Mapping Crimean-Congo Haemorrhagic Fever in Africa with BART

Getting Started

So you're interested in using embarcadero to do species distribution modeling with Bayesian additive regression trees! That's great. BARTs are a powerful way to do machine learning and, while not a new method per se, they are very new for SDMs.

In this advanced tutorial, I'm going to assume that you've seen the main paper and the embarcadero package vignette, which goes over basic functions and some of the internal structure of BART models. This tutorial, and the associated repo (cjcarlson/pier39), are intended for advanced users interested in seeing what a professional workflow might look like for an applied use.

```
library(embarcadero)
set.seed(12345)
```

Our goal here is to do a few things:

- 1. Build a species distribution model for *Hyalomma truncatum*, a tick we think might be a vector of Crimean-Congo haemorrhagic fever (CCHF) in Africa.
- 2. To build a transmission risk map, again using species distribution modeling/ecological niche modeling, for CCHF in Africa.
- 3. To see if we learn anything from that model about CCHF we didn't know before.
- 4. To see some of the advanced visualization and workflow tricks in the embarcadero package!

Mapping Hyalomma

We're going to make a suitability layer for *Hyalomma truncatum*, a possible CCHF vector. The tick occurrence data, which is a set of presence points, comes from Cumming et al. (1998) *Bulletin of Entomological Research*, one of the most detailed datasets in the world on parasite distributions. We're going to build a species distribution model using those points, some climate data, and a few other convenient layers that might be relevant.

Data entry

Let's start by loading in the climate data. We'll use three main sources:

- 1. WorldClim v1.4 (Hijmans et al. 2005), a harmonized dataset of bioclimatic variables mostly used for species distribution modeling
- 2. Two layers describing long-term averages of the normalized difference vegetation index (NDVI), which captures the greenness of a landscape (taken from previous use in Carlson et al. 2018). This can be an important variable for measuring invertebrate distributions, for example, or partitioning landscapes into different biomes.
- 3. A layer produced by the NASA SEDAC center describing the percent cropland by grid cell. If we think ruminants are spreading CCHF, this might matter! (https://sedac.ciesin.columbia.edu/data/set/aglands-croplands-2000)

Based on some expert opinion, I picked a handful of variables I thought might work well (and it's already reduced down to minimize redunandancy):

- BIO1: Mean annual temperature
- BIO2: Mean diurnal range
- BIO5: Max temperature of warmest month
- BIO6: Min temperature of coldest month
- BIO12: Mean annual precipitation
- BIO13: Precipitation of wettest month
- BIO14: Precipitation of driest month
- BIO15: Precipitation seasonality
- Mean NDVI (normalized difference vegetation index)
- NDVI amplitude
- Percent cropland by pixel

Let's read in the covariates.

```
load(file = "~/Github/pier39/covsraw.rda"); covs <- covsraw
class(covs)
#> [1] "RasterStack"
#> attr(, "package")
#> [1] "raster"
covs@crs <- crs('+proj=longlat +ellps=WGS84 +datum=WGS84 +no_defs +towgs84=0,0,0')</pre>
```

Next, let's load in our tick occurrence dataset, which is a set of longitude/latitude coordinates.

```
load(file = "~/Github/pier39/ticks.rda")
head(ticks)
#>
        X
              ScientificName Longitude.X Latitude.Y
#> 1 16471 Hyalomma truncatum
                                   16
#> 2 16472 Hyalomma truncatum
                                             -12.5
                                     14
#> 3 16473 Hyalomma truncatum
                                              -5.5
                                     12
#> 4 16474 Hyalomma truncatum
                                    15
                                             -16.5
#> 5 16475 Hyalomma truncatum
                                    14
                                             -13.6
#> 6 16476 Hyalomma truncatum
                                     15
                                             -12.4
nrow(ticks)
#> [1] 1794
```

Presence points are usually a little aggregated in space, so as a normal data cleaning practice, I like to thin my occurrence data to one point per raster grid cell.

```
# Make a spatial point data frame
mod <- SpatialPointsDataFrame(ticks[,3:4],data.frame(ticks[,1]))</pre>
names(mod@data) <- 'Presence'</pre>
nrow(mod)
#> [1] 1794
# Rasterizing and converting back to points makes one unique point per grid cell
tmp=rasterize(mod, covs[[1]], field="Presence", fun="min")
pts.sp1=rasterToPoints(tmp, fun=function(x){x>0})
nrow(pts.sp1)
#> [1] 1716
# Extract the climate data at each point
pres.cov <- raster::extract(covs, pts.sp1[,1:2])</pre>
head(pres.cov)
        bio1 bio12 bio13 bio14 bio15 bio2 bio5 bio6 crop ndvi.amp
#> [1,] 29 0.0064 0.0090 0.0042 0.15 23 47 8.0 0.000
#> [2,] 30 0.0061 0.0092 0.0037 0.17 23 48 8.7 0.000
                                                                0.056
```

```
#> [3,] 32 0.0090 0.0117 0.0064 0.16
                                       22 48 11.3 0.000
                                                            0.105
#> [4,]
         32 0.0085 0.0145 0.0036 0.38
                                       21
                                            47 14.7 0.031
                                                            0.032
#> [5,] 31 0.0081 0.0154 0.0028 0.45 21 46 12.4 0.011
                                                            0.014
#> [6,] 32 0.0087 0.0148 0.0037 0.38 21 47 14.9 0.044
                                                             0.151
#>
       ndvi.mean
#> [1,]
          -0.015
#> [2,]
           0.016
#> [3,]
           0.095
#> [4,]
           0.127
#> [5,]
           0.122
#> [6,]
          -0.039
```

