5.Gene_expression_analysis

Fay

2022-08-09

load libraries

```
library(ggplot2)
library(tidyverse)
## -- Attaching packages -----
                                             ----- tidyverse 1.3.1 --
## v tibble 3.1.8
                 v dplyr
                          1.0.9
## v tidyr 1.2.0
                 v stringr 1.4.0
         2.1.2
                  v forcats 0.5.1
## v readr
## v purrr
         0.3.4
## -- Conflicts -----
                                  ## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                 masks stats::lag()
library(optimx)
```

Import data:

Here, we have the experimental / field data, inlcuding imputed data

```
lab <- read.csv("output_data/gene_expression/data_products/lab_imputed_gene_expression.csv")
field <- read.csv("output_data/gene_expression/data_products/field_imputed_gene_expression.csv")</pre>
```

Selecting genes

It is time to apply the package of Alice Balard et al. on our predictions!

Let's see if we indeed have differences across the hybrid index with our predicted weight loss.

Install the package

```
##
## * checking for file '/tmp/RtmpAt7KCX/remotes3475b43382dd1d/alicebalard-parasiteLoad-1b43216/DESCRIPT
## * preparing 'parasiteLoad':
## * checking DESCRIPTION meta-information ... OK
## * checking for LF line-endings in source and make files and shell scripts
## * checking for empty or unneeded directories
## * building 'parasiteLoad_0.1.0.tar.gz'
```

Applying Alice's package on every gene

```
x <- field$ID01
# Define function to be used to test, get the log lik and aic
tryDistrib <- function(x, distrib){</pre>
  # deals with fitdistr error:
 fit <- tryCatch(MASS::fitdistr(x, distrib), error=function(err) "fit failed")</pre>
  return(list(fit = fit,
              loglik = tryCatch(fit$loglik, error=function(err) "no loglik computed"),
              AIC = tryCatch(fit$aic, error=function(err) "no aic computed")))
}
findGoodDist <- function(x, distribs, distribs2){</pre>
  l =lapply(distribs, function(i) tryDistrib(x, i))
  names(1) <- distribs
  print(1)
  listDistr <- lapply(distribs2, function(i){</pre>
    if (i %in% "t"){
      fitdistrplus::fitdist(x, i, start = list(df =2))
    } else {
      fitdistrplus::fitdist(x,i)
    }}
  par(mfrow=c(2,2))
  denscomp(listDistr, legendtext=distribs2)
  cdfcomp(listDistr, legendtext=distribs2)
  qqcomp(listDistr, legendtext=distribs2)
  ppcomp(listDistr, legendtext=distribs2)
  par(mfrow=c(1,1))
tryDistrib(x, "normal")
```

Functions for testing distributions

```
## $fit
##
         mean
                       sd
                   4.4097438
##
     15.0575257
## ( 0.2409300) ( 0.1703633)
##
## $loglik
## [1] -972.423
##
## $AIC
## NULL
tryDistrib(x, "binomial")
## $fit
## [1] "fit failed"
##
## $loglik
## [1] "no loglik computed"
##
## $AIC
## [1] "no aic computed"
tryDistrib(x, "student")
## $fit
## [1] "fit failed"
## $loglik
## [1] "no loglik computed"
##
## $AIC
## [1] "no aic computed"
tryDistrib(x, "weibull")
## Warning in densfun(x, parm[1], parm[2], ...): NaNs produced
## Warning in densfun(x, parm[1], parm[2], ...): NaNs produced
## $fit
##
        shape
                     scale
##
      3.5549795
                 16.6979975
## ( 0.1421786) ( 0.2724857)
## $loglik
## [1] -976.5934
##
## $AIC
## NULL
tryDistrib(x, "weibullshifted")
## $fit
## [1] "fit failed"
##
## $loglik
```

```
## [1] "no loglik computed"
##
## $AIC
## [1] "no aic computed"
field$Sex <- as.factor(field$Sex)</pre>
speparam <- parasiteLoad::getParamBounds("weibull", data = field, response = "IFNy")</pre>
IFNy <- parasiteLoad::analyse(data = field,</pre>
                        response = "IFNy",
                        model = "weibull",
                        group = "Sex")
## [1] "Analysing data for response: IFNy"
## [1] "Fit for the response: IFNy"
## [1] "Fitting for all"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupA : F"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
```

```
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
                pvalue
##
      dLL dDF
## 1 0.67
           1 0.2464973
## [1] "Testing H1 no alpha vs alpha"
##
     dLL dDF
                 pvalue
## 1 0.46
           1 0.3364279
## [1] "Testing H2 groupA no alpha vs alpha"
   dLL dDF
                pvalue
          1 0.9640624
## 1 0
## [1] "Testing H2 groupB no alpha vs alpha"
     dLL dDF
                  pvalue
## 1 1.64
          1 0.06993746
## [1] "Testing H3 groupA no alpha vs alpha"
     dLL dDF
                 pvalue
## 1 0.01
            1 0.9198046
## [1] "Testing H3 groupB no alpha vs alpha"
     dLL dDF
                pvalue
## 1 1.25
           1 0.1142013
## [1] "Testing H1 vs H0"
     dLL dDF
                pvalue
## 1 0.14
           1 0.5927666
## [1] "Testing H2 vs H0"
##
     dLL dDF
                 pvalue
## 1 1.51
            3 0.3873923
## [1] "Testing H3 vs H1"
     dLL dDF
                 pvalue
## 1 2.21
           4 0.3529437
## [1] "Testing H3 vs H2"
     dLL dDF
                pvalue
## 1 0.84
            2 0.4334773
##A11
print(IFNy)
```

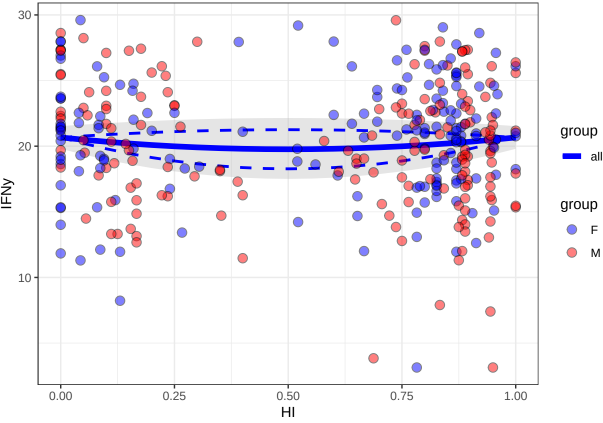
\$HO

```
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
   Coefficients:
##
            L1
                     alpha
                               myshape
               0.08694602
                            5.00000000
##
  20.66436487
##
## Log-likelihood: -985.61
## Best method: bobyqa
##
## $H1
##
## Call:
  bbmle::mle2(minuslogl = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
       control = config$control)
##
##
  Coefficients:
##
                        L2
                                  alpha
                                            myshape
  20.80840234 20.45982142 0.07573465 5.00000000
## Log-likelihood: -985.47
## Best method: bobyga
## $H2
## $H2$groupA
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
##
  Coefficients:
##
                       alpha
                                  myshape
  20.397246611 -0.004673532 5.000000000
##
## Log-likelihood: -483.7
## Best method: bobyga
```

```
##
## $H2$groupB
##
## Call:
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
  Coefficients:
##
           L1
                   alpha
                             myshape
## 20.9652515 0.2076485
                          4.8276501
##
## Log-likelihood: -500.4
## Best method: bobyqa
##
##
## $H3
## $H3$groupA
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
  Coefficients:
##
                                  alpha
##
                        L2
                                            myshape
  20.26647283 20.62522076
                            0.01117735 5.00000000
##
##
## Log-likelihood: -483.63
## Best method: bobyqa
##
## $H3$groupB
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
## Coefficients:
##
           L1
                      L2
                              alpha
                                        myshape
```

Scale for 'fill' is already present. Adding another scale for 'fill', which ## will replace the existing scale.

Scale for 'colour' is already present. Adding another scale for 'colour', ## which will replace the existing scale.



[1] "Analysing data for response: CXCR3"

```
## [1] "Fit for the response: CXCR3"
## [1] "Fitting for all"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupA : F"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupB : M"
```

[1] "Fitting model basic without alpha"

```
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
##
     dLL dDF
                pvalue
            1 0.4941464
## 1 0.23
## [1] "Testing H1 no alpha vs alpha"
     dLL dDF
                pvalue
## 1 0.26
            1 0.4693894
## [1] "Testing H2 groupA no alpha vs alpha"
     dLL dDF
                pvalue
## 1 0.13
           1 0.6142887
## [1] "Testing H2 groupB no alpha vs alpha"
     dLL dDF
                pvalue
## 1 0.11
           1 0.6444226
## [1] "Testing H3 groupA no alpha vs alpha"
     dLL dDF
              pvalue
           1 0.492078
## 1 0.24
## [1] "Testing H3 groupB no alpha vs alpha"
     dLL dDF
              pvalue
           1 0.687482
## 1 0.08
## [1] "Testing H1 vs H0"
##
     dLL dDF
               pvalue
## 1 0.03
            1 0.802799
## [1] "Testing H2 vs H0"
               pvalue
   dLL dDF
## 1 0
           3 0.9998966
## [1] "Testing H3 vs H1"
     dLL dDF
                pvalue
## 1 0.25
           4 0.9732499
## [1] "Testing H3 vs H2"
              pvalue
     dLL dDF
## 1 0.28
          2 0.7559332
```

```
##A11
print(CXCR3)
## $HO
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
            L1
                     alpha
                               myshape
##
  21.13757351 0.05184824
                            5.00000000
##
## Log-likelihood: -887.03
## Best method: bobyqa
##
## $H1
##
## Call:
  bbmle::mle2(minuslogl = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
  Coefficients:
##
           T.1
                        L2
                                  alpha
                                            myshape
## 21.06456046 21.22999921 0.05633644 5.00000000
##
## Log-likelihood: -887
## Best method: bobyga
##
## $H2
  $H2$groupA
##
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
## Coefficients:
```

