Worked Example

April 21, 2021

1 Brief user guide for the RandomForestDist package

This companion notebook briefly explains how to use the main functionalities provided by the R package RandomForestDist and presents the code needed to reproduce part of the experiments presented in the article Adapting Random Forests for Stochastic Downscaling of Rainfall Based on Local Estimated Probability Distributions, submitted to Water Resources Research by Legasa et al. in 2021. In that work, random forests (RFs) are applied to the problem of statistical downscaling of rainfall. In particular, the authors analyze the suitability of different split functions which allow to work with non-normally distributed variables and propose a novel a posteriori (AP) approach which permits to accurately estimate the shape and rate parameters of the underlying rainfall distribution, which in turn allows for generating reliable stochastic rainfall series.

RandomForestDist requires a modified version of rpart which can be found here. In order to run the examples provided below, these two packages need to be first installed. This can be easily done from GitHub using the devtools package:

```
[]: devtools::install_github("MNLR/rpart")
  install.packages(c("progressr", "qmap", "fitdistrplus"))
  devtools::install_github("MNLR/RandomForestDist")
```

We start by loading RandomForestDist and VALUE9, a dataset included in the package which contains daily series of rainfall for 9 illustrative meteorological stations over Europe along with the 17 large-scale reanalysis predictors used for statistical downscaling (see the paper for details).

```
[1]: library(RandomForestDist) data(VALUE9)
```

Loading required package: rpart

Loading required package: progressr

VALUE9 is a list of 9 elements (one per station), being the data already divided into a stratified 5-fold (F1 = 1979-1984, F2 = 1985-1990, F3 = 1991-1996, F4 = 1997-2002, F5 = 2003-2008). \$train.y(training set) and \$test.y (test set) correspond to the predictands; \$train.x (training set) and \$test.x (test set) are the predictors. For example, for the first station and fold:

```
[2]: head(cbind(VALUE9[[1]]$f1_79_84$train.y, VALUE9[[1]]$f1_79_84$train.x))
```

```
101127.6
                            5.4
                                  100934.56
                                             101076.31
                                                         101029.38
                                                                               0.9725891
                                                                                           -0.5552429
                            3.0
                                  99861.81
                                             100098.66
                                                         99890.75
                                                                     100026.2
                                                                               4.0751587
                                                                                          2.3417603
                                  100774.52
                                             101000.39
                                                         101343.70
                                                                    101556.2
                                                                               1.7720581
                                                                                           0.8526245
                            5.0
A matrix: 6 Œ 69 of type dbl
                            12.9
                                  100790.47
                                             100810.84
                                                         101059.16
                                                                    101159.5
                                                                               4.7946411
                                                                                           4.9987427
                            11.0
                                  99843.41
                                             99968.94
                                                         100306.59
                                                                    100510.2
                                                                               8.2443481
                                                                                           8.8000122
                                             99915.36
                            1.0
                                  99936.30
                                                         100108.30
                                                                    100281.5
                                                                               7.5472656
                                                                                          9.0443359
```

1.1 Training the models

To illustrate the training process, let's just focus on the sixth station and first fold:

```
[3]: train.x <- VALUE9[[6]]$f1_79_84$train.x # predictors for the train period train.y <- VALUE9[[6]]$f1_79_84$train.y # predictand (rainfall) for the train_ period
```

The function randomForestTrain() builds the model for the predictors x and predictands y.

```
[4]: rf <- randomForestTrain(x = train.x, y = train.y) # building/training a_{\square} \rightarrow default RF
```

The user can specify the desired values for the parameters mtry (number of predictors to randomly use as candidate split variables), ntree (number of trees in the forest), minbucket (minimum number of elements each leaf is required to have), minsplit (minimum number of elements each node is required to have to attempt the splitting) and maxdepth (maximum depth allowed for each tree).

```
[5]: rf <- randomForestTrain(x = train.x, y = train.y, ntree = 25, mtry = 22, minbucket = 10, minsplit = 30, maxdepth = 10) #⊔

→building/training a customized RF
```

By default, RFs consider the root mean squared error as split function. However, RandomForestDist allows for using different split functions (split.function argument), which have been defined in the modified version of rpart. Currently supported options are "anova", "poisson", "class", "exp", "gammaLLMME", "gammaLLmean", "bernoulliGammaLLMME", "gammaDeviation", "gammaLLBC3", "bernoulliLL" and "binaryCrossEntropyGammaDeviation".