Next, let's generate an equal number of pseudoabsences around Africa to the number of presences we have. BARTs are like BRTs in that they are sensitive to assumed prevalence; anecdotally, I strongly suggest using an equal number of presences and absences in your training data. You can experiment with the demo data by changing "nrow(ticks)" to "5000" below if you want to see some weirder model behavior.

```
#Generate the data
absence <- randomPoints(covs,nrow(ticks))</pre>
#> Warning in .couldBeLonLat(x, warnings = warnings): CRS is NA. Assuming it
#> is longitude/latitude
abs.cov <- raster::extract(covs, absence)</pre>
#Code the response
pres.cov <- data.frame(pres.cov); pres.cov$tick <- 1</pre>
abs.cov <- data.frame(abs.cov); abs.cov$tick <- 0
# And one to bind them
all.cov <- rbind(pres.cov, abs.cov)</pre>
head(all.cov)
     bio1 bio12 bio13 bio14 bio15 bio2 bio5 bio6 crop ndvi.amp ndvi.mean
      29 0.0064 0.0090 0.0042 0.15 23 47 8.0 0.000
                                                             0.050
                                                                      -0.015
      30 0.0061 0.0092 0.0037 0.17 23 48 8.7 0.000
                                                             0.056
                                                                       0.016
#> 3
      32 0.0090 0.0117 0.0064 0.16 22 48 11.3 0.000
                                                             0.105
                                                                       0.095
#> 4 32 0.0085 0.0145 0.0036 0.38 21 47 14.7 0.031
                                                             0.032
                                                                       0.127
#> 5
      31 0.0081 0.0154 0.0028 0.45
                                       21 46 12.4 0.011
                                                             0.014
                                                                       0.122
#> 6 32 0.0087 0.0148 0.0037 0.38
                                       21 47 14.9 0.044
                                                                      -0.039
                                                             0.151
#>
   tick
#> 1
       1
#> 2
        1
#> 3
       1
#> 4
       1
#> 5
        1
#> 6
# Let's just clean it up a little bit - remove any missing data
all.cov <- all.cov[complete.cases(all.cov),]</pre>
```

Now we have a dataset ready to do some modeling!

Running models with dbarts

We could try something really simple on defaults, right out the gate. The bart function in dbarts can just be run on defaults:

```
xvars <- names(all.cov)[!(names(all.cov)=='tick')]</pre>
xvars
#> [1] "bio1"
                    "bio12"
                                 "bio13"
                                             "bio14"
                                                         "bio15"
#> [6] "bio2"
                    "bio5"
                                 "bio6"
                                             "crop"
                                                        "ndvi.amp"
#> [11] "ndvi.mean"
first.model <- bart(all.cov[,xvars], all.cov[,'tick'], keeptrees=TRUE)</pre>
#> Running BART with binary y
#>
#> number of trees: 200
#> number of chains: 1, number of threads 1
#> Prior:
#> k: 2.000000
#> power and base for tree prior: 2.000000 0.950000
#> use quantiles for rule cut points: false
#> data:
#> number of training observations: 3478
#> number of test observations: 0
#> number of explanatory variables: 11
#>
\#> Cutoff rules c in x \le c vs x \ge c
#> Number of cutoffs: (var: number of possible c):
#> (1: 100) (2: 100) (3: 100) (4: 100) (5: 100)
#> (6: 100) (7: 100) (8: 100) (9: 100) (10: 100)
#> (11: 100)
#>
#> offsets:
#> reg : 0.00 0.00 0.00 0.00 0.00
#> Running mcmc loop:
#> iteration: 100 (of 1000)
#> iteration: 200 (of 1000)
#> iteration: 300 (of 1000)
#> iteration: 400 (of 1000)
#> iteration: 500 (of 1000)
#> iteration: 600 (of 1000)
#> iteration: 700 (of 1000)
#> iteration: 800 (of 1000)
#> iteration: 900 (of 1000)
#> iteration: 1000 (of 1000)
#> total seconds in loop: 10.667233
#> Tree sizes, last iteration:
#> [1] 2 2 2 2 3 3 2 3 2 2 2 2 3 3 4 3 3 2
#> 3 3 4 1 4 2 2 3 2 4 3 5 5 2 3 2 3 2 3 1
#> 3 2 1 3 2 2 3 3 3 3 3 4 2 2 2 3 2 2 2 2
#> 2 3 1 2 2 4 2 2 2 2 1 2 3 3 2 3 2 3 2 2
#> 3 2 2 1 2 2 3 2 3 2 4 2 2 2 2 2 3 1 2 3
#> 4 1 3 2 1 2 2 2 3 2 2 2 3 2 4 2 2 2 3 4
#> 3 2 2 2 2 3 2 3 2 4 2 3 2 4 2 2 3 3 2 2
#> 2 2 2 3 2 3 2 1 2 2 3 2 3 2 2 4 3 2 3 4
#> 2 3 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 3 2 2 2 2 4
#> 3 2 2 3 2 2 3 2 2 2 3 3 3 3 3 2 2 2 3 1
#> 2 3
```

```
#> Variable Usage, last iteration (var:count):

#> (1: 32) (2: 30) (3: 24) (4: 22) (5: 34)

#> (6: 18) (7: 27) (8: 30) (9: 22) (10: 18)

#> (11: 28)

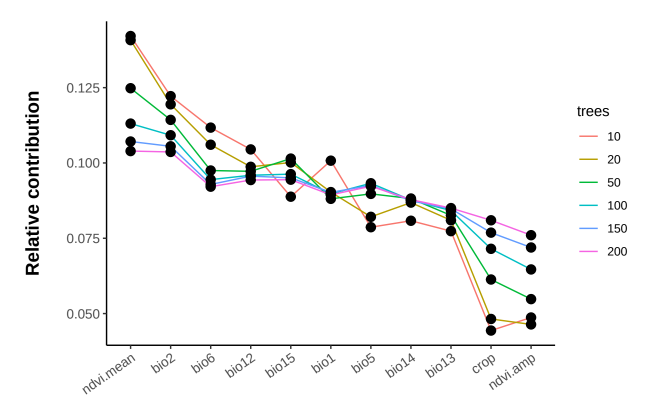
#> DONE BART
```

In reality, we know we want to do variable set reduction, and tune the model a bit from there. In the main vignette you can see the different component parts of that process. Here, we're going to use our most powerful shortcut: bart.step. Inside bart.step, five things happen:

- 1. The variable importance diagnostic from varimp.diag is generated
- 2. The stepwise variable set reduction in variable.step is run
- 3. The final model is run with the reduced predictor set
- 4. The variable importance plot from varimp is generated
- 5. The model summary from summary is returned

This is slow but very easy to simply fire off if you're running a large workflow (you can speed it up, in a demo, by reducing iter.step and even more so by reducing iter.plot). Let's do it:

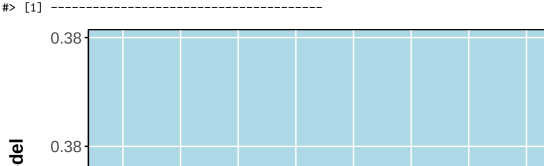
```
sdm <- bart.step(x.data = all.cov[,xvars],</pre>
                  y.data = all.cov[,'tick'],
                 full = TRUE,
                 quiet = TRUE)
#>
#>
    10 tree models: 100 iterations
#>
#>
    20 tree models: 100 iterations
#>
#>
    50 tree models: 100 iterations
#>
    100 tree models: 100 iterations
#>
#>
    150 tree models: 100 iterations
#>
#>
   200 tree models: 100 iterations
```

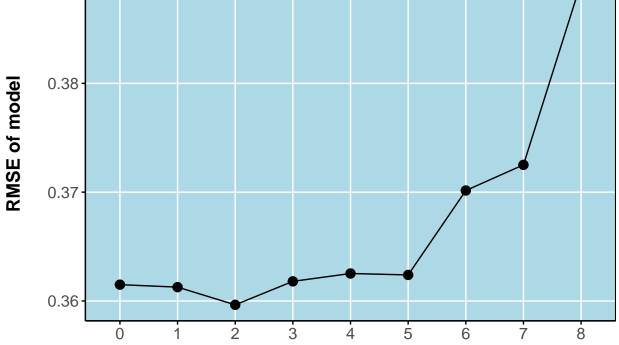


Variables dropped

#> [1] Number of variables included: 11 #> [1] Dropped: #> [1] **#>** [1] -----#> [1] Number of variables included: 10 #> [1] Dropped: #> [1] crop **#>** [1] -----#> [1] Number of variables included: 9 #> [1] Dropped: #> [1] crop ndvi.amp #> [1] -----#> [1] Number of variables included: 8 #> [1] Dropped: #> [1] crop ndvi.amp bio5 #> [1] -----#> [1] Number of variables included: 7 #> [1] Dropped: #> [1] crop ndvi.amp bio5 bio14 #> [1] -----#> [1] Number of variables included: 6 #> [1] Dropped: #> [1] -----#> [1] Number of variables included: 5 #> [1] Dropped:

```
#> [1] crop
          ndvi.amp bio5 bio14 bio13
#> [1] -----
#> [1] Number of variables included: 4
#> [1] Dropped:
#> [1] crop
          ndvi.amp bio5 bio14
                               bio13
                                      bio15
                                             bio6
#> [1] -----
#> [1] Number of variables included: 3
#> [1] Dropped:
#> [1] crop
         ndvi.amp bio5 bio14 bio13
                                      bio15
                                             bio6
                                                    bio2
```

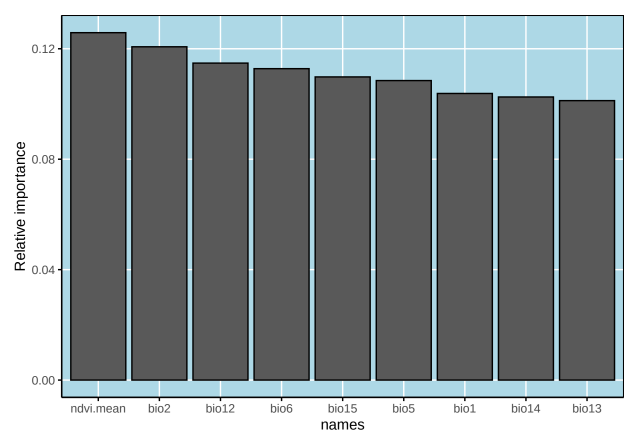




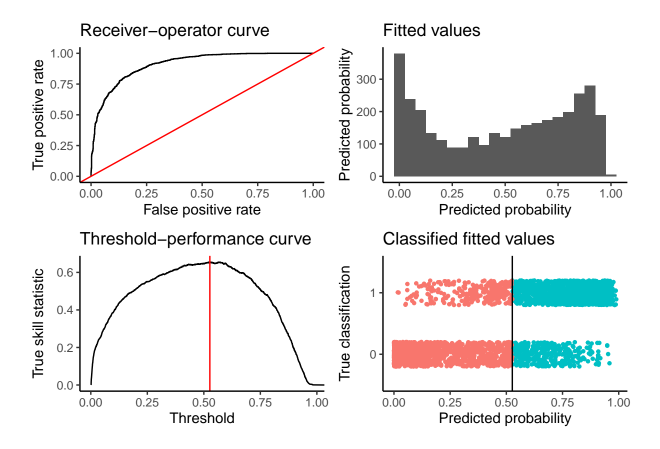
Variables dropped

```
#> [1] -----
#> [1] Final recommended variable list
#> [1] bio1
               bio12
                        bio13 bio14 bio15 bio2
                                                            bio5
#> [8] bio6
               ndvi.mean
#>
#> Running BART with binary y
#> number of trees: 200
#> number of chains: 1, number of threads 1
#> Prior:
#> k: 2.000000
#> power and base for tree prior: 2.000000 0.950000
#> use quantiles for rule cut points: false
#> number of training observations: 3478
#> number of test observations: 0
```

```
#> number of explanatory variables: 9
#>
#> Cutoff rules c in x<=c vs x>c
#> Number of cutoffs: (var: number of possible c):
#> (1: 100) (2: 100) (3: 100) (4: 100) (5: 100)
#> (6: 100) (7: 100) (8: 100) (9: 100)
#>
#> offsets:
#> reg : 0.00 0.00 0.00 0.00 0.00
#> Running mcmc loop:
#> iteration: 100 (of 1000)
#> iteration: 200 (of 1000)
#> iteration: 300 (of 1000)
#> iteration: 400 (of 1000)
#> iteration: 500 (of 1000)
#> iteration: 600 (of 1000)
#> iteration: 700 (of 1000)
#> iteration: 800 (of 1000)
#> iteration: 900 (of 1000)
#> iteration: 1000 (of 1000)
#> total seconds in loop: 8.798700
#> Tree sizes, last iteration:
#> [1] 2 2 2 2 2 3 2 2 1 2 3 1 3 3 3 2 2 2
#> 2 2 4 2 2 4 3 2 2 2 4 2 3 2 2 3 3 2 2 3
#> 2 2 2 3 3 3 3 1 2 4 2 3 3 2 2 2 3 2 2 2
#> 3 2 3 2 2 2 3 2 2 3 2 3 2 2 2 2 2 2 3 3
#> 2 2 3 2 3 2 4 2 3 2 3 3 2 2 2 3 3 3 3
#> 3 3 2 6 2 1 2 2 2 2 2 2 2 3 1 3 2 1 2 4
#> 3 3 2 2 2 3 2 2 3 3 2 2 2 2 3 2 2 3 2
#> 3 2 4 3 2 2 3 2 2 3 3 3 5 2 2 3 2 2 2 2
#> 2 2 5 3 3 2 4 4 2 2 4 2 3 3 2 2 2 2 2 2
#> 2 2 1 2 2 3 2 2 4 3 2 2 2 3 3 2 2 2 2 2
#> 4 3
#> Variable Usage, last iteration (var:count):
#> (1: 31) (2: 37) (3: 30) (4: 27) (5: 33)
#> (6: 33) (7: 26) (8: 30) (9: 39)
#> DONE BART
```



```
#> Call: bart x.data[, vs] y.data TRUE
#>
#> Predictor list:
#> bio1 bio12 bio13 bio14 bio15 bio2 bio5 bio6 ndvi.mean
#>
#> Area under the receiver-operator curve
#> AUC = 0.91
#>
#> Recommended threshold (maximizes true skill statistic)
#> Cutoff = 0.53
#> TSS = 0.66
#> Resulting type I error rate: 0.16
#> Resulting type II error rate: 0.19
```