##

L1

alpha

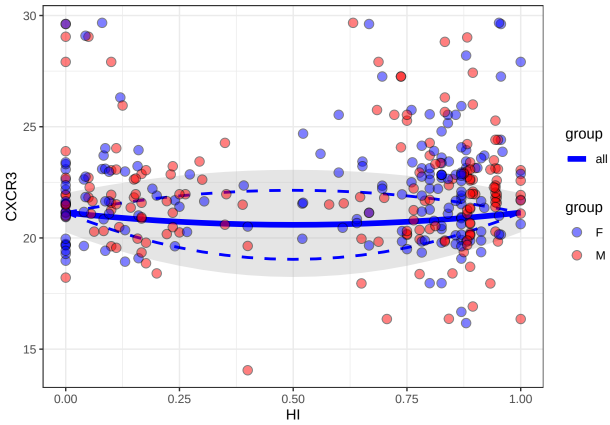
myshape

```
## 21.16012651 0.05309422 5.00000000
##
## Log-likelihood: -441.11
## Best method: bobyqa
## $H2$groupB
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
  Coefficients:
##
            L1
                     alpha
                               myshape
  21.11417149 0.05042914 5.00000000
##
##
## Log-likelihood: -445.92
## Best method: bobyqa
##
##
## $H3
##
  $H3$groupA
##
##
  Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
##
  Coefficients:
                                  alpha
                                            myshape
##
                        1.2
  20.89486805 21.54677628 0.07518605 5.00000000
## Log-likelihood: -440.88
## Best method: L-BFGS-B
##
## $H3$groupB
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
```

```
control = config$control)
##
##
## Coefficients:
##
            L1
                        L2
                                 alpha
                                           myshape
## 21.25139030 20.96471235 0.04478293 5.00000000
##
## Log-likelihood: -445.87
## Best method: L-BFGS-B
bananaPlot(mod = CXCR3$H0,
             data = field,
             response = "CXCR3",
             group = "Sex") +
  scale_fill_manual(values = c("blue", "red")) +
  scale_color_manual(values = c("blue", "red")) +
 theme_bw()
```

Scale for 'fill' is already present. Adding another scale for 'fill', which
will replace the existing scale.

Scale for 'colour' is already present. Adding another scale for 'colour', ## which will replace the existing scale.



```
group = "Sex")
## [1] "Analysing data for response: IL.6"
## [1] "Fit for the response: IL.6"
## [1] "Fitting for all"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupA : F"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
```

model = "weibull"

```
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
   dLL dDF
             pvalue
          1 0.976873
## 1 0
## [1] "Testing H1 no alpha vs alpha"
   dLL dDF
                pvalue
## 1 0.01
           1 0.9058748
## [1] "Testing H2 groupA no alpha vs alpha"
   dLL dDF
               pvalue
## 1 0
         1 0.9545683
## [1] "Testing H2 groupB no alpha vs alpha"
## dLL dDF
               pvalue
## 1 0
          1 0.9881301
## [1] "Testing H3 groupA no alpha vs alpha"
## dLL dDF
               pvalue
## 1 0
          1 0.9770896
## [1] "Testing H3 groupB no alpha vs alpha"
   dLL dDF
               pvalue
          1 0.9771681
## 1 0
## [1] "Testing H1 vs H0"
     dLL dDF
                pvalue
##
## 1 0.12
           1 0.6259119
## [1] "Testing H2 vs H0"
                pvalue
     dLL dDF
## 1 0.18
           3 0.9485655
## [1] "Testing H3 vs H1"
     dLL dDF
                pvalue
## 1 0.37
           4 0.9459875
## [1] "Testing H3 vs H2"
```

```
dLL dDF
                 pvalue
## 1 0.31
            2 0.7330492
##A11
print(IL.6)
## $HO
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
  Coefficients:
##
##
             T.1
                       alpha
                                   myshape
##
  22.358033634 -0.002280237 5.000000000
##
## Log-likelihood: -904.04
## Best method: bobyga
##
## $H1
##
## Call:
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
  Coefficients:
##
             T.1
                          L2
                                     alpha
                                                myshape
## 22.512843288 22.176246390 -0.009510556 5.000000000
##
## Log-likelihood: -903.92
## Best method: bobyqa
##
## $H2
## $H2$groupA
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
```

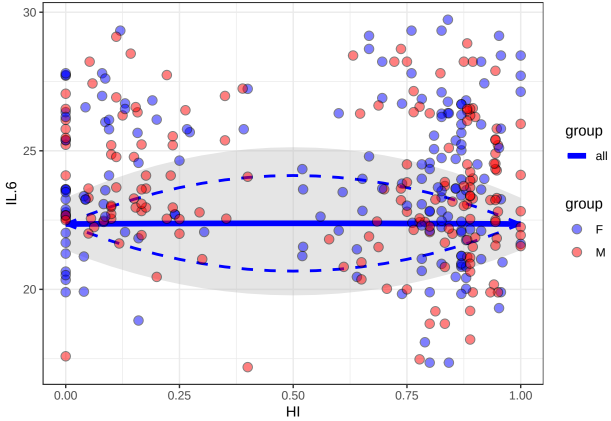
##

```
## Coefficients:
##
             T.1
                       alpha
                                   myshape
##
  22.482401564 -0.006154352 5.000000000
##
## Log-likelihood: -452.81
## Best method: bobyga
## $H2$groupB
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
  Coefficients:
##
##
                       alpha
                                   myshape
##
  22.231268733 0.001715418 5.000000000
##
## Log-likelihood: -451.05
## Best method: bobyga
##
## $H3
##
   $H3$groupA
##
## Call:
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
                          1.2
                                     alpha
                                                myshape
##
  22.448010801 22.534765599 -0.003246811 5.000000000
## Log-likelihood: -452.81
## Best method: bobyqa
##
## $H3$groupB
##
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
```

```
upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
## Coefficients:
             L1
                          L2
                                                myshape
##
                                    alpha
## 22.634336663 21.883191191 -0.003330059 5.000000000
##
## Log-likelihood: -450.74
## Best method: bobyqa
bananaPlot(mod = IL.6$H0,
             data = field,
             response = "IL.6",
             group = "Sex") +
  scale_fill_manual(values = c("blue", "red")) +
  scale_color_manual(values = c("blue", "red")) +
 theme_bw()
```

Scale for 'fill' is already present. Adding another scale for 'fill', which ## will replace the existing scale.

Scale for 'colour' is already present. Adding another scale for 'colour', ## which will replace the existing scale.



```
field$Sex <- as.factor(field$Sex)
speparam <- parasiteLoad::getParamBounds("weibull", data = field, response = "IL.10")</pre>
```

```
IL.10 <- parasiteLoad::analyse(data = field,</pre>
                        response = "IL.10",
                        model = "weibull",
                        group = "Sex")
## [1] "Analysing data for response: IL.10"
## [1] "Fit for the response: IL.10"
## [1] "Fitting for all"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupA : F"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
```

```
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
     dLL dDF
                 pvalue
## 1 0.28
           1 0.4523162
## [1] "Testing H1 no alpha vs alpha"
     dLL dDF
               pvalue
## 1 0.15
           1 0.580233
## [1] "Testing H2 groupA no alpha vs alpha"
     dLL dDF
                pvalue
           1 0.5646307
## 1 0.17
## [1] "Testing H2 groupB no alpha vs alpha"
##
     dLL dDF
               pvalue
## 1 0.12
           1 0.625714
## [1] "Testing H3 groupA no alpha vs alpha"
     dLL dDF
                pvalue
## 1 0.06
            1 0.7246361
## [1] "Testing H3 groupB no alpha vs alpha"
   dLL dDF
                pvalue
## 1 0.1
           1 0.6590938
## [1] "Testing H1 vs H0"
##
     dLL dDF
                 pvalue
## 1 0.36
           1 0.3961769
## [1] "Testing H2 vs H0"
     dLL dDF
                pvalue
## 1 0.04
          3 0.9934943
```

```
## [1] "Testing H3 vs H1"
##
      dLL dDF
                 pvalue
## 1 0.05
            4 0.9987232
## [1] "Testing H3 vs H2"
      dLL dDF
                 pvalue
## 1 0.37
            2 0.6917753
##A11
print(IL.10)
## $HO
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
  Coefficients:
##
##
                     alpha
                               myshape
## 23.57396321 0.05860765 5.00000000
##
## Log-likelihood: -915.52
## Best method: bobyqa
##
## $H1
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
  Coefficients:
##
                              alpha
                                        myshape
  23.8477593 23.2281731 0.0444881 5.0000000
##
## Log-likelihood: -915.16
## Best method: bobyqa
##
## $H2
## $H2$groupA
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibul1(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
```

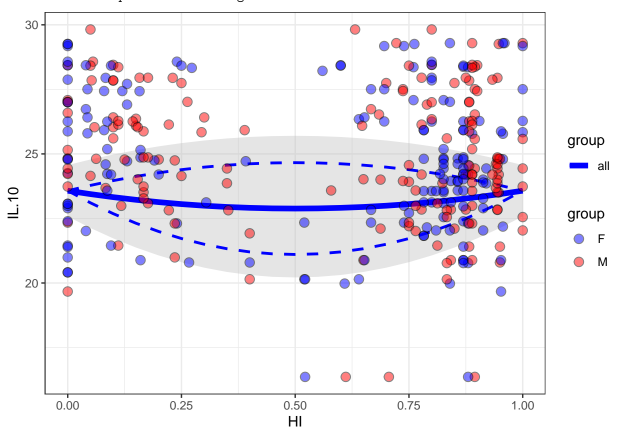
##

```
myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
            T.1
                     alpha
                               myshape
  23.50707937 0.06024782 5.00000000
##
## Log-likelihood: -457.14
## Best method: bobyqa
## $H2$groupB
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
  Coefficients:
##
##
            L1
                     alpha
                               mvshape
## 23.63920972 0.05694935 5.00000000
## Log-likelihood: -458.34
  Best method: bobyqa
##
##
##
## $H3
  $H3$groupA
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
  Coefficients:
##
##
                        L2
                                  alpha
                                            myshape
  23.71854225 23.14229445 0.03961731 5.00000000
##
##
## Log-likelihood: -457
## Best method: bobyqa
## $H3$groupB
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibul1(shape = myshape,
```

```
scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
            L1
                        L2
                                 alpha
                                            myshape
                            0.05194454
  24.00122226 23.32014918
                                        5.00000000
##
## Log-likelihood: -458.11
## Best method: bobyqa
bananaPlot(mod = IL.10$H0,
             data = field,
             response = "IL.10",
             group = "Sex") +
  scale_fill_manual(values = c("blue", "red")) +
  scale_color_manual(values = c("blue", "red")) +
  theme_bw()
```

Scale for 'fill' is already present. Adding another scale for 'fill', which
will replace the existing scale.