In *Legasa et al.* 2021, we restrict ourselves to the 2-parameter gamma distribution family, with different estimators (see the paper for details).

```
[6]: # NOTE: This chunk may take a while:

rf.dev <- randomForestTrain(x = train.x, y = train.y, ntree = 100, minbucket = 100, method = "gammaDeviation")

rf.LLMME <- randomForestTrain(x = train.x, y = train.y, ntree = 100, minbucket = 10, method = "gammaLLMME")

rf.LLBC3 <- randomForestTrain(x = train.x, y = train.y, ntree = 100, minbucket = 10, method = "gammaLLBC3")
```

1.2 Parallelization

The training process can be parallelized by means of the package future.apply. This package has to be installed independently, otherwise parallelization will not be an option. Use install.packages(future.apply) to install it, which will also install the dependency future.

The simplest way to use parallelization is by setting parallel.plan = "auto", which will automatically use all the available cores to train the model in parallel.

```
[7]: rf <- randomForestTrain(x = train.x, y = train.y, ntree = 100, minbucket = 10, method = "gammaDeviation", parallel.plan = "auto") # training a RF with

→parallelization
```

If not set to "auto", paralell.plan will use the execution plan stored in the strategy argument of future::plan(). The interested reader is referred to the packages future and future.apply for details on the available options. As an example, the following block of code shows how parallelization in 4 cores can be achieved in combination with the argument workers and setting parallel.plan to multisession:

Also, if missing or set to NULL (default), the function will use the current execution plan. This is useful to configure a customized plan outside the function, as we do in the next example. In particular, we set the execution to multisession() prior to launching the loop and train several models using this plan (do not forget to close the sessions by reverting to sequential()).

Note that using parallel.plan = NA will avoid parallelization altogether. This is the default if future.apply is not available.

1.3 Generating the predictions

Once the RF is trained, out-of-sample predictors (\$test.x) can be used to obtain the corresponding downscaled predictions with the function randomForestPredict(). The default configuration of randomForestPredict() considers the mean as aggregation function (this is the standard procedure in RFs; see the paper for details):

```
[10]: test.x <- VALUE9[[6]]$f1_79_84$test.x # predictors for the test period test.y <- VALUE9[[6]]$f1_79_84$test.y # predictand (rainfall) for the train_ period
```

```
# We use the RF with 100 trees and gammaDeviation split function that we have

→already learnt to predict daily rainfall:

pr <- randomForestPredict(rf, newdata = test.x) # predictions for the test

→period, considering the mean as aggregation function

head(pr)
```

1 6.69288220375256 **2** 2.60350666864955 **3** 4.30276790536605 **4** 2.22515326732951 **5** 4.25758322163732 **6** 10.8015920530665

The parameter bagging.function can be used to apply other aggregation functions (e.g.the median). Also, if set to NA, randomForestPredict() will provide the predictions returned by all the individual trees.

```
[11]: pr.median <- randomForestPredict(rf, newdata = test.x, bagging.function = median) # median as aggregation function

pr.alltrees <- randomForestPredict(rf, newdata = test.x, bagging.function = NA)

pr.alltrees[1:10,] # 10 first predictions returned by the 100 trees
```

1 1.8250000 5.2857143 2.640000 5.9166667 4.6578947 7.18750 2 0.9636364 4.282353 2.0363636 0.7727273 1.5142857 4.80000 3 4.3529412 5.2857143 2.133333 5.9166667 4.8800000 1.96153 4 1.1357143 5.2857143 2.133333 1.2909091 0.9818182 1.61666 2.4230769 2.0700000 2.761538 1.9100000 3.2315789 1.60909 A matrix: 10 Œ 100 of type dbl 13.3000000 9.5800000 8.190909 9.7200000 6.3125000 13.5235 12.5000000 11.6894737 9.240000 14.2166667 2.1142857 13.5235 8 1.7764706 0.8666667 2.747059 2.4857143 1.5916667 4.68000 9 8.8666667 6.4153846 10.507143 12.2454545 9.6153846 4.09000 10 | 11.2888889 16.2900 14.5333333 12.300000 14.0666667 13.6181818

The parameter method controls the *a posteriori* approach (the reader is referred to the paper for details). The available options are described in the help page (?randomForestPredict), and we summarize them here. If method is not NULL, the argument bagging.function is ignored.