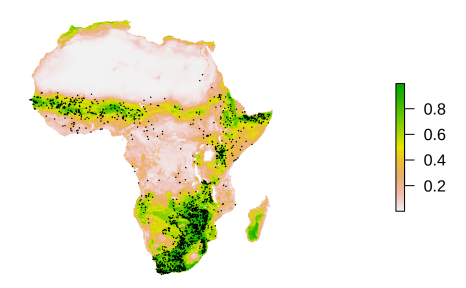


Out of all the variables, "crop" seems to be especially undesirable as a predictor. A few other variables seem like they might not be helping the model either, but we probably need a systematic way to deal with that.

A high AUC value indicates our model performs well. The summary function also returns an optimal threshold that maximizes the true skill statistic (TSS), and the sensitivity/specificity of the model at that cutoff (alpha).

What do the predictions look like? To make a predicted raster, we have to use embarcadero's wrapper for the native predict function in dbarts.

Hyalomma truncatum



This model seems okay. We're getting predictions in places we don't have any records, like North Africa. That could be good if we think that's suitable climatic space (and if you know *Hyalomma*, you know there's definitely some species there, though posibly not truncatum), but with much of the inhabited area not being predicted, let's revisit that later.

Mapping CCHF

Alright. Now let's get back to business by building the CCHF map. We're going to use the same predictors as we used for *H. truncatum*, plus the suitability layer for the ticks. (Using one SDM layer in another map is a sort of finnicky business - sometimes it improves predictions and is epistemologically valid, other times it predetermines a certain outcome in the final model, given issues with colinearity. That's a broader discussion than what we're doing here! If the final map predicts some areas that aren't where the ticks are, I'll call that a win. If they were identical, I would worry.)

Running the CCHF model

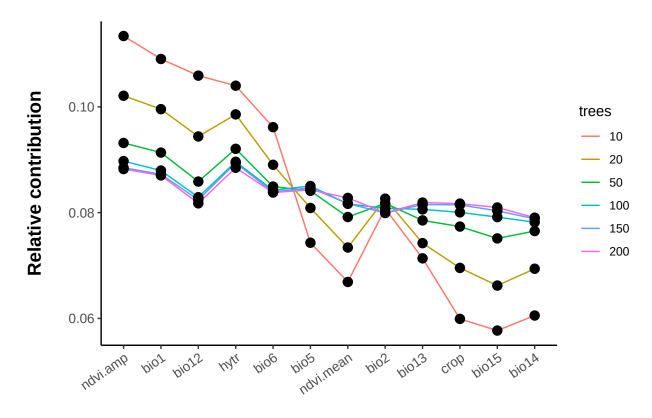
This time, let's just do the variable selection up front. Same plan as before:

```
# Update the covariate stack

covs <- stack(covs, hytr.layer)
names(covs)[12]='hytr' # for Hyalomma truncatum
xvars <- c(xvars, 'hytr')</pre>
```

```
# Read in the data
load(file = "~/Github/pier39/cchf.rda")
head(cchf)
#> OCCURRENCE ID LOCATION TYPE ADMIN LEVEL GAUL AD1 GAUL AD2
#> 1
               1
                        point
                               -999 1282
                                                   16397
#> 2
               2
                       point
                                   -999
                                            1282
                                                    16397
#> 3
               3
                                   -999
                                            1282 16397
                       point
#> 4
               4
                                   -999
                                            1278 16376
                        point
#> 5
               5
                                    -999
                                            1278
                        point
                                                    16376
                        point
                                                    16376
#> 6
               6
                                    -999
                                            1278
#> UNIQUE_LOCATION YEAR LATITUDE LONGITUDE COUNTRY REGION
#> 1
              535 1953 38
                                   69 Tajikistan Asia
                           38
#> 2
              1178 1953
                                      70 Tajikistan
                                                     Asia
                          42
#> 3
              620 1954
                                     21
                                            Serbia Asia
#> 4
             1182 1954
                           37
                                    69 Tajikistan Asia
                                69 Tajikistan Asia
70 Tajikistan Asia
#> 5
              1165 1954
                           37
#> 6
              1178 1954
                           38
nrow(cchf)
#> [1] 1721
# Spatial thinning checks; this also limits it to African points
# (CCHF is found elsewhere in the world too,
# and our data includes other records!)
cchf <- cchf[,c('LONGITUDE','LATITUDE')]; cchf$Presence = 1</pre>
cchf <- SpatialPointsDataFrame(cchf[,1:2],data.frame(Presence=cchf[,3]))</pre>
tmp=rasterize(cchf, covs[[1]], field="Presence", fun="min")
pts.sp1=rasterToPoints(tmp, fun=function(x){x>0})
nrow(pts.sp1)
#> [1] 147
# Extract presence values
pres.cov <- raster::extract(covs, pts.sp1[,1:2])</pre>
pres.cov <- na.omit(pres.cov)</pre>
head(pres.cov)
      bio1 bio12 bio13 bio14 bio15 bio2 bio5 bio6 crop ndvi.amp
#> [1,] 31 0.0076 0.011 0.0044 0.20 23 48 9.9 0.046 0.183
#> [2,] 31 0.0076 0.011 0.0044 0.21 23 48 9.9 0.009
                                                          0.101
#> [3,] 29 0.0106 0.016 0.0070 0.30 15 41 16.9 0.000
                                                         0.031
#> [4,] 31 0.0098 0.015 0.0053 0.33 21 46 14.7 0.036
                                                       0.019
#> [5,] 32 0.0085 0.014 0.0036 0.38 21 47 14.5 0.002
                                                        0.023
#> [6,] 31 0.0082 0.015 0.0032 0.43 20 47 13.3 0.048
                                                          0.025
#> ndvi.mean hytr
#> [1,]
         0.033 0.021
#> [2,]
         0.023 0.034
#> [3,]
         0.069 0.177
#> [4,] 0.133 0.381
#> [5,] 0.124 0.126
#> [6,] 0.133 0.187
#Generate pseudoabsences
```