Scale for 'colour' is already present. Adding another scale for 'colour',
which will replace the existing scale.

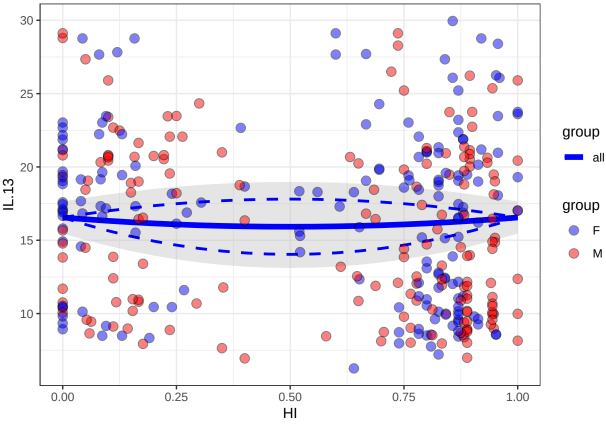


```
field$Sex <- as.factor(field$Sex)</pre>
speparam <- parasiteLoad::getParamBounds("weibull", data = field, response = "IL.13")</pre>
IL.13 <- parasiteLoad::analyse(data = field,</pre>
                        response = "IL.13",
                        model = "weibull",
                        group = "Sex")
## [1] "Analysing data for response: IL.13"
## [1] "Fit for the response: IL.13"
## [1] "Fitting for all"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Fitting for groupA : F"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
     dLL dDF
##
                 pvalue
            1 0.5200098
## 1 0.21
## [1] "Testing H1 no alpha vs alpha"
     dLL dDF
                 pvalue
            1 0.7073971
## 1 0.07
## [1] "Testing H2 groupA no alpha vs alpha"
   dLL dDF
               pvalue
           1 0.5291421
## 1 0.2
## [1] "Testing H2 groupB no alpha vs alpha"
##
      dLL dDF
                 pvalue
## 1 0.02
            1 0.8375112
## [1] "Testing H3 groupA no alpha vs alpha"
      dLL dDF
                 pvalue
##
## 1 0.17
            1 0.5569559
## [1] "Testing H3 groupB no alpha vs alpha"
```

```
dLL dDF
                pvalue
       0
           1 0.9623027
## 1
## [1] "Testing H1 vs H0"
      dLL dDF
                 pvalue
##
## 1 1.19
            1 0.1231585
## [1] "Testing H2 vs H0"
      dLL dDF
                 pvalue
## 1 1.61
            3 0.3579706
## [1] "Testing H3 vs H1"
      dLL dDF
                 pvalue
## 1 2.72
            4 0.2448702
## [1] "Testing H3 vs H2"
    dLL dDF
                pvalue
## 1 2.3
           2 0.1006604
##A11
print(IL.13)
## $HO
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
## Coefficients:
##
            L1
                     alpha
                               myshape
## 16.55284103 0.07616907 3.09803071
## Log-likelihood: -1055.58
## Best method: bobyqa
##
## $H1
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibul1(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
                                       myshape
                      L2
                              alpha
## 17.1494569 15.8898297
                         0.0456834 3.1023105
## Log-likelihood: -1054.39
## Best method: bobyqa
##
```

```
## $H2
## $H2$groupA
##
## Call:
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
  Coefficients:
##
            L1
                     alpha
                                myshape
  17.20862483 0.09777046
                            3.20485373
##
## Log-likelihood: -527.65
## Best method: bobyqa
##
## $H2$groupB
##
## Call:
## bbmle::mle2(minuslogl = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
##
  Coefficients:
##
            T.1
                     alpha
                                myshape
   15.83805312 0.03732138
                            3.02065015
##
## Log-likelihood: -526.32
## Best method: bobyqa
##
##
## $H3
  $H3$groupA
##
##
## Call:
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
            T.1
                        1.2
                                  alpha
                                            myshape
## 17.25774384 17.14480046 0.09423363 3.20425311
```

```
##
## Log-likelihood: -527.64
## Best method: bobyqa
##
## $H3$groupB
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
## Coefficients:
                                    alpha
                                               myshape
## 17.110706241 14.713148410 0.008718456 3.056444824
## Log-likelihood: -524.03
## Best method: bobyqa
bananaPlot(mod = IL.13$HO,
             data = field,
             response = "IL.13",
             group = "Sex") +
  scale_fill_manual(values = c("blue", "red")) +
  scale_color_manual(values = c("blue", "red")) +
 theme_bw()
## Scale for 'fill' is already present. Adding another scale for 'fill', which
## will replace the existing scale.
## Scale for 'colour' is already present. Adding another scale for 'colour',
## which will replace the existing scale.
```



- ## [1] "Analysing data for response: IL1RN"
- ## [1] "Fit for the response: IL1RN"
- ## [1] "Fitting for all"
- ## [1] "Fitting model basic without alpha"
- ## [1] "Did converge"
- ## [1] "Fitting model basic with alpha"
- ## [1] "Did converge"
- ## [1] "Fitting model advanced without alpha"
- ## [1] "Did converge"
- ## [1] "Fitting model advanced with alpha"
- ## [1] "Did converge"
- ## [1] "Fitting for groupA : F"
- ## [1] "Fitting model basic without alpha"
- ## [1] "Did converge"
- ## [1] "Fitting model basic with alpha"
- ## [1] "Did converge"
- ## [1] "Fitting model advanced without alpha"
- ## [1] "Did converge"

```
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
               pvalue
      dLL dDF
            1 0.079619
## 1 1.54
## [1] "Testing H1 no alpha vs alpha"
##
      dLL dDF
                  pvalue
            1 0.09573598
## 1 1.39
## [1] "Testing H2 groupA no alpha vs alpha"
     dLL dDF
                pvalue
## 1 0.9
           1 0.1785661
## [1] "Testing H2 groupB no alpha vs alpha"
      dLL dDF
                 pvalue
## 1 0.65
            1 0.2535602
## [1] "Testing H3 groupA no alpha vs alpha"
##
      dLL dDF
                 pvalue
## 1 0.58
            1 0.2822587
## [1] "Testing H3 groupB no alpha vs alpha"
##
      dLL dDF
                pvalue
## 1 0.75
            1 0.221989
## [1] "Testing H1 vs H0"
      dLL dDF
                 pvalue
            1 0.8594302
## 1 0.02
## [1] "Testing H2 vs H0"
      dLL dDF
                 pvalue
## 1 1.79
            3 0.3114628
## [1] "Testing H3 vs H1"
##
      dLL dDF
                pvalue
## 1 2.21
            4 0.352885
## [1] "Testing H3 vs H2"
##
      dLL dDF
                 pvalue
## 1 0.44
            2 0.6464306
##A11
print(IL1RN)
## $HO
##
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
```

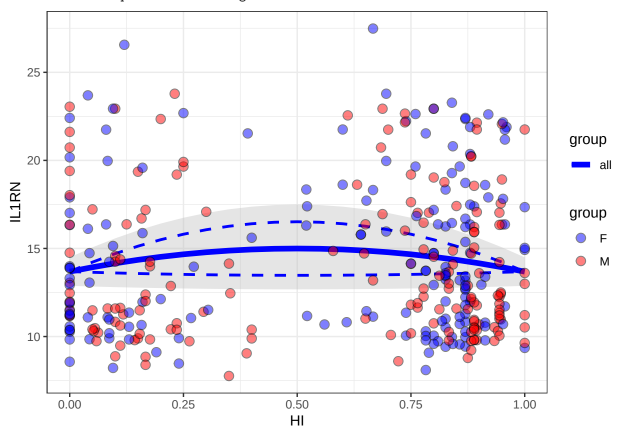
```
##
## Coefficients:
##
                   alpha
                            myshape
## 13.6859746 -0.1914273 3.5477182
##
## Log-likelihood: -961.86
## Best method: bobyga
## $H1
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
                      1.2
                              alpha
                                        myshape
## 13.6383627 13.7456601 -0.1865142 3.5469568
## Log-likelihood: -961.84
## Best method: bobyqa
##
## $H2
## $H2$groupA
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
           T.1
                   alpha
                            myshape
  13.9590093 -0.2154608 3.5120743
##
## Log-likelihood: -485.85
## Best method: bobyqa
##
## $H2$groupB
##
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
```

```
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
  Coefficients:
##
##
                   alpha
                            myshape
## 13.4130727 -0.1676594
                          3.6279845
## Log-likelihood: -474.22
## Best method: bobyga
##
##
## $H3
## $H3$groupA
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
       control = config$control)
##
##
##
  Coefficients:
                      L2
                              alpha
                                        myshape
##
   13.6959865 14.3516835 -0.1775076 3.5107781
##
## Log-likelihood: -485.59
## Best method: bobyqa
##
## $H3$groupB
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
  Coefficients:
                      L2
                              alpha
                                        myshape
## 13.6461956 13.1720271 -0.1827691 3.6358216
##
## Log-likelihood: -474.05
## Best method: bobyqa
bananaPlot(mod = IL1RN$H0,
             data = field,
             response = "IL1RN",
             group = "Sex") +
```

```
scale_fill_manual(values = c("blue", "red")) +
scale_color_manual(values = c("blue", "red")) +
theme_bw()
```

Scale for 'fill' is already present. Adding another scale for 'fill', which
will replace the existing scale.

Scale for 'colour' is already present. Adding another scale for 'colour',
which will replace the existing scale.