- method = "leaves" provides all the observations for each prediction.
- method = "random.sample" draws a random sample from the leaves for each prediction.
- method = "aposteriori" returns the parameter of the probability distribution of train.y | train.x, as estimated using Moments Matching Estimation (refer to fitdistrplus::fitdist()). In the case of the gamma distribution, its shape and rate parameters are returned. method = "bc3" uses BC3 estimators for the gamma distribution. Additionally, this parameter can be used to select the estimation method (see fitdistrplus::fitdist()), i.e., one of the following: "mme" (Moments Matching Estimation; default), "mle" (Maximum Likelihood Estimation), "qme" (Quantile Matching Estimation), "mge" (Maximum Goodness-of-fit Estimation), "mse" (Maximum Spacing Estimation). Note that some of these options require additional parameters (use . . .), and some of them may not converge.

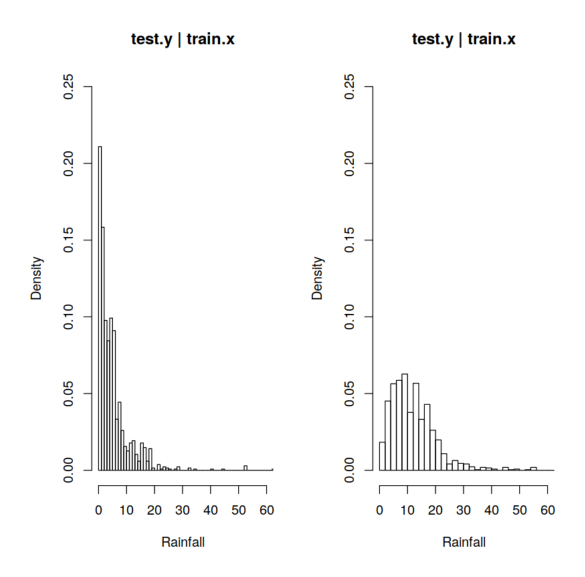
We provide some examples below.

Setting the parameter method = "leaves" allows us to plot train.y | train.x, for instance:

```
[12]: pr.leaves <- randomForestPredict(rf, newdata = test.x, method = "leaves") # we<sub>□</sub>

→ predict the observations falling on leaves
```

```
# and plot the histogram of their distribution:
par(mfrow = c(1,2))
hist(pr.leaves[[100]], breaks = 50, xlab = "Rainfall", freq = FALSE,
    main = "test.y | train.x", ylim = c(0, 0.25), xlim = c(0,60))
hist(pr.leaves[[300]], breaks = 50, xlab = "Rainfall", freq = FALSE,
    main = "test.y | train.x",ylim = c(0, 0.25), xlim = c(0,60))
```



As the model was trained using the *deviation* for the 2 parameter gamma distribution, using method = "aposteriori" provides the shape and rate parameters for each prediction:

```
[13]: pr.distr <- randomForestPredict(rf, newdata = test.x, method = "aposteriori") pr.distr[1:10,] # We show the first 10 predicted outcomes
```

	shape	rate
A matrix: 10 Œ 2 of type dbl	0.9677957	0.1419155
	0.7862478	0.2982816
	1.3759968	0.3128242
	0.7537063	0.3403316
	1.0151516	0.2371556
	1.4834291	0.1391585
	1.5673480	0.1938595
	1.2676882	0.3518086
	2.6937874	0.2916631
	4.7145554	0.3804880

As stated above, the user can also specify the estimator used. For instance, rather than the Moments Matching Estimation (default) we could use the Maximum Likelihood Estimation by setting method = "mle":

```
[14]: pr.mle <- randomForestPredict(rf, newdata = test.x, method = "mle") pr.distr[1:10,] # 10 first predicted outcomes
```

rate

	orier p	10100
A matrix: 10 Œ 2 of type dbl	0.9677957	0.1419155
	0.7862478	0.2982816
	1.3759968	0.3128242
	0.7537063	0.3403316
	1.0151516	0.2371556
	1.4834291	0.1391585
	1.5673480	0.1938595
	1.2676882	0.3518086
	2.6937874	0.2916631
	4.7145554	0.3804880