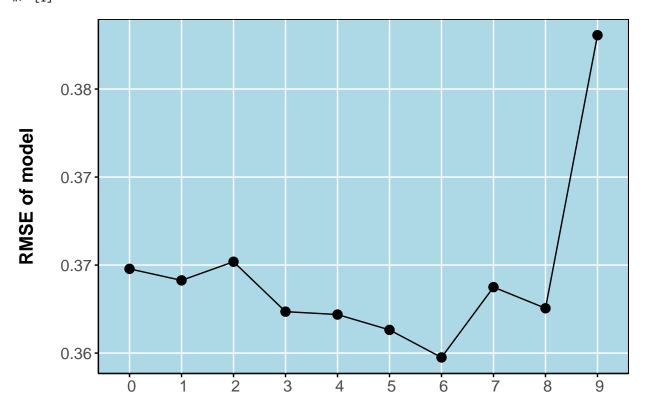
```
absence <- randomPoints(covs,nrow(pres.cov))</pre>
#> Warning in .couldBeLonLat(x, warnings = warnings): CRS is NA. Assuming it
#> is longitude/latitude
abs.cov <- raster::extract(covs, absence)</pre>
#Code the response
pres.cov <- data.frame(pres.cov); pres.cov$cchf <- 1</pre>
abs.cov <- data.frame(abs.cov); abs.cov$cchf <- 0
# And one to bind them
all.cov <- rbind(pres.cov, abs.cov)</pre>
all.cov <- all.cov[complete.cases(all.cov),]; nrow(all.cov)</pre>
#> [1] 184
head(all.cov)
#> bio1 bio12 bio13 bio14 bio15 bio2 bio5 bio6 crop ndvi.amp ndvi.mean
#> 1 31 0.0076 0.011 0.0044 0.20 23 48 9.9 0.046
                                                        0.183
                                                                  0.033
#> 2 31 0.0076 0.011 0.0044 0.21 23 48 9.9 0.009
                                                        0.101
                                                                  0.023
0.031
                                                                  0.069
#> 4 31 0.0098 0.015 0.0053 0.33 21 46 14.7 0.036
                                                        0.019
                                                                  0.133
#> 5 32 0.0085 0.014 0.0036 0.38 21 47 14.5 0.002 0.023
                                                                  0.124
#> 6 31 0.0082 0.015 0.0032 0.43 20 47 13.3 0.048 0.025
                                                                  0.133
#> hytr cchf
#> 1 0.021
#> 2 0.034
#> 3 0.177
           1
#> 4 0.381 1
#> 5 0.126
            1
#> 6 0.187
# This part automates the variable selection and returns the model
cchf.model <- bart.step(x.data = all.cov[,xvars],</pre>
                      y.data = all.cov[,'cchf'],
                      full = TRUE,
                      quiet = TRUE)
#>
#> 10 tree models: 100 iterations
#>
#> 20 tree models: 100 iterations
#>
#> 50 tree models: 100 iterations
#> 100 tree models: 100 iterations
#>
#> 150 tree models: 100 iterations
#>
#> 200 tree models: 100 iterations
```



Variables dropped

#> [1] Number of variables included: 12 #> [1] Dropped: #> [1] **#>** [1] -----#> [1] Number of variables included: 11 #> [1] Dropped: #> [1] bio15 **#>** [1] -----#> [1] Number of variables included: 10 #> [1] Dropped: #> [1] bio15 crop **#>** [1] -----#> [1] Number of variables included: 9 #> [1] Dropped: #> [1] bio15 crop bio14 **#>** [1] -----#> [1] Number of variables included: 8 #> [1] Dropped: crop bio14 ndvi.mean #> [1] bio15 #> [1] -----#> [1] Number of variables included: 7 #> [1] Dropped: #> [1] bio15 crop bio14 ndvi.mean bio13 **#>** [1] -----#> [1] Number of variables included: 6 #> [1] Dropped:

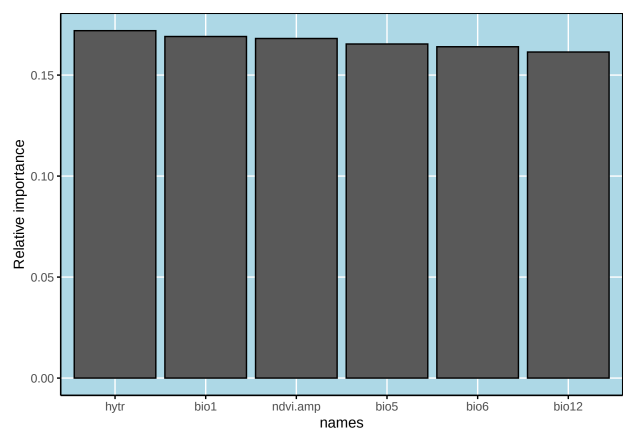
```
#> [1] bio15
          crop bio14 ndvi.mean bio13 bio2
#> [1] -----
#> [1] Number of variables included: 5
#> [1] Dropped:
#> [1] bio15
            crop
                 bio14 ndvi.mean bio13 bio2
                                                 bio5
#> [1] -----
#> [1] Number of variables included: 4
#> [1] Dropped:
#> [1] bio15
          crop
                   bio14 ndvi.mean bio13
                                        bio2
                                                 bio5
#> [8] hytr
#> [1] -----
#> [1] Number of variables included: 3
#> [1] Dropped:
#> [1] bio15
                   bio14 ndvi.mean bio13
                                       bio2
            crop
                                                 bio5
#> [8] hytr
            bio1
#> [1] -----
```