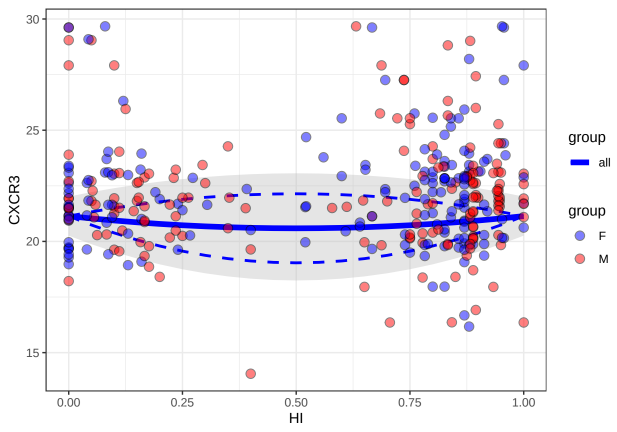
- ## [1] "Analysing data for response: CXCR3"
- ## [1] "Fit for the response: CXCR3"
- ## [1] "Fitting for all"
- ## [1] "Fitting model basic without alpha"
- ## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
- ## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
- ## calculations based on Hessian may be unreliable
- ## [1] "Did converge"

```
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupA : F"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
```

```
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
      dLL dDF
                 pvalue
## 1 0.23
           1 0.4941464
## [1] "Testing H1 no alpha vs alpha"
      dLL dDF
##
                 pvalue
## 1 0.26
            1 0.4693894
## [1] "Testing H2 groupA no alpha vs alpha"
      dLL dDF
                 pvalue
## 1 0.13
            1 0.6142887
## [1] "Testing H2 groupB no alpha vs alpha"
     dLL dDF
                 pvalue
            1 0.6444226
## 1 0.11
## [1] "Testing H3 groupA no alpha vs alpha"
##
      dLL dDF
               pvalue
## 1 0.24
            1 0.492078
## [1] "Testing H3 groupB no alpha vs alpha"
                pvalue
      dLL dDF
## 1 0.08
            1 0.687482
## [1] "Testing H1 vs H0"
     dLL dDF
               pvalue
## 1 0.03
            1 0.802799
## [1] "Testing H2 vs H0"
    dLL dDF
               pvalue
           3 0.9998966
## 1
## [1] "Testing H3 vs H1"
##
      dLL dDF
                 pvalue
## 1 0.25
            4 0.9732499
## [1] "Testing H3 vs H2"
      dLL dDF
                 pvalue
## 1 0.28
            2 0.7559332
##A11
print(CXCR3)
## $HO
##
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
```

```
data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
           T.1
                     alpha
                               myshape
## 21.13757351 0.05184824
                            5.00000000
##
## Log-likelihood: -887.03
## Best method: bobyqa
##
## $H1
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
       control = config$control)
##
##
  Coefficients:
                        1.2
                                 alpha
                                            myshape
##
  21.06456046 21.22999921
                            0.05633644 5.00000000
## Log-likelihood: -887
## Best method: bobyqa
##
## $H2
## $H2$groupA
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
  Coefficients:
            L1
                     alpha
                               myshape
## 21.16012651 0.05309422 5.00000000
##
## Log-likelihood: -441.11
## Best method: bobyqa
## $H2$groupB
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
```

```
scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
## Coefficients:
##
            L1
                     alpha
                               myshape
## 21.11417149 0.05042914
                            5.00000000
## Log-likelihood: -445.92
## Best method: bobyqa
##
##
## $H3
## $H3$groupA
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
  Coefficients:
##
##
            L1
                        L2
                                  alpha
                                            myshape
                            0.07518605 5.00000000
## 20.89486805 21.54677628
##
## Log-likelihood: -440.88
## Best method: L-BFGS-B
## $H3$groupB
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
                        L2
                                  alpha
                                            myshape
##
  21.25139030 20.96471235
                            0.04478293 5.00000000
##
## Log-likelihood: -445.87
## Best method: L-BFGS-B
```



```
## [1] "Analysing data for response: CASP1"
## [1] "Fit for the response: CASP1"
## [1] "Fitting for all"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
```

```
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupA : F"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
```

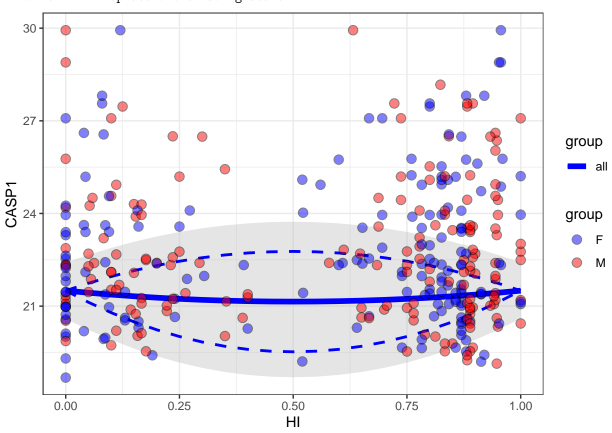
```
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
      dLL dDF
                pvalue
## 1 0.09
           1 0.6713713
## [1] "Testing H1 no alpha vs alpha"
      dLL dDF
##
                 pvalue
## 1 0.13
            1 0.6058085
## [1] "Testing H2 groupA no alpha vs alpha"
      dLL dDF
                pvalue
## 1 0.02
            1 0.8234527
## [1] "Testing H2 groupB no alpha vs alpha"
     dLL dDF
                pvalue
            1 0.7057156
## 1 0.07
## [1] "Testing H3 groupA no alpha vs alpha"
##
      dLL dDF
                pvalue
## 1 0.11
            1 0.6453932
## [1] "Testing H3 groupB no alpha vs alpha"
      dLL dDF
                pvalue
           1 0.710826
## 1 0.07
## [1] "Testing H1 vs H0"
     dLL dDF
                 pvalue
## 1 0.13
            1 0.6067087
## [1] "Testing H2 vs H0"
     dLL dDF
                pvalue
## 1 0.02
            3 0.9982353
## [1] "Testing H3 vs H1"
##
      dLL dDF
                 pvalue
## 1 0.21
            4 0.9803564
## [1] "Testing H3 vs H2"
      dLL dDF
                pvalue
## 1 0.33
            2 0.720831
##A11
print(CASP1)
## $HO
##
```

Call:

```
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
            L1
                     alpha
                               myshape
  21.50099623 0.03297412
                            5.00000000
##
## Log-likelihood: -885.58
## Best method: bobyqa
## $H1
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
  Coefficients:
##
##
                        L2
                                  alpha
            L1
                                            myshape
## 21.34079401 21.68489302 0.04068176 5.00000000
##
## Log-likelihood: -885.45
## Best method: bobyqa
##
## $H2$groupA
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
## Coefficients:
##
            L1
                     alpha
                               myshape
  21.49396241 0.02443874 5.00000000
## Log-likelihood: -442.38
## Best method: bobyqa
##
## $H2$groupB
```

```
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
   Coefficients:
##
            L1
                     alpha
                               myshape
  21.50902537
               0.04166358
                            5.00000000
##
##
## Log-likelihood: -443.18
## Best method: bobyqa
##
##
## $H3
## $H3$groupA
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
                      L2
                              alpha
                                        myshape
  21.1797205 21.9665025
                          0.0522642
                                     5.0000000
##
## Log-likelihood: -442.06
## Best method: bobyga
##
## $H3$groupB
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
       control = config$control)
##
##
##
  Coefficients:
                        L2
                                  alpha
                                            myshape
## 21.54566954 21.47719556 0.04106758 5.00000000
##
```

```
## Log-likelihood: -443.18
## Best method: bobyqa
```



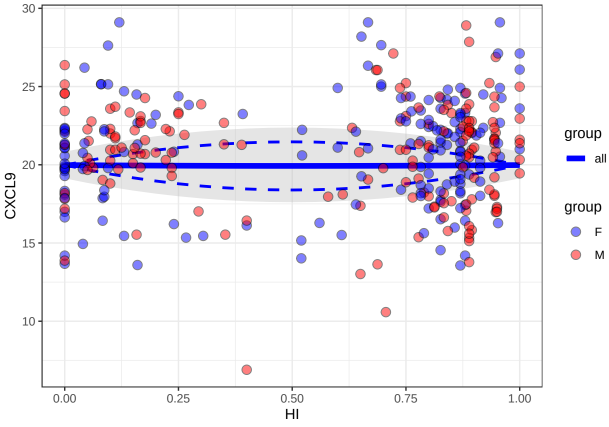
- ## [1] "Analysing data for response: CXCL9"
- ## [1] "Fit for the response: CXCL9"
- ## [1] "Fitting for all"

```
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupA : F"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
```

```
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
    dLL dDF
              pvalue
          1 0.9447407
## 1
       0
## [1] "Testing H1 no alpha vs alpha"
     dLL dDF
                pvalue
## 1 0.02
           1 0.8563652
## [1] "Testing H2 groupA no alpha vs alpha"
##
      dLL dDF
                pvalue
## 1 0.14
           1 0.5997119
## [1] "Testing H2 groupB no alpha vs alpha"
                 pvalue
     dLL dDF
## 1 0.26
           1 0.4744637
## [1] "Testing H3 groupA no alpha vs alpha"
     dLL dDF
                pvalue
## 1 0.04
           1 0.7645265
## [1] "Testing H3 groupB no alpha vs alpha"
     dLL dDF
                pvalue
## 1 0.26
           1 0.4733661
## [1] "Testing H1 vs H0"
##
     dLL dDF
               pvalue
## 1 0.28
            1 0.452086
## [1] "Testing H2 vs H0"
##
     dLL dDF
                 pvalue
## 1 0.72
            3 0.6981508
## [1] "Testing H3 vs H1"
     dLL dDF
                pvalue
## 1 0.86
           4 0.7881596
## [1] "Testing H3 vs H2"
     dLL dDF
               pvalue
## 1 0.42
            2 0.654451
##A11
print(CXCL9)
```

```
## $HO
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
       control = config$control)
##
##
##
   Coefficients:
##
                       alpha
                                   myshape
             L1
   19.981630484
                0.005451623 5.000000000
##
##
## Log-likelihood: -906.19
## Best method: bobyqa
##
## $H1
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
##
  Coefficients:
##
            T.1
                        L2
                                  alpha
                                            myshape
  19.75363306 20.21762770 0.01433242 5.00000000
##
## Log-likelihood: -905.91
## Best method: bobyqa
##
## $H2
## $H2$groupA
##
## Call:
  bbmle::mle2(minuslogl = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
##
  Coefficients:
##
            T.1
                     alpha
                                myshape
## 19.86502220 -0.05730404 5.00000000
##
## Log-likelihood: -455
```

```
## Best method: bobyqa
##
##
  $H2$groupB
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
       control = config$control)
##
##
##
   Coefficients:
##
            L1
                     alpha
                               myshape
##
  20.15461841 0.08152077
                           5.00000000
##
## Log-likelihood: -450.47
## Best method: bobyqa
##
## $H3
## $H3$groupA
##
## Call:
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
   Coefficients:
##
##
           T.1
                      L2
                              alpha
                                        myshape
##
  19.5035914 20.3150787 -0.0333836 5.0000000
##
## Log-likelihood: -454.58
## Best method: bobyqa
##
## $H3$groupB
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
## Coefficients:
```