shape

Finally, the predicted distribution parameters can be used to simulate a series of stochastic rainfall values using the function randomForestSimulate(), which takes as inputs the object returned by randomForestPredict() and the number of simulated series, n. The parameter distr allows for manually selecting the distribution of interest (by default, it is automatically selected based on the training split function considered).

```
[15]: simulation <- randomForestSimulate(pr.distr)
        We simulate 1000 series:
[16]: simulations <- randomForestSimulate(pr.distr, n = 1000)
     head(simulations)
                                      3.3957831
                                                   2.6261201
                                                              4.2000766
                                                                          23.5097911
                                                                                      0.2244423
                                                                                                   2.4337384
                                      0.9729068
                                                   0.3553539
                                                              7.1575646
                                                                          0.7050184
                                                                                      3.0437166
                                                                                                   0.4486807
                                      13.5193071
                                                  1.1896116
                                                              1.8297097
                                                                          4.7100415
                                                                                      1.8269712
                                                                                                   1.5602529
        A matrix: 6 Œ 1000 of type dbl
                                      1.0754117
                                                              0.1027564
                                                   0.4053252
                                                                          6.8429978
                                                                                      0.8373085
                                                                                                   0.1429419
                                      5.5282513
                                                   5.4668850
                                                              6.2051241
                                                                          0.5221806
                                                                                       13.2525883
                                                                                                   0.1781004
                                      17.9463134
                                                  0.6897609
                                                              3.2023356
                                                                          4.5290871
                                                                                      6.5932239
                                                                                                   21.9272342
```

2 Reproducibility code for Figure 6 in Legasa et al. 2021

The code provided below can be used to obtain a similar to Figure 6 in *Legasa et al.* 2021 (we reproduce here only the continuous part of rainfall) based on a RF with 25 trees and minimum leaf size 5 which considers the gammaDeviation as split function.

```
[17]: future::plan(future::multisession, workers = 6) # parallelizing using 6 cores
     predictions <-
     lapply(VALUE9, function(meteorological.station){
         predictions <- lapply(meteorological.station, function(fold){</pre>
              # training the model
             md <- randomForestTrain(x = fold$train.x, y = fold$train.y,</pre>
                                       ntree = 25, minbucket = 5,
                                       method = "gammaDeviation",
                                       parallel.plan = NULL)
             # applying the trained model to test.x and returning the observed
      \rightarrow values $test.y:
             return(list(predicted = randomForestPredict(md, newdata = fold$test.x,_
      →method = "mme"),
                           observed = fold$test.y))
         })
         # rearranging the series
         predicted <- do.call(rbind, lapply(predictions, function(pr) pr$predicted))</pre>
         attr(predicted, "class") <- "RandomForestDist.prediction.simulable" #_</pre>
      \rightarrowrecovers the class
         observed <- do.call(c, lapply(predictions, function(pr) pr$observed))
         return(list(predicted = predicted, observed = observed))
     })
     future::plan(future::sequential) # reverting plan to normal
```

We end up with the downscaled probability distribution (parameters *shape* and *rate*) for each station and day, which are then used to generate an ensemble of 250 stochastic rainfall values. With illustrative purposes, these values are subsequently validated in terms of the SDII, SD, P95 and P95A metrics (see the paper for details).

```
# validating each simulation
        stochastic.validation <-
            apply(simulations, MARGIN = 2, FUN = function(simulation){# for_
 \rightarrow each simulation
                 P95.simulated <- quantile(simulation, probs = 0.95)
                 P95.observed <- quantile(observed, probs = 0.95)
                 return(
                     c(SDII = mean(simulation)/mean(observed),
                       SD = sd(simulation)/sd(observed),
                       P95 = P95.simulated/P95.observed,
                       P95A = sum(simulation[simulation >= P95.simulated])/
 →sum(observed[observed >= P95.observed])
                 )
            })
        # averaging the 250 values obtained for each validation metric
        stochastic.validation <- rowMeans(stochastic.validation)</pre>
        # correlation is computed for the deterministic series of predicted_{f \sqcup}
 \rightarrow rainfall; recall E[X] = alpha/beta
        alphas <- station.predictions$predicted[ , 1]</pre>
        betas <- station.predictions$predicted[ , 2]</pre>
        correlation <- cor(alphas/betas, observed, method = "spearman")</pre>
        return(c(stochastic.validation, Cor.Det = correlation))
    })
validation <- t(simplify2array(validation))</pre>
# plotting the results
boxplot(validation)
abline(h = 1, lty = 2)
```