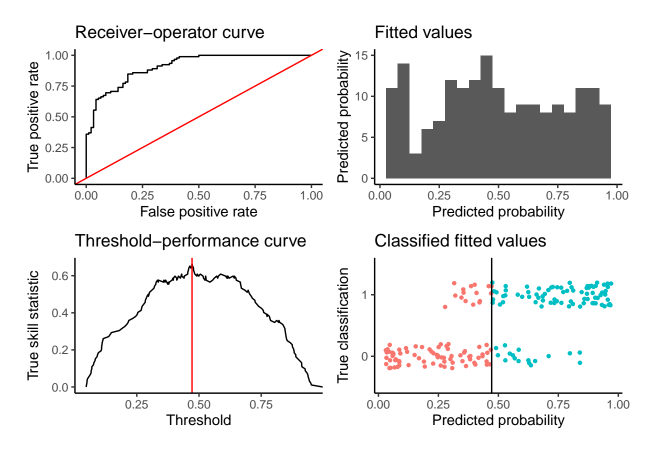
Variables dropped

```
#> [1] ------
#> [1] Final recommended variable list
#> [1] bio1 bio12 bio5 bio6 ndvi.amp hytr
#>
#> Running BART with binary y
#>
#> number of trees: 200
#> number of chains: 1, number of threads 1
#> Prior:
#> k: 2.000000
```

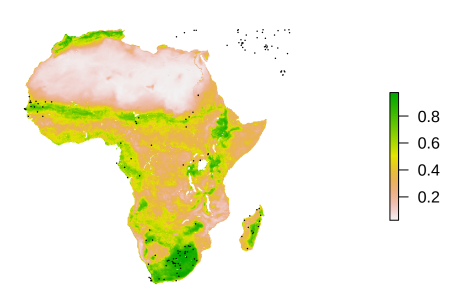
```
#> power and base for tree prior: 2.000000 0.950000
#> use quantiles for rule cut points: false
#> number of training observations: 184
#> number of test observations: 0
#> number of explanatory variables: 6
#> Cutoff rules c in x<=c vs x>c
#> Number of cutoffs: (var: number of possible c):
#> (1: 100) (2: 100) (3: 100) (4: 100) (5: 100)
#> (6: 100)
#>
#> offsets:
#> reg : 0.00 0.00 0.00 0.00 0.00
#> Running mcmc loop:
#> iteration: 100 (of 1000)
#> iteration: 200 (of 1000)
#> iteration: 300 (of 1000)
#> iteration: 400 (of 1000)
#> iteration: 500 (of 1000)
#> iteration: 600 (of 1000)
#> iteration: 700 (of 1000)
#> iteration: 800 (of 1000)
#> iteration: 900 (of 1000)
#> iteration: 1000 (of 1000)
#> total seconds in loop: 1.149624
#> Tree sizes, last iteration:
#> [1] 4 2 3 2 2 2 2 2 2 2 6 2 3 2 1 2 3 2
#> 2 2 1 2 5 3 2 4 2 2 3 3 3 2 2 2 4 3 2 3
#> 1 2 4 4 1 4 3 2 2 3 4 3 3 2 3 2 3 2 2 1
#> 2 3 2 2 2 2 3 3 6 2 3 2 2 2 2 2 4 2 4 2
#> 2 2 3 3 4 2 4 2 2 2 3 2 2 2 2 2 2 4 2 2
#> 3 3 3 4 1 2 2 2 2 1 2 3 2 3 3 2 2 2 2 1
#> 3 4 3 2 2 3 2 2 2 3 2 4 2 3 2 2 2 2 3
#> 2 2 2 2 2 2 2 2 2 2 2 2 2 3 1 1 3 3 3
#> 2 2 2 2 4 2 2 3 3 2 4 2 2 2 2 5 4 3 2 2
#> 3 2
#>
#> Variable Usage, last iteration (var:count):
#> (1: 50) (2: 50) (3: 54) (4: 39) (5: 46)
#> (6: 44)
#> DONE BART
```



```
#> Call: bart x.data[, vs] y.data TRUE
#>
#> Predictor list:
#> bio1 bio12 bio5 bio6 ndvi.amp hytr
#>
#> Area under the receiver-operator curve
#> AUC = 0.91
#>
#> Recommended threshold (maximizes true skill statistic)
#> Cutoff = 0.47
#> TSS = 0.66
#> Resulting type I error rate: 0.15
#> Resulting type II error rate: 0.18
```







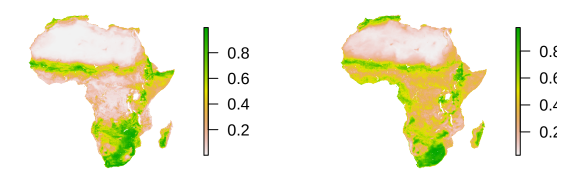
OK. Nice model! Let's see what we can do to unpack it.

First, let's compare it against the tick map.

```
par(mfrow=c(1,2))
plot(hytr.layer, box=FALSE, axes=FALSE, main='H. truncatum')
plot(cchf.map[[1]], box=FALSE, axes=FALSE, main='CCHF')
```

H. truncatum

CCHF



Our model seems to be different from the previous one (published by Messina et al., using this dataset which they generously provide online) in three major ways.

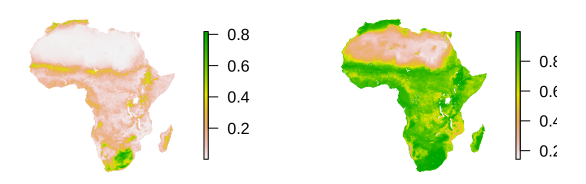
- 1. First, using the tick vector has increased the amount of predicted suitable area in South Africa, Namibia, Botswana, and Zimbabwe. That makes sense, overall—if there are vectors present, CCHF seems plausible.
- 2. The model predicts the area in coastal Cameroon, Gabon, and Equatorial Guinea that we know has some CCHF records but has previously been underpredicted. Weirdly, we don't have good evidence *Hyalomma truncatum* is there. So, the model is doing well, but the ecology is still unclear.
- 3. Finally, the northern coast of Africa is predicted to be highly suitable. There's plenty of *Hyalomma* species up there, though not *H. truncatum* as far as our data suggests. It's possible we should think more about the possibility of CCHF in Morocco and Algeria.