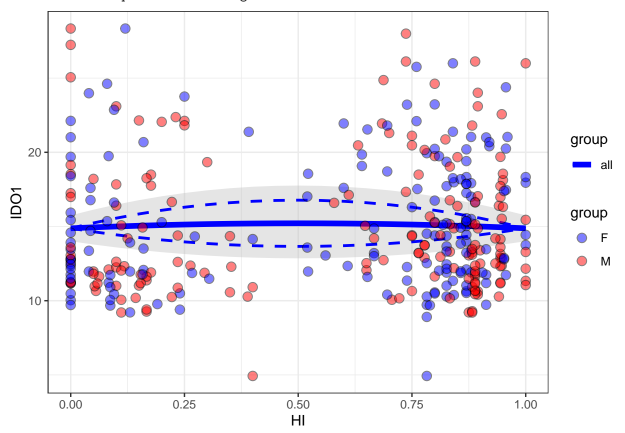


```
## [1] "Analysing data for response: ID01"
## [1] "Fit for the response: ID01"
## [1] "Fitting for all"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Fitting for groupA : F"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
     dLL dDF
                 pvalue
## 1 0.09
            1 0.6683474
## [1] "Testing H1 no alpha vs alpha"
     dLL dDF pvalue
## 1 0.09
           1 0.668404
## [1] "Testing H2 groupA no alpha vs alpha"
     dLL dDF
                 pvalue
## 1 0.32
           1 0.4266815
## [1] "Testing H2 groupB no alpha vs alpha"
    dLL dDF pvalue
           1 0.937834
## 1 0
## [1] "Testing H3 groupA no alpha vs alpha"
     dLL dDF
                pvalue
## 1 0.22
           1 0.5080867
## [1] "Testing H3 groupB no alpha vs alpha"
## dLL dDF
               pvalue
## 1 0
          1 0.9951221
## [1] "Testing H1 vs H0"
## dLL dDF
               pvalue
## 1 0
           1 0.9561119
## [1] "Testing H2 vs H0"
     dLL dDF
                pvalue
## 1 0.98
           3 0.5798636
## [1] "Testing H3 vs H1"
```

```
##
      dLL dDF
                 pvalue
            4 0.6926173
## 1 1.12
  [1] "Testing H3 vs H2"
      dLL dDF
                pvalue
## 1 0.14
            2 0.872036
##A11
print(ID01)
## $HO
##
## Call:
##
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
## Coefficients:
##
           T.1
                   alpha
                            myshape
## 14.8795486 -0.0453282 3.5547258
##
## Log-likelihood: -976.5
## Best method: bobyqa
##
## $H1
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
       control = config$control)
##
##
  Coefficients:
##
##
                        1.2
                                  alpha
                                            myshape
  14.89546892 14.85915732 -0.04688506 3.55516770
##
##
## Log-likelihood: -976.5
## Best method: bobyqa
##
## $H2
  $H2$groupA
##
## Call:
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
```

```
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
  Coefficients:
##
##
                   alpha
                            myshape
## 14.7557708 -0.1199757
                          3.7550175
## Log-likelihood: -481.26
## Best method: bobyga
##
## $H2$groupB
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
  Coefficients:
##
            T.1
                     alpha
                                myshape
## 14.95514457 0.01171584 3.39652130
##
## Log-likelihood: -494.26
## Best method: bobyqa
##
##
## $H3
## $H3$groupA
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
##
          T.1
                    L2
                           alpha
                                    myshape
  14.646811 14.921180 -0.104478 3.750816
##
## Log-likelihood: -481.21
## Best method: bobyqa
##
## $H3$groupB
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
```

```
start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
## Coefficients:
##
                            L2
                                       alpha
                                                    myshape
## 15.1510029708 14.7414325430 -0.0009399171 3.4010182803
## Log-likelihood: -494.17
## Best method: bobyqa
bananaPlot(mod = IDO1$HO,
             data = field,
             response = "ID01",
             group = "Sex") +
  scale_fill_manual(values = c("blue", "red")) +
  scale_color_manual(values = c("blue", "red")) +
  theme_bw()
```

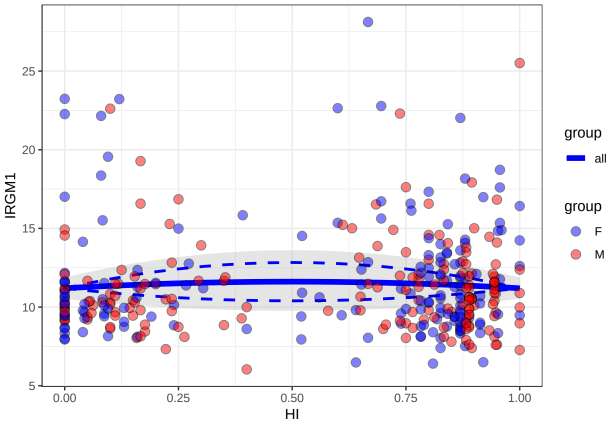


```
field$Sex <- as.factor(field$Sex)</pre>
speparam <- parasiteLoad::getParamBounds("weibull", data = field, response = "IRGM1")</pre>
IRGM1 <- parasiteLoad::analyse(data = field,</pre>
                        response = "IRGM1",
                        model = "weibull",
                        group = "Sex")
## [1] "Analysing data for response: IRGM1"
## [1] "Fit for the response: IRGM1"
## [1] "Fitting for all"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Fitting for groupA : F"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
##
     dLL dDF
                 pvalue
## 1 0.24
            1 0.4927543
## [1] "Testing H1 no alpha vs alpha"
     dLL dDF
                 pvalue
## 1 0.27
            1 0.4609059
## [1] "Testing H2 groupA no alpha vs alpha"
      dLL dDF
                 pvalue
            1 0.4978844
## 1 0.23
## [1] "Testing H2 groupB no alpha vs alpha"
##
   dLL dDF
                pvalue
           1 0.9569908
## [1] "Testing H3 groupA no alpha vs alpha"
      dLL dDF
##
                 pvalue
## 1 0.47
            1 0.3297621
## [1] "Testing H3 groupB no alpha vs alpha"
```

```
dLL dDF
                pvalue
           1 0.9999517
## 1
       0
## [1] "Testing H1 vs H0"
      dLL dDF
                 pvalue
##
## 1 0.06
            1 0.7262511
## [1] "Testing H2 vs H0"
      dLL dDF
                   pvalue
## 1 5.85
            3 0.008523426
## [1] "Testing H3 vs H1"
      dLL dDF
                   pvalue
## 1 6.74
            4 0.009134627
## [1] "Testing H3 vs H2"
      dLL dDF
                 pvalue
            2 0.3834328
## 1 0.96
##A11
print(IRGM1)
## $HO
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
## Coefficients:
##
            L1
                     alpha
                               myshape
## 11.19559941 -0.07493732 3.30034466
## Log-likelihood: -891.91
## Best method: bobyqa
##
## $H1
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibul1(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
                        L2
                                  alpha
                                            myshape
## 11.28436166 11.10078855 -0.08197282 3.30240109
## Log-likelihood: -891.85
## Best method: bobyqa
##
```

```
## $H2
## $H2$groupA
##
## Call:
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
  Coefficients:
##
##
           L1
                   alpha
                             myshape
## 11.3905791 -0.1090206
                          3.0999499
##
## Log-likelihood: -459.72
## Best method: bobyqa
##
## $H2$groupB
##
## Call:
## bbmle::mle2(minuslogl = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
##
  Coefficients:
##
             T.1
                       alpha
                                   myshape
   11.126190811 0.007833011 3.648704592
##
## Log-likelihood: -426.34
## Best method: bobyqa
##
##
## $H3
  $H3$groupA
##
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
           T.1
                      L2
                               alpha
                                        myshape
## 11.7250901 10.8968237 -0.1679786 3.1220485
```

```
##
## Log-likelihood: -459.22
## Best method: bobyqa
##
## $H3$groupB
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
## Coefficients:
##
                            L2
                                       alpha
                                                    myshape
   1.074277e+01 1.137318e+01 -9.759829e-06 3.662007e+00
##
## Log-likelihood: -425.88
## Best method: bobyqa
bananaPlot(mod = IRGM1$H0,
             data = field,
             response = "IRGM1",
             group = "Sex") +
  scale_fill_manual(values = c("blue", "red")) +
  scale_color_manual(values = c("blue", "red")) +
 theme_bw()
## Scale for 'fill' is already present. Adding another scale for 'fill', which
## will replace the existing scale.
## Scale for 'colour' is already present. Adding another scale for 'colour',
## which will replace the existing scale.
```



