3))
 model <- keras_model(inputs = inputs, outputs = outputs)</pre>
}
if (architecture == "CNN10") {
  inputs <- layer_input(shape = input_shape)</pre>
 x = inputs
 11 = layer_conv_2d(x ,filters = 50, kernel_size = c(3,3), activation = 'relu', padding = "valid")
 12 = layer_conv_2d(11,filters = 25, kernel_size = c(3,3), activation = 'relu', padding = "valid")
 13 = layer_conv_2d(12, filters = 10, kernel_size = c(3,3), activation = 'relu', padding = "valid")
 14 = layer_flatten(13)
 parameter1 = layer_dense(14,units = output_shape, activation = "sigmoid")
 parameter2 = layer_dense(14,units = output_shape)
 parameter3 = layer_dense(14,units = output_shape)
 outputs = layer concatenate(list(parameter1, parameter2, parameter3))
 model <- keras_model(inputs = inputs, outputs = outputs)</pre>
}
if (architecture == "CNN-PR") {
 inputs <- layer_input(shape = input_shape)</pre>
 x = inputs
 11 = layer_conv_2d(x ,filters = 10, kernel_size = c(3,3), activation = 'relu', padding = "valid")
 12 = layer_conv_2d(11,filters = 25, kernel_size = c(3,3), activation = 'relu', padding = "valid")
 13 = layer_conv_2d(12, filters = 50, kernel_size = c(3,3), activation = 'relu', padding = "valid")
 14 = layer_flatten(13)
 parameter1 = layer_dense(14,units = output_shape, activation = "sigmoid")
 parameter2 = layer_dense(14,units = output_shape)
 parameter3 = layer_dense(14,units = output_shape)
 outputs = layer_concatenate(list(parameter1,parameter2,parameter3))
  model <- keras_model(inputs = inputs, outputs = outputs)</pre>
}
if (architecture == "CNNdense") {
  inputs <- layer_input(shape = input_shape)</pre>
 x = inputs
 11 = layer_conv_2d(x ,filters = 50, kernel_size = c(3,3), activation = 'relu', padding = "valid")
 12 = layer_conv_2d(11,filters = 25, kernel_size = c(3,3), activation = 'relu', padding = "valid")
 13 = layer_conv_2d(12, filters = 10, kernel_size = c(3,3), activation = 'relu', padding = "valid")
 14 = layer_flatten(13)
 15 = layer_dense(14,units = 50, activation = "relu")
 16 = layer_dense(15,units = 50, activation = "relu")
 parameter1 = layer_dense(16,units = output_shape, activation = "sigmoid")
 parameter2 = layer_dense(16,units = output_shape)
```

```
parameter3 = layer_dense(16,units = output_shape)
  outputs = layer_concatenate(list(parameter1,parameter2,parameter3))
  model <- keras_model(inputs = inputs, outputs = outputs)
}
return(model)
}</pre>
```

To train the latter architectures we have encapsulated them into a more general funcion called trainDEEP.

```
trainDEEP <- function(x,y,architecture, epochs=10000,patience=30,
                      learning_rate = 0.0001,loss_function = discreteGamma){
  x <- x$Data
  x <- x \% aperm(c(2,3,4,1))
  y <- y$Data
  dim(y) \leftarrow c(dim(y)[1], dim(y)[2]*dim(y)[3])
  indLand <- (!apply(y,MARGIN = 2,anyNA)) %>% which()
  y <- y[,indLand]
  y < -y - 1
  y[which(y < 0, arr.ind = TRUE)] < 0
  callbacks <- list(callback_early_stopping(patience = patience),</pre>
                    callback_model_checkpoint(filepath=paste0('./models/precip/',architecture,'.h5'),
                                               monitor='val_loss', save_best_only=TRUE)
  )
  model <- architectures(architecture,input_shape = dim(x)[-1],output_shape = ncol(y))</pre>
  model %>% compile(optimizer = optimizer_adam(lr = learning_rate), loss = loss_function)
  model %>% fit(x, y, epochs = epochs, batch size = 100,
                validation_split = 0.1, callbacks = callbacks, verbose = 0)
  k clear session()
}
```

Finally, we train the models and save the model in the path specified in the trainDEEP function, according to the early-stopping criteria.

```
trainDEEP(xT,yT,architecture = "CNN-LM")
trainDEEP(xT,yT,architecture = "CNN1")
trainDEEP(xT,yT,architecture = "CNN10")
trainDEEP(xT,yT,architecture = "CNN-PR")
trainDEEP(xT,yT,architecture = "CNN-PR")
```

2.2.2 Prediction

Once the models are trained and saved, we predict onto the train and test datasets. To do so, we first define a function called predictDEEP that encapsulates the deterministic and stochastic predictions. Recall that, according to the loss function, the net estimates the parameters p, α and β of a Bernouilli-Gamma distribution. In the deterministic way, the prediction for a given day is the expectance of the conditional Bernouilli-Gamma distribution infered. In the stochastic way, we sample from the conditional Bernouilli-Gamma distribution. This sampling is coded in the functions simulateOcu and simulateReg.