Let's take a look at the uncertainty in the model:

```
par(mfrow=c(1,2))
plot(cchf.map[[2]], box=FALSE, axes=FALSE, main='2.5% bound')
plot(cchf.map[[3]], box=FALSE, axes=FALSE, main='97.5% bound')
```

2.5% bound

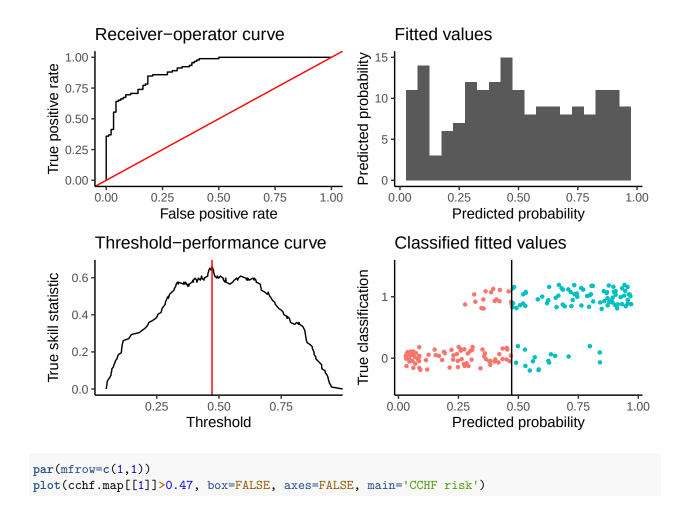
97.5% bound



This may be more interesting to think about in the context of binary maps of presence/absence. We can produce those using summary:

```
summary(cchf.model)
#> Call: bart x.data[, vs] y.data TRUE

#>
#> Predictor list:
#> bio1 bio12 bio5 bio6 ndvi.amp hytr
#>
#> Area under the receiver-operator curve
#> AUC = 0.91
#>
#> Recommended threshold (maximizes true skill statistic)
#> Cutoff = 0.47
#> TSS = 0.66
#> Resulting type I error rate: 0.15
#> Resulting type II error rate: 0.18
```



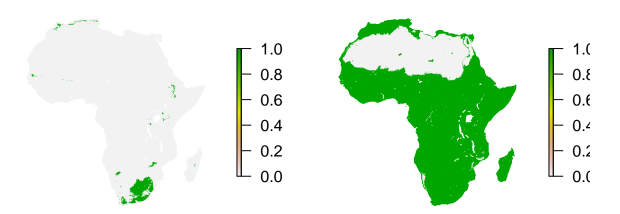
CCHF risk



```
par(mfrow=c(1,2))
plot(cchf.map[[2]]>0.47, box=FALSE, axes=FALSE, main='2.5% bound')
plot(cchf.map[[3]]>0.47, box=FALSE, axes=FALSE, main='97.5% bound')
```

2.5% bound

97.5% bound

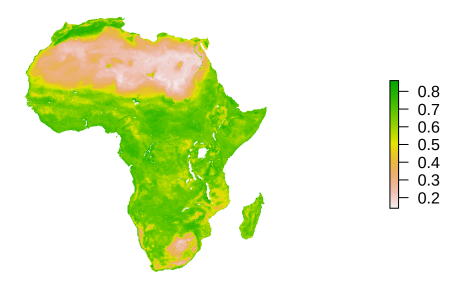


The outer bounds here are *incredibly* wide - which highlights exactly how little data we have about the distribution of CCHF, and how much uncertainty presentation matters.

Where is the uncertainty highest?

```
par(mfrow=c(1,1))
plot(cchf.map[[3]]-cchf.map[[2]],
    box=FALSE, axes=FALSE, main='Posterior width, scaled')
```

Posterior width, scaled

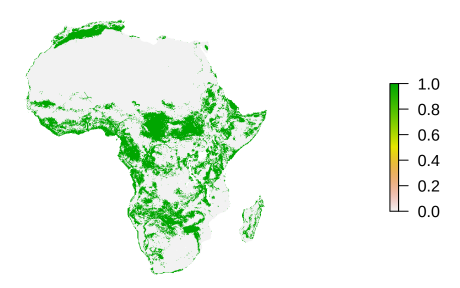


Posterior width isn't always the most informative measure of uncertainty. But it does tell us we're most confident in the Sahara, where we're quite sure there's no real CCHF risk, and in the parts of South Africa where spillover is most common. We can further investigate by mapping the places with the most uncertainty:

```
quant <- quantile(values(cchf.map[[3]]-cchf.map[[2]]), 0.75, na.rm=TRUE)

par(mfrow=c(1,1))
plot((cchf.map[[3]]-cchf.map[[2]])>quant,
    box=FALSE, axes=FALSE,
    main = "Highest uncertainty zones")
```

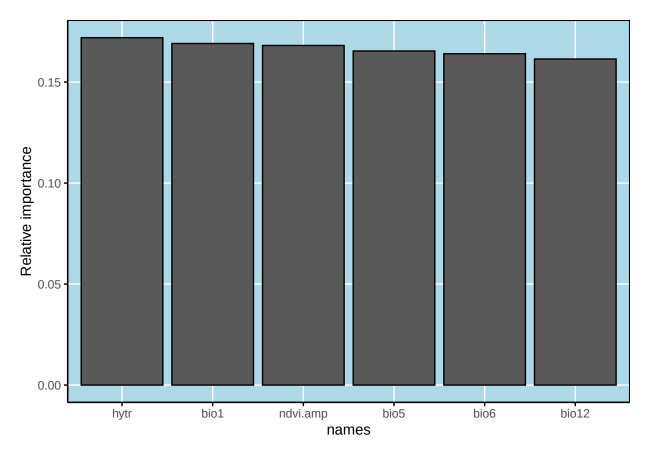
Highest uncertainty zones



The most interesting takehome is that uncertainty is particularly high in the Sahel and on the Western coast of Africa; along the equator, CCHF spillover has previously occurred but the tick vector is absent. This may be worth further investigation. Similarly, there are no spillovers along the southern coast of West Africa, but uncertainty is very high, and it may be worth investigating this more (especially given tick records in that region).