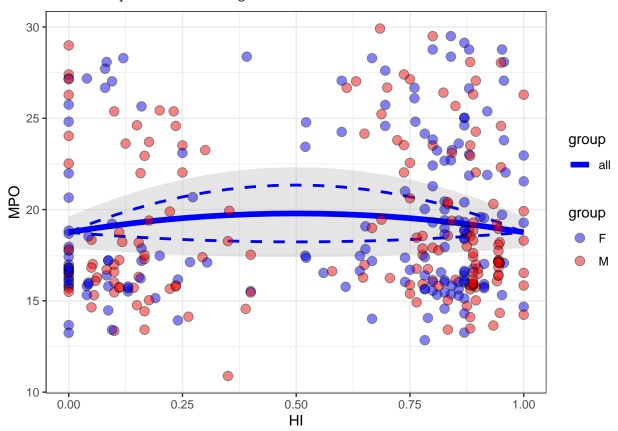
- ## [1] "Analysing data for response: MPO"
- ## [1] "Fit for the response: MPO"
- ## [1] "Fitting for all"
- ## [1] "Fitting model basic without alpha"
- ## [1] "Did converge"
- ## [1] "Fitting model basic with alpha"
- ## [1] "Did converge"
- ## [1] "Fitting model advanced without alpha"
- ## [1] "Did converge"
- ## [1] "Fitting model advanced with alpha"
- ## [1] "Did converge"
- ## [1] "Fitting for groupA : F"
- ## [1] "Fitting model basic without alpha"
- ## [1] "Did converge"
- ## [1] "Fitting model basic with alpha"
- ## [1] "Did converge"
- ## [1] "Fitting model advanced without alpha"
- ## [1] "Did converge"

```
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
      dLL dDF
                 pvalue
            1 0.1836948
## 1 0.88
## [1] "Testing H1 no alpha vs alpha"
##
      dLL dDF
                 pvalue
            1 0.2431323
## 1 0.68
## [1] "Testing H2 groupA no alpha vs alpha"
      dLL dDF
               pvalue
## 1 0.36
            1 0.397991
## [1] "Testing H2 groupB no alpha vs alpha"
      dLL dDF
                 pvalue
## 1 0.56
            1 0.2887894
## [1] "Testing H3 groupA no alpha vs alpha"
##
      dLL dDF
                 pvalue
## 1 0.12
            1 0.6208357
## [1] "Testing H3 groupB no alpha vs alpha"
##
      dLL dDF
                 pvalue
## 1 0.64
            1 0.2592147
## [1] "Testing H1 vs H0"
      dLL dDF
                pvalue
## 1 0.11
            1 0.633915
## [1] "Testing H2 vs H0"
              pvalue
      dLL dDF
## 1 0.98
            3 0.579382
## [1] "Testing H3 vs H1"
##
      dLL dDF
                 pvalue
## 1 1.51
            4 0.5534792
## [1] "Testing H3 vs H2"
##
      dLL dDF
                 pvalue
## 1 0.64
            2 0.5256382
##A11
print(MPO)
## $HO
##
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
```

```
##
## Coefficients:
                 alpha
##
          T.1
                         myshape
## 18.754871 -0.110056 4.629210
## Log-likelihood: -976.21
## Best method: bobyga
## $H1
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
                        1.2
                                  alpha
                                            myshape
## 18.62382075 18.92884013 -0.09929406 4.62917360
## Log-likelihood: -976.1
## Best method: bobyqa
##
## $H2
## $H2$groupA
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
           T.1
                   alpha
                            myshape
##
  19.0549443 -0.1032107 4.5931937
## Log-likelihood: -491.09
## Best method: bobyqa
##
## $H2$groupB
##
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
```

```
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
  Coefficients:
##
##
          T.1
                 alpha
                         myshape
## 18.452376 -0.117932 4.693600
## Log-likelihood: -484.14
## Best method: bobyga
##
##
## $H3
## $H3$groupA
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
       control = config$control)
##
##
##
  Coefficients:
                        1.2
                                  alpha
                                            myshape
##
   18.66472726 19.65178839 -0.06199369 4.59998139
##
## Log-likelihood: -490.54
## Best method: bobyqa
##
## $H3$groupB
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
  Coefficients:
                      L2
                              alpha
                                        myshape
## 18.6295086 18.2485621 -0.1288016 4.6979042
##
## Log-likelihood: -484.05
## Best method: bobyqa
bananaPlot(mod = MPO$HO,
             data = field,
             response = "MPO",
             group = "Sex") +
```

```
scale_fill_manual(values = c("blue", "red")) +
scale_color_manual(values = c("blue", "red")) +
theme_bw()
```



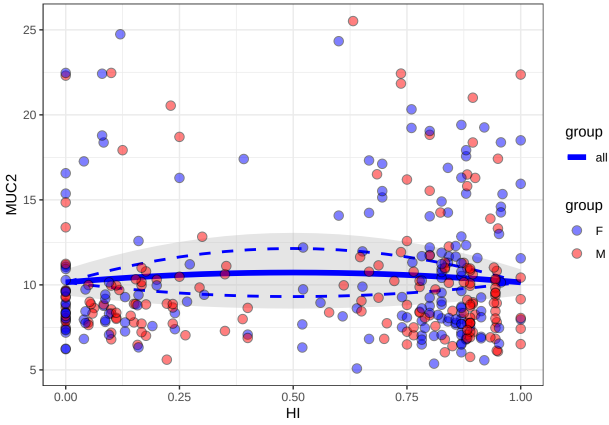
- ## [1] "Analysing data for response: MUC2"
- ## [1] "Fit for the response: MUC2"
- ## [1] "Fitting for all"
- ## [1] "Fitting model basic without alpha"
- ## [1] "Did converge"
- ## [1] "Fitting model basic with alpha"
- ## [1] "Did converge"
- ## [1] "Fitting model advanced without alpha"
- ## [1] "Did converge"

```
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Fitting for groupA : F"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
      dLL dDF
                pvalue
            1 0.4119459
## 1 0.34
## [1] "Testing H1 no alpha vs alpha"
##
                pvalue
      dLL dDF
## 1 0.31
            1 0.4293845
## [1] "Testing H2 groupA no alpha vs alpha"
     dLL dDF
               pvalue
           1 0.818322
## 1 0.03
## [1] "Testing H2 groupB no alpha vs alpha"
   dLL dDF
               pvalue
## 1 0.5
           1 0.3168754
## [1] "Testing H3 groupA no alpha vs alpha"
      dLL dDF
                pvalue
## 1 0.03
          1 0.8167992
## [1] "Testing H3 groupB no alpha vs alpha"
      dLL dDF
                 pvalue
## 1 0.47
            1 0.3318034
## [1] "Testing H1 vs H0"
    dLL dDF
               pvalue
           1 0.9439104
## 1 0
## [1] "Testing H2 vs H0"
     dLL dDF
                pvalue
## 1 0.91
           3 0.6091541
## [1] "Testing H3 vs H1"
##
      dLL dDF
                pvalue
## 1 0.94
            4 0.756035
## [1] "Testing H3 vs H2"
      dLL dDF
                 pvalue
## 1 0.03
            2 0.9665639
##A11
print(MUC2)
```

\$HO

```
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
  Coefficients:
##
           L1
                   alpha
                            myshape
  10.1482011 -0.1138808
##
                          2.6959899
##
## Log-likelihood: -927.03
## Best method: bobyqa
##
## $H1
##
## Call:
  bbmle::mle2(minuslogl = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
       control = config$control)
##
##
  Coefficients:
##
                      L2
                              alpha
                                        myshape
  10.1288812 10.1700892 -0.1118408 2.6956908
## Log-likelihood: -927.02
## Best method: bobyga
## $H2
## $H2$groupA
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
##
  Coefficients:
##
                   alpha
                            myshape
  10.5576903 -0.0442586 2.6942756
##
## Log-likelihood: -468
## Best method: L-BFGS-B
```

```
##
## $H2$groupB
##
## Call:
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
   Coefficients:
##
          L1
                 alpha
                         myshape
    9.712777 -0.200091 2.709400
##
##
## Log-likelihood: -458.11
## Best method: bobyqa
##
##
## $H3
## $H3$groupA
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
  Coefficients:
##
                                  alpha
                        L2
                                            myshape
  10.57250704 10.53754890 -0.04652781 2.69471223
##
##
## Log-likelihood: -468
## Best method: bobyqa
##
## $H3$groupB
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
## Coefficients:
##
           L1
                      L2
                              alpha
                                        myshape
```



[1] "Analysing data for response: MUC5AC"

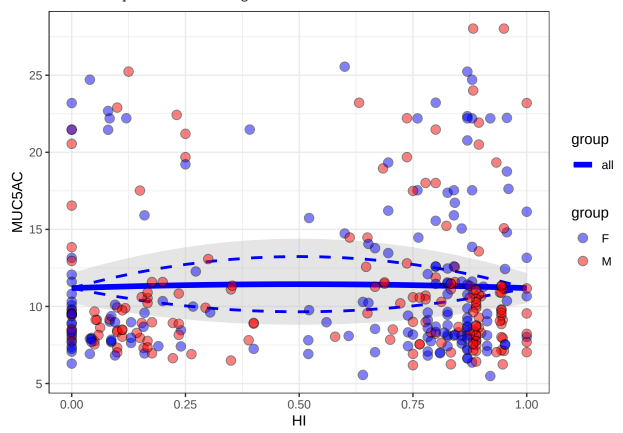
```
## [1] "Fit for the response: MUC5AC"
## [1] "Fitting for all"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Fitting for groupA : F"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
##
      dLL dDF
                 pvalue
## 1 0.04
            1 0.7774737
## [1] "Testing H1 no alpha vs alpha"
      dLL dDF
                 pvalue
## 1 0.01
            1 0.9163143
## [1] "Testing H2 groupA no alpha vs alpha"
     dLL dDF
                 pvalue
## 1 0.01
            1 0.9163009
## [1] "Testing H2 groupB no alpha vs alpha"
##
      dLL dDF
                 pvalue
## 1 0.05
            1 0.7572557
## [1] "Testing H3 groupA no alpha vs alpha"
      dLL dDF
                 pvalue
## 1 0.01
            1 0.8757905
## [1] "Testing H3 groupB no alpha vs alpha"
     dLL dDF
                 pvalue
            1 0.7811917
## 1 0.04
## [1] "Testing H1 vs H0"
    dLL dDF
                pvalue
## 1 0.4
           1 0.3710776
## [1] "Testing H2 vs H0"
##
      dLL dDF
                 pvalue
## 1 0.42
           3 0.8390002
## [1] "Testing H3 vs H1"
##
      dLL dDF
                 pvalue
```

```
## 1 0.48
            4 0.9147444
## [1] "Testing H3 vs H2"
      dLL dDF
                 pvalue
            2 0.6301823
## 1 0.46
##A11
print(MUC5AC)
## $HO
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
  Coefficients:
##
            L1
                     alpha
                               myshape
## 11.19018063 -0.04570668 2.43259875
##
## Log-likelihood: -984.71
## Best method: bobyqa
##
## $H1
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
                        L2
                                  alpha
            L1
                                            myshape
  10.89482803 11.53606022 -0.01706037 2.43489799
##
## Log-likelihood: -984.31
## Best method: bobyqa
##
## $H2
## $H2$groupA
##
## Call:
##
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
```

```
##
       control = config$control)
##
##
  Coefficients:
##
            T.1
                     alpha
                               myshape
##
  11.43407356 -0.02433222
                           2.40856703
##
## Log-likelihood: -496.3
## Best method: bobyqa
##
## $H2$groupB
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
                     alpha
                               myshape
## 10.93963605 -0.06977184
                           2.46002231
## Log-likelihood: -487.98
## Best method: bobyqa
##
##
## $H3
## $H3$groupA
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
  Coefficients:
##
                        L2
                                  alpha
                                            myshape
  11.08524393 11.97805583 0.03706757 2.40963793
##
## Log-likelihood: -495.96
## Best method: bobyqa
##
## $H3$groupB
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibul1(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
```

```
data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
       control = config$control)
##
##
##
  Coefficients:
##
                        L2
                                 alpha
                                           myshape
## 10.68317077 11.15961692 -0.06233649 2.46303965
##
## Log-likelihood: -487.86
## Best method: bobyqa
bananaPlot(mod = MUC5AC$HO,
             data = field,
             response = "MUC5AC",
             group = "Sex") +
  scale_fill_manual(values = c("blue", "red")) +
  scale_color_manual(values = c("blue", "red")) +
  theme_bw()
```