```
predictDEEP <- function(x,template,architecture,dataset) {
  loss_function <- discreteGamma
  model <- load_model_hdf5(filepath = paste0("./models/precip/",architecture,".h5"),</pre>
```

```
custom_objects = c("custom_loss" = loss_function))
x \leftarrow xData %>% aperm(c(2,3,4,1))
pred <- model$predict(x)</pre>
D \leftarrow ncol(pred)/3
ntime <- dim(template$Data)[1]</pre>
nlat <- dim(template$Data)[2]</pre>
nlon <- dim(template$Data)[3]</pre>
# Deterministic Prediction
predDET_ocu <- pred[,1:D]</pre>
predDET_reg \leftarrow exp(pred[,(D+1):(D*2)])*exp(pred[,(D*2+1):(D*3)]) + 1
# Stochastic Prediction
simulate_ocu <- function(dat,model,D){</pre>
  ocu <- model$predict(dat)[,1:D,drop = FALSE]</pre>
  sim <- matrix(runif(length(ocu),min = 0,max = 1), nrow = nrow(ocu), ncol = ncol(ocu))</pre>
  cond <- ocu > sim
  return(cond*1)
}
simulate_reg <- function(dat,model,D) {</pre>
  shape <- exp(model$predict(dat)[,(D+1):(D*2),drop = FALSE])</pre>
  scale <- exp(model$predict(dat)[,(D*2+1):(D*3),drop = FALSE])</pre>
  p <- matrix(nrow = nrow(dat),ncol = D)</pre>
  for (i in 1:D) {
    p[,i] <- rgamma(n = nrow(dat), shape = shape[,i], scale = scale[,i])</pre>
  }
  return(p)
predSTO_ocu <- simulate_ocu(x,model,D)</pre>
predSTO_reg <- simulate_reg(x,model,D) + 1</pre>
# Converting to 2D-map (including sea)
yT1D <- yT$Data
dim(yT1D) <- c(dim(yT1D)[1],nlat*nlon)</pre>
indLand <- (!apply(yT1D,MARGIN = 2,anyNA)) %>% which()
convert2map2D <- function(grid, template) {</pre>
  dim(template$Data) <- c(ntime,nlat*nlon)</pre>
  out <- template
  out$Data[,indLand] <- grid</pre>
  dim(out$Data) <- c(ntime,nlat,nlon)</pre>
  return(out)
}
predDET_ocu <- convert2map2D(predDET_ocu,template)</pre>
predDET_reg <- convert2map2D(predDET_reg,template)</pre>
predSTO_ocu <- convert2map2D(predSTO_ocu,template)</pre>
predSTO_reg <- convert2map2D(predSTO_reg,template)</pre>
save(predDET_ocu,predDET_reg,predSTO_ocu,predSTO_reg,
     file = paste0("./Data/precip/predictions_",dataset,"_",architecture,".rda"))
k_clear_session()
```

Then we predict and save the predictions.

```
# Train
predictDEEP(xT,architecture = "CNN-LM",template = yT,dataset = "train")
predictDEEP(xT,architecture = "CNN1",template = yT,dataset = "train")
predictDEEP(xT,architecture = "CNN10",template = yT,dataset = "train")
predictDEEP(xT,architecture = "CNN-PR",template = yT,dataset = "train")
predictDEEP(xT,architecture = "CNN-PR",template = yT,dataset = "train")