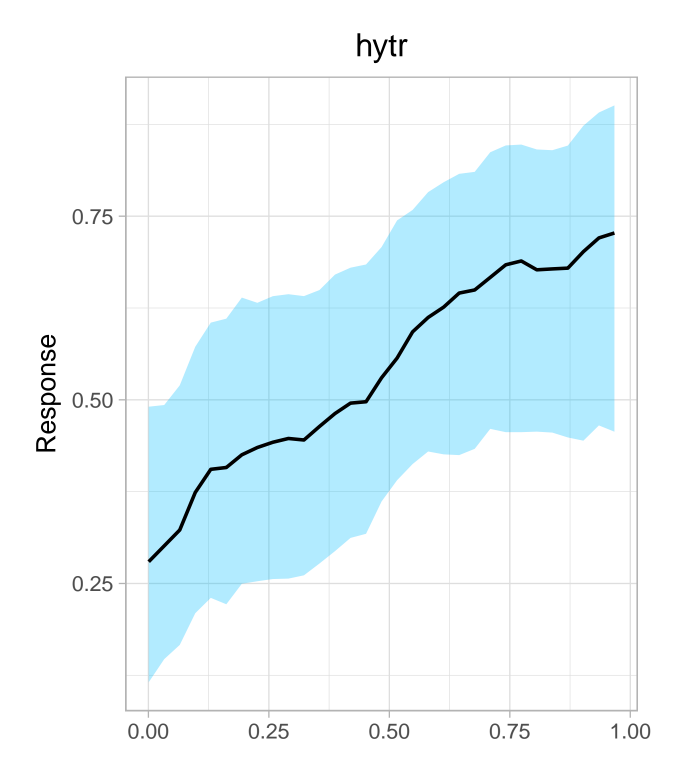
Analytics

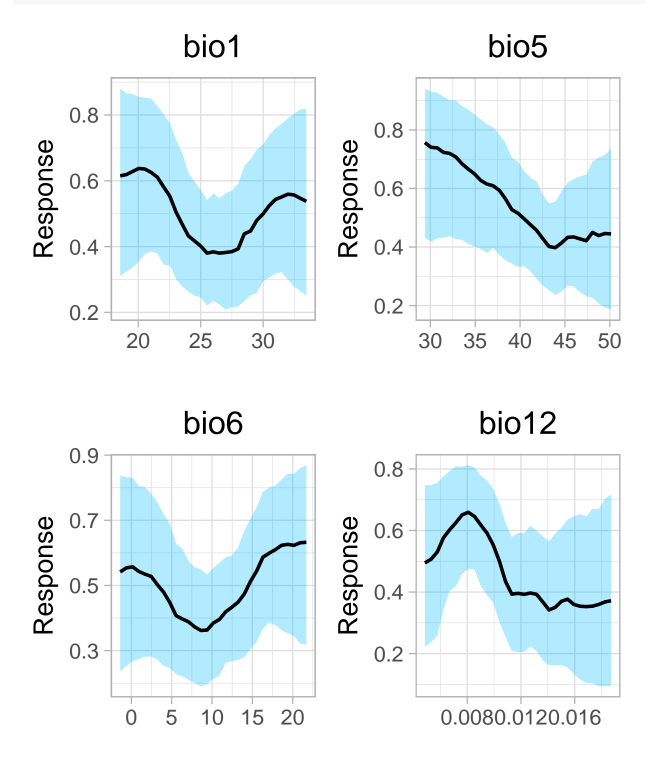
Finally, let's unpack some of what's under the hood in the model. First let's look at the variable contributions: varimp(cchf.model, plots=TRUE)



```
#>
        names varimps
#> 1
         bio1
                  0.17
                  0.16
#> 2
        bio12
#> 3
         bio5
                  0.17
                  0.16
#> 4
         bio6
#> 5 ndvi.amp
                  0.17
#> 6
                  0.17
         hytr
```

The tick vector comes out on top, unsurprisingly! Let's look at the response functions a little bit. First, let's look at the partial dependence plots for a couple individual variables.



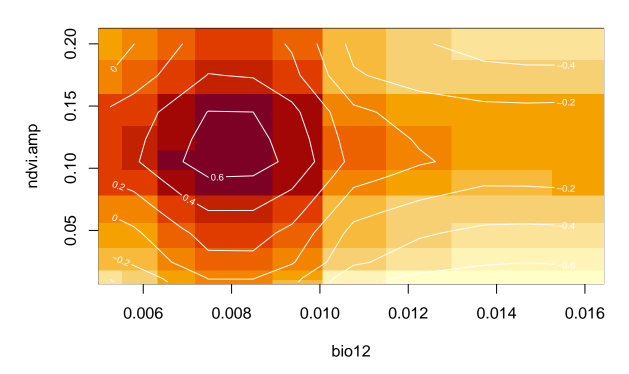


Some of these patterns are pretty clear - suitability declines with the maximum temperature of the warmest month, and increases with the probability of the tick.

A great feature of BART is that we can pretty easily do two-dimensional partial dependence plots, and we can pretty easily visualize the optimum (I haven't added a wrapper for this yet):

```
p <- pd2bart(cchf.model, xind=c('bio12', 'ndvi.amp'), pl=TRUE)</pre>
```

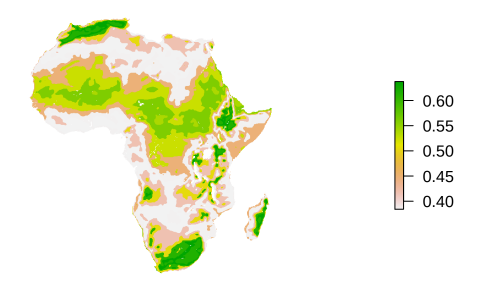
Median



This is probably my favorite feature of dbarts - these plots are a really nice way to visualize the Hutchinsonian niche sort of like how NicheA does, but within the familiar framework of classification trees. Plus it's not just a cross-product of the individual partials - the BART framework allows for interactions and sometimes you'll see them show up in these plots.

One last cool trick: spatial partial dependence plots

```
spartial(cchf.model, covs, x.vars = 'bio1', equal=TRUE)
```



#> class : RasterStack

#> dimensions : 867, 999, 866133, 1 (nrow, ncol, ncell, nlayers)

#> resolution : 0.083, 0.083 (x, y)

#> extent : -25, 58, -35, 37 (xmin, xmax, ymin, ymax)

#> crs : NA
#> names : bio1
#> min values : 0.38
#> max values : 0.64

This map projects the partial dependence plot onto the raster data for the bio1 layer, and can be interpreted as answering the question "Where are mean temperatures most conducive for viral transmission?"

Finally...

A big thanks to Jane Messina et al. for publicly sharing their CCHF data, and to Graeme Cumming for tick data.

References

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