Scale for 'colour' is already present. Adding another scale for 'colour',
which will replace the existing scale.



field\$Sex <- as.factor(field\$Sex)</pre>

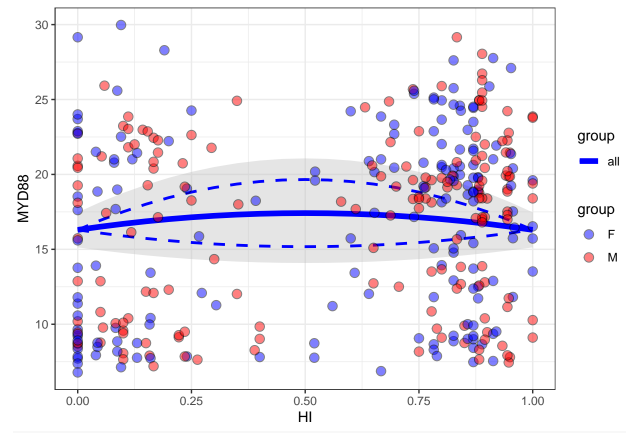
```
speparam <- parasiteLoad::getParamBounds("weibull", data = field, response = "MYD88")
MYD88 <- parasiteLoad::analyse(data = field,
                        response = "MYD88",
                        model = "weibull",
                        group = "Sex")
## [1] "Analysing data for response: MYD88"
## [1] "Fit for the response: MYD88"
## [1] "Fitting for all"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Fitting for groupA : F"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
##
     dLL dDF
                pvalue
           1 0.3130745
## 1 0.51
## [1] "Testing H1 no alpha vs alpha"
     dLL dDF
              pvalue
           1 0.611203
## 1 0.13
## [1] "Testing H2 groupA no alpha vs alpha"
##
     dLL dDF
                 pvalue
## 1 0.81
           1 0.2034581
## [1] "Testing H2 groupB no alpha vs alpha"
                 pvalue
     dLL dDF
## 1 0.01
           1 0.9031706
## [1] "Testing H3 groupA no alpha vs alpha"
     dLL dDF
                pvalue
## 1 0.24
           1 0.4902722
## [1] "Testing H3 groupB no alpha vs alpha"
## dLL dDF pvalue
```

1 0 1 0.942868

```
## [1] "Testing H1 vs H0"
                  pvalue
##
      dLL dDF
            1 0.02625978
## 1 2.47
## [1] "Testing H2 vs H0"
      dLL dDF
                 pvalue
## 1 1.17
            3 0.5060472
## [1] "Testing H3 vs H1"
##
      dLL dDF
                 pvalue
## 1 1.14
            4 0.6835506
## [1] "Testing H3 vs H2"
      dLL dDF
                  pvalue
            2 0.08674323
## 1 2.44
##A11
print (MYD88)
## $HO
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
       control = config$control)
##
##
## Coefficients:
           L1
                   alpha
                            myshape
## 16.2986324 -0.1374078 3.2122525
##
## Log-likelihood: -1064.23
## Best method: bobyqa
##
## $H1
##
## bbmle::mle2(minuslogl = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
## Coefficients:
                        L2
                                  alpha
                                            myshape
  15.50631266 17.28908021 -0.06862561 3.21673665
##
## Log-likelihood: -1061.76
## Best method: bobyga
##
## $H2
## $H2$groupA
```

```
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
  Coefficients:
##
          L1
                 alpha
                         myshape
  15.749626 -0.261844
                        3.042196
##
## Log-likelihood: -536.29
## Best method: bobyqa
##
## $H2$groupB
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
  Coefficients:
            L1
                     alpha
                               myshape
## 16.84579190 -0.02174222 3.41265792
##
## Log-likelihood: -526.78
## Best method: bobyqa
##
##
## $H3
## $H3$groupA
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
## Coefficients:
           T.1
                      L2
                              alpha
                                        myshape
## 15.2607632 16.7667979 -0.1492716 3.0359027
##
## Log-likelihood: -535.55
```

```
## Best method: bobyqa
##
## $H3$groupB
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
## Coefficients:
##
            L1
                        L2
                                 alpha
                                           myshape
## 15.72515033 17.69501413 -0.01248977 3.43791893
## Log-likelihood: -525.07
## Best method: bobyga
bananaPlot(mod = MYD88$H0,
             data = field,
             response = "MYD88",
             group = "Sex") +
  scale_fill_manual(values = c("blue", "red")) +
  scale_color_manual(values = c("blue", "red")) +
 theme_bw()
## Scale for 'fill' is already present. Adding another scale for 'fill', which
## will replace the existing scale.
## Scale for 'colour' is already present. Adding another scale for 'colour',
## which will replace the existing scale.
```



```
field$Sex <- as.factor(field$Sex)</pre>
speparam <- parasiteLoad::getParamBounds("weibull", data = field, response = "NCR1")</pre>
NCR1 <- parasiteLoad::analyse(data = field,</pre>
                        response = "NCR1",
                        model = "weibull",
                        group = "Sex")
## [1] "Analysing data for response: NCR1"
## [1] "Fit for the response: NCR1"
## [1] "Fitting for all"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
```

```
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupA : F"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
```

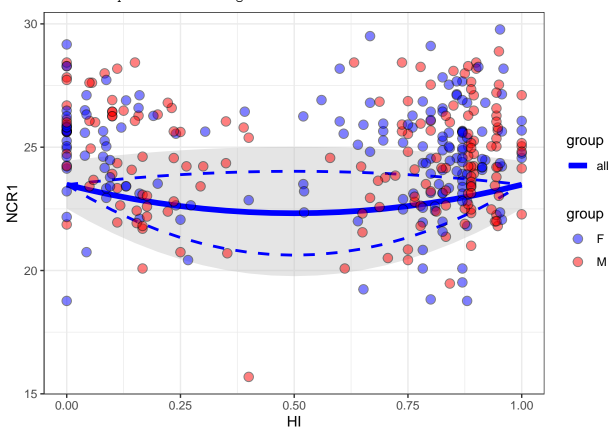
```
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
      dLL dDF
                 pvalue
## 1 0.85
            1 0.1926329
## [1] "Testing H1 no alpha vs alpha"
      dLL dDF
                 pvalue
## 1 0.76
            1 0.2183278
## [1] "Testing H2 groupA no alpha vs alpha"
##
      dLL dDF
                 pvalue
## 1 0.14
            1 0.5949384
## [1] "Testing H2 groupB no alpha vs alpha"
      dLL dDF
                 pvalue
            1 0.1757295
## 1 0.92
## [1] "Testing H3 groupA no alpha vs alpha"
     dLL dDF
                 pvalue
            1 0.6105077
## 1 0.13
## [1] "Testing H3 groupB no alpha vs alpha"
##
      dLL dDF
                 pvalue
## 1 0.87
            1 0.1875805
## [1] "Testing H1 vs H0"
      dLL dDF pvalue
##
## 1 0.01
            1 0.87117
## [1] "Testing H2 vs H0"
      dLL dDF
                 pvalue
## 1 0.21
            3 0.9351275
## [1] "Testing H3 vs H1"
      dLL dDF
                 pvalue
## 1 0.27
            4 0.9696534
## [1] "Testing H3 vs H2"
      dLL dDF
              pvalue
## 1 0.07
            2 0.932305
##A11
print(NCR1)
## $HO
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
## Coefficients:
                     alpha
                               myshape
## 23.47968070 0.09858866 5.00000000
## Log-likelihood: -898.46
```

```
## Best method: bobyqa
##
## $H1
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
  Coefficients:
##
##
            L1
                        L2
                                  alpha
                                            myshape
## 23.53250341 23.41385461 0.09587579 5.00000000
##
## Log-likelihood: -898.45
## Best method: bobyga
##
## $H2
## $H2$groupA
##
## Call:
  bbmle::mle2(minuslogl = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
  Coefficients:
##
                     alpha
                               myshape
##
##
  23.26482586 0.05503497
                            5.00000000
##
## Log-likelihood: -447.4
## Best method: bobyqa
##
## $H2$groupB
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
## Coefficients:
##
           T.1
                            myshape
                   alpha
## 23.7424488 0.1502043 5.0000000
```

```
##
## Log-likelihood: -450.85
## Best method: L-BFGS-B
##
##
## $H3
## $H3$groupA
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
       control = config$control)
##
##
##
  Coefficients:
##
                              alpha
                                        myshape
##
  23.2495452 23.2909717 0.0565567 5.0000000
##
## Log-likelihood: -447.4
## Best method: bobyga
##
## $H3$groupB
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
## Coefficients:
                              alpha
##
                      L2
                                        myshape
## 23.9458833 23.5635635 0.1473126 5.0000000
## Log-likelihood: -450.78
## Best method: bobyga
bananaPlot(mod = NCR1$H0,
             data = field,
             response = "NCR1",
             group = "Sex") +
  scale_fill_manual(values = c("blue", "red")) +
  scale_color_manual(values = c("blue", "red")) +
  theme_bw()
## Scale for 'fill' is already present. Adding another scale for 'fill', which
```

will replace the existing scale.

```
## Scale for 'colour' is already present. Adding another scale for 'colour',
## which will replace the existing scale.
```



```
field$Sex <- as.factor(field$Sex)</pre>
speparam <- parasiteLoad::getParamBounds("weibull", data = field, response = "PRF1")</pre>
PRF1 <- parasiteLoad::analyse(data = field,
                        response = "PRF1",
                        model = "weibull",
                        group = "Sex")
## [1] "Analysing data for response: PRF1"
## [1] "Fit for the response: PRF1"
## [1] "Fitting for all"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
```

[1] "Fitting model advanced without alpha"

```
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupA : F"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
```