# Test
predictDEEP(xt,architecture = "CNN-LM",template = yt,dataset = "test")
predictDEEP(xt,architecture = "CNN1",template = yt,dataset = "test")
predictDEEP(xt,architecture = "CNN10",template = yt,dataset = "test")
predictDEEP(xt,architecture = "CNN-PR",template = yt,dataset = "test")
predictDEEP(xt,architecture = "CNN-PR",template = yt,dataset = "test")
predictDEEP(xt,architecture = "CNN-PR",template = yt,dataset = "test")
```

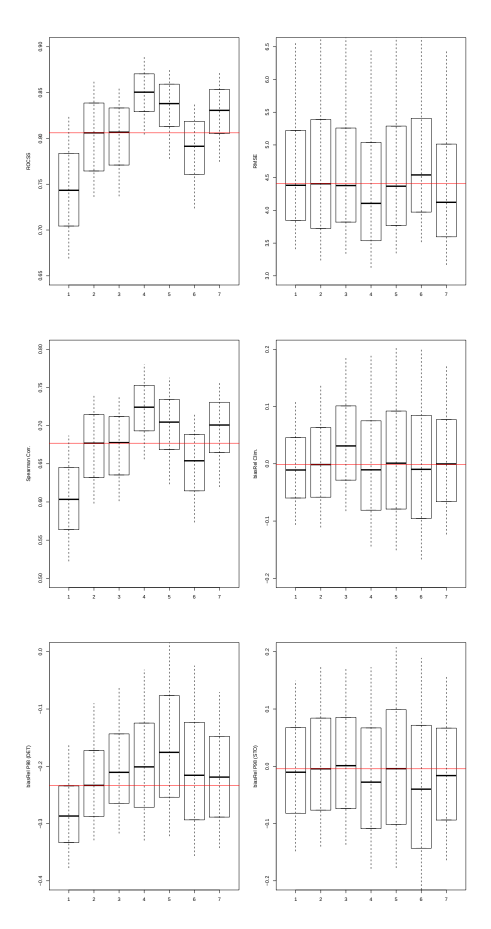
3 Validation of the Results

3.1 Intercomparison of Methods

In this figure, we calculate the validation indices by using the library climate4R.value of climate4R. In particular the indices used are: the Roc Skill Score (ROCSS), the Root Mean Squared Error (RMSE), the spearman correlation and the relative biases of the climatology and of the percentile 98 of the rainy days distribution (both deterministic and stochastic predictions). The indices are plotted with the function violinPlot of climate4R's package visualizeR.

```
# file_name <- paste0("./figures/fig04_precipitation.pdf")</pre>
# pdf(file = file_name, width = 5, height = 10)
par(mfrow = c(3,2))
filepreds=list("glm1", "glm4",
                "CNN-LM", "CNN1", "CNN10",
                "CNN-PR", "CNNdense")
datasetspred <- length(filepreds)</pre>
indexNames <- c("ROCSS", "RMSE", "Spearman Corr.",</pre>
                 "biasRel Clim.", "biasRel P98 (DET)", "biasRel P98 (STO)")
pplot <- list(); n <- 0</pre>
for (indexName in indexNames) {
  index <- array(dim = c(datasetspred,dim(yT$Data)[2:3]))</pre>
  for (i in 1:datasetspred) {
    load(paste0("./Data/precip/predictions_train_",filepreds[[i]],".rda"))
    predDET_ocu_train <- predDET_ocu</pre>
    load(paste0("./Data/precip/predictions_test_",filepreds[[i]],".rda"))
    predDET_bin <- binaryGrid(predDET_ocu,ref.obs = yT_bin,ref.pred = predDET_ocu_train)</pre>
    predDET <- gridArithmetics(predDET_bin,predDET_reg,operator = "*")</pre>
    predSTO <- gridArithmetics(predSTO_ocu,predSTO_reg,operator = "*")</pre>
    if (indexName == "ROCSS") {
      index[i,,] <- valueMeasure(yt bin,predDET ocu,</pre>
                                   measure.code="ts.rocss")$Measure$Data
      ylim < -c(0.65,0.9)
    }
    if (indexName == "RMSE") {
```

```
index[i,,] <- valueMeasure(yt,predDET_reg,measure.code="ts.RMSE",</pre>
                                condition = "GT", threshold = 1,
                                which.wetdays = "Observation")$Measure$Data
    ylim < -c(3,6.5)
 }
 if (indexName == "Spearman Corr.") {
    index[i,,] <- valueMeasure(yt,predDET,measure.code="ts.rs")$Measure$Data</pre>
    vlim < c(0.5, 0.8)
 }
 if (indexName == "biasRel Clim.") {
    index[i,,] <- valueMeasure(yt,predDET,measure.code="biasRel",index.code="Mean")$Measure$Data</pre>
    ylim < -c(-0.2,0.2)
 if (indexName == "biasRel P98 (DET)") {
    index[i,,] <- valueMeasure(yt,predDET,measure.code="biasRel",index.code="P98")$Measure$Data
   ylim <- c(-0.4,0.0)
 }
 if (indexName == "biasRel P98 (STO)") {
    index[i,,] <- valueMeasure(yt,predSTO,measure.code="biasRel",index.code="P98")$Measure$Data</pre>
    ylim < c(-0.2, 0.2)
n < - n + 1
dim(index) <- c(datasetspred,prod(dim(yT$Data)[2:3]))</pre>
indLand <- (!apply(index,MARGIN = 2,anyNA)) %>% which()
index <- index[,indLand] %>% t()
mglm4 <- median(index[,2],na.rm = TRUE)</pre>
perc <- apply(index,MARGIN = 2,FUN = function(z) quantile(z,probs = c(0.1,0.9)))</pre>
boxplot(index, outline = FALSE, asp = 1, ylim = ylim, range = 0.0001, ylab = indexName, asp = 1)
lines(c(0,8),c(mglm4,mglm4), col = "red", asp = 1)
for (i in 1:datasetspred) lines(c(i,i),perc[,i], lty = 2)
```



3.2 Spatial Maps

The above indices can be visualized spatially per model, this is, observing the results per gridbox. To do so, we define a function called experiment1 that encapsulates the code needed for plotting. The spatial map is plotted with the function spatialPlot of visualizeR, which we remind is a package of climate4R.

```
experiment1 <- function(model) {</pre>
  cb <- colorRampPalette(brewer.pal(9, "OrRd"))(80)</pre>
  colsindex <- rev(brewer.pal(n = 9, "RdBu"))</pre>
  cb2 <- colorRampPalette(colsindex)</pre>
  load(paste0("./Data/precip/predictions_train_",model,".rda"))
  predDET_ocu_train <- predDET_ocu</pre>
  load(paste0("./Data/precip/predictions_test_",model,".rda"))
  predDET_bin <- binaryGrid(predDET_ocu,ref.obs = yT_bin,ref.pred = predDET_ocu_train)</pre>
  predDET <- gridArithmetics(predDET_bin,predDET_reg,operator = "*")</pre>
  indexNames <- c("ROCSS", "RMSE", "Spearman Corr.",</pre>
                    "biasRel Clim.", "biasRel P98 (DET)")
  pplot <- list(); n <- 0</pre>
  for (indexName in indexNames) {
    if (indexName == "ROCSS") {
      index <- valueMeasure(yt_bin,predDET_ocu,</pre>
                               measure.code="ts.rocss")$Measure %>% redim()
      at \leftarrow seq(0.5, 1, 0.01); colorbar \leftarrow cb
    }
    if (indexName == "RMSE") {
      index <- valueMeasure(yt,predDET_reg,</pre>
                               measure.code="ts.RMSE",
                               condition = "GT", threshold = 1,
                               which.wetdays = "Observation")$Measure %>% redim()
      at \leftarrow seq(2, 6.5, 0.25); colorbar \leftarrow cb
    }
    if (indexName == "Spearman Corr.") {
      index <- valueMeasure(yt,predDET,</pre>
                               measure.code="ts.rs")$Measure %>% redim()
      at \leftarrow seq(0.5, 1, 0.02); colorbar \leftarrow cb
    }
    if (indexName == "biasRel Clim.") {
      index <- valueMeasure(yt,predDET,</pre>
                               measure.code="biasRel",index.code="Mean")$Measure %>% redim()
      at \leftarrow seq(-0.5, 0.5, 0.01); colorbar \leftarrow cb2
    if (indexName == "biasRel P98 (DET)") {
      index <- valueMeasure(yt,predDET,</pre>
                               measure.code="biasRel",index.code="P98") $ Measure %>% redim()
       at \leftarrow seq(-0.5, 0.5, 0.01); colorbar \leftarrow cb2
    }
    n < -n + 1
```

Now we call the experiment 1 to obtain the figures for the following methods: glm1, glm4 and CNN1.

```
figure ([1]] <- experiment1(model = "glm1")
figure [[3]] <- experiment1(model = "glm4")
figure [[3]] <- experiment1(model = "CNN1")

lay = rbind(1,2,3)
grid.arrange(grobs = figure,layout_matrix = lay)

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```

4 Technical Aspects

Due to the different computational requirements at the different stages of the study we have relied on two different machines, one local (HP-ProDesk-600-G2-MT) and other accessed remotely (virtual machine). Specific characteristics of both machines are indicated below. The former was used to load, process, validate and visualize the results but lacked from memory to perform downscaling with generalized linear models. Despite the downscaleChunk function is a wrapper of downscaleTrain and downscalePredict (we refer the reader to the to the github donwscaleR repository for more details concerning these functions) where downscaling is performed by chunking into latitudes (i.e., avoiding to work with the whole predictand domain simultaneously) and therefore suitable for high demanding computational downscaling, this stage could not

be done locally. For this reason we relied on a virtual machine with more memory capacity. On the other hand, deep learning models were successfully trained locally without the need to access remotely to the virtual machine. Moreover there were differences in the training times. Calling the function trainPredictGLM for downscaling the GLMs took a magnitude of hours whereas the deep learning models were trained in a maximum of 1 hour despite their overparameterized structure in comparison with the linear models.

1. Local Machine (HP-ProDesk-600-G2-MT)

• Operating system: ubuntu 16.04 LTS

• Memory: 15.6 GiB

- Processor: Intel® Core
™ i7-6700 CPU @ 3.40GHz × 8

SO: 64 bitsDisc: 235.1 GiB

2. Virtual Machine

• Operating system: ubuntu 16.04 LTS

• Memory: 244.1 GiB

- Processor: Intel® Xeon
TM CPU E7-4870 @ 2.40GHz × 4

• SO: 64 bits

• Disc: 1 TB for users