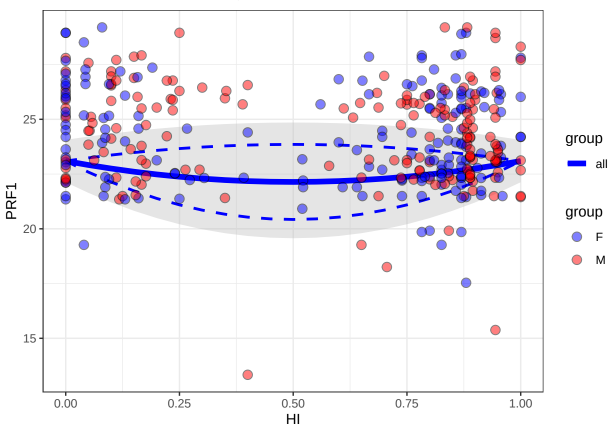
```
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
##
      dLL dDF
                 pvalue
## 1 0.57
            1 0.2839376
## [1] "Testing H1 no alpha vs alpha"
      dLL dDF
                pvalue
## 1 0.39
            1 0.374593
## [1] "Testing H2 groupA no alpha vs alpha"
     dLL dDF
                 pvalue
## 1 0.92
            1 0.1751379
## [1] "Testing H2 groupB no alpha vs alpha"
      dLL dDF
              pvalue
            1 0.900514
## 1 0.01
## [1] "Testing H3 groupA no alpha vs alpha"
##
      dLL dDF
                 pvalue
## 1 0.66
            1 0.2494584
## [1] "Testing H3 groupB no alpha vs alpha"
##
      dLL dDF
                 pvalue
## 1 0.01
            1 0.9025852
## [1] "Testing H1 vs H0"
    dLL dDF pvalue
## 1 0.3
           1 0.435094
## [1] "Testing H2 vs H0"
##
      dLL dDF
                 pvalue
## 1 0.37
            3 0.8655757
## [1] "Testing H3 vs H1"
    dLL dDF
                pvalue
## 1 0.3
           4 0.9640454
## [1] "Testing H3 vs H2"
     dLL dDF
                 pvalue
## 1 0.23
            2 0.7913691
##A11
print(PRF1)
## $HO
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
## Coefficients:
            L1
                     alpha
                               myshape
## 23.09514365 0.08269669 5.00000000
```

```
##
## Log-likelihood: -898.26
## Best method: bobyqa
##
## $H1
##
## bbmle::mle2(minuslog1 = response ~ dweibul1(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
  Coefficients:
                                  alpha
##
            L1
                        L2
                                            myshape
  23.34591766 22.78849290 0.07060213 5.00000000
##
##
## Log-likelihood: -897.96
## Best method: bobyqa
##
## $H2
## $H2$groupA
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
##
           T.1
                   alpha
                            myshape
## 23.3547596 0.1405258 5.0000000
##
## Log-likelihood: -446.86
## Best method: bobyga
##
## $H2$groupB
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
## Coefficients:
```

```
L1
                     alpha
                               myshape
## 22.77637819 0.01448086 5.00000000
##
## Log-likelihood: -451.03
## Best method: bobyqa
##
##
## $H3
## $H3$groupA
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
       control = config$control)
##
##
##
  Coefficients:
##
                      L2
                              alpha
                                        myshape
## 23.4503911 23.1790306 0.1301858 5.0000000
##
## Log-likelihood: -446.83
## Best method: bobyqa
##
## $H3$groupB
##
## Call:
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
                        L2
                                 alpha
                                            myshape
  23.13315274 22.50826018 0.01425636 5.00000000
##
## Log-likelihood: -450.83
## Best method: L-BFGS-B
bananaPlot(mod = PRF1$H0,
             data = field,
             response = "PRF1",
             group = "Sex") +
  scale_fill_manual(values = c("blue", "red")) +
  scale_color_manual(values = c("blue", "red")) +
  theme_bw()
```

Scale for 'fill' is already present. Adding another scale for 'fill', which

```
\mbox{\tt \#\#} will replace the existing scale.
```



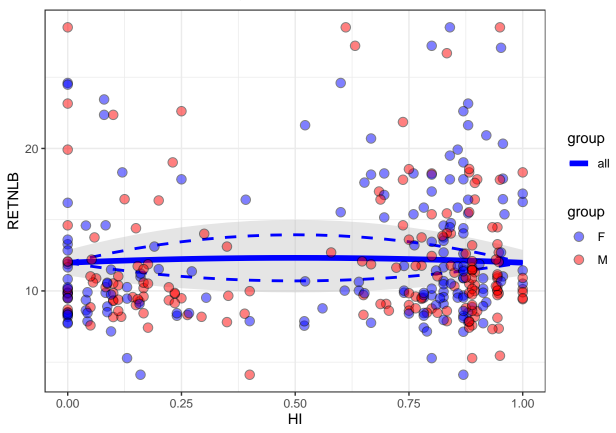
- ## [1] "Analysing data for response: RETNLB"
- ## [1] "Fit for the response: RETNLB"
- ## [1] "Fitting for all"
- ## [1] "Fitting model basic without alpha"
- ## [1] "Did converge"
- ## [1] "Fitting model basic with alpha"
- ## [1] "Did converge"
- ## [1] "Fitting model advanced without alpha"
- ## [1] "Did converge"
- ## [1] "Fitting model advanced with alpha"
- ## [1] "Did converge"
- ## [1] "Fitting for groupA : F"
- ## [1] "Fitting model basic without alpha"
- ## [1] "Did converge"

```
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
##
      dLL dDF
                 pvalue
## 1 0.09
            1 0.6752996
## [1] "Testing H1 no alpha vs alpha"
   dLL dDF
                pvalue
## 1
       0
           1 0.9602801
## [1] "Testing H2 groupA no alpha vs alpha"
      dLL dDF
##
                 pvalue
## 1 0.02
            1 0.8274163
## [1] "Testing H2 groupB no alpha vs alpha"
      dLL dDF
                 pvalue
## 1 0.08
            1 0.6940514
## [1] "Testing H3 groupA no alpha vs alpha"
     dLL dDF
                 pvalue
## 1 0.12
            1 0.6274213
## [1] "Testing H3 groupB no alpha vs alpha"
##
      dLL dDF
                 pvalue
## 1 0.09
            1 0.6773944
## [1] "Testing H1 vs H0"
      dLL dDF
                 pvalue
## 1 0.93
            1 0.1729385
## [1] "Testing H2 vs H0"
     dLL dDF
                 pvalue
## 1 0.81
            3 0.6567631
## [1] "Testing H3 vs H1"
      dLL dDF
                 pvalue
## 1 1.97
            4 0.4140644
## [1] "Testing H3 vs H2"
      dLL dDF
                 pvalue
## 1 2.09
            2 0.1232809
##A11
print(RETNLB)
## $HO
##
## bbmle::mle2(minuslog1 = response ~ dweibul1(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
```

```
data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
                     alpha
                               myshape
## 11.98025381 -0.05703314
                            2.69643002
##
## Log-likelihood: -979.82
## Best method: bobyqa
##
## $H1
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
       control = config$control)
##
##
  Coefficients:
                          L2
                                     alpha
                                                myshape
##
  11.560733567 12.523947050 -0.006854438
                                            2.699198022
## Log-likelihood: -978.89
## Best method: bobyga
##
## $H2
## $H2$groupA
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
  Coefficients:
                     alpha
                               myshape
## 12.31694795 -0.04355707
                           2.69496495
##
## Log-likelihood: -494.58
## Best method: bobyqa
## $H2$groupB
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibul1(shape = myshape,
```

```
scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
## Coefficients:
##
           T.1
                   alpha
                            myshape
## 11.6371020 -0.0727024
                         2.7070206
## Log-likelihood: -484.43
## Best method: bobyqa
##
##
## $H3
## $H3$groupA
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibul1(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
  Coefficients:
##
##
            L1
                        L2
                                  alpha
                                            myshape
## 11.47963497 13.65113222 0.09626788 2.71933993
##
## Log-likelihood: -492.5
## Best method: bobyqa
## $H3$groupB
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
                        L2
                                  alpha
                                            myshape
## 11.71141177 11.55903491 -0.07836459
                                        2.70810144
## Log-likelihood: -484.42
## Best method: bobyga
```

Scale for 'fill' is already present. Adding another scale for 'fill', which ## will replace the existing scale.



```
## [1] "Analysing data for response: SOCS1"
## [1] "Fit for the response: SOCS1"
```

^{## [1] &}quot;Fitting for all"

^{## [1] &}quot;Fitting model basic without alpha"

^{## [1] &}quot;Did converge"

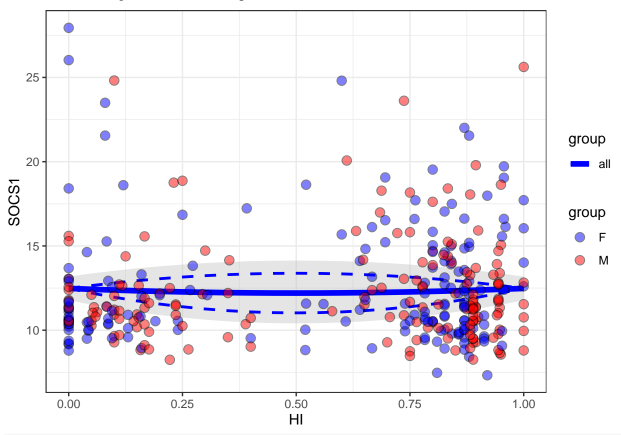
```
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Fitting for groupA : F"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
   dLL dDF
               pvalue
## 1 0.1
           1 0.6485448
## [1] "Testing H1 no alpha vs alpha"
     dLL dDF
                 pvalue
## 1 0.08
           1 0.6827193
## [1] "Testing H2 groupA no alpha vs alpha"
##
      dLL dDF
                 pvalue
## 1 0.52
            1 0.3095676
## [1] "Testing H2 groupB no alpha vs alpha"
     dLL dDF
                 pvalue
## 1 0.31
            1 0.4302486
## [1] "Testing H3 groupA no alpha vs alpha"
     dLL dDF
                 pvalue
## 1 0.26
            1 0.4733257
## [1] "Testing H3 groupB no alpha vs alpha"
      dLL dDF
                pvalue
## 1 0.33
          1 0.4189738
## [1] "Testing H1 vs H0"
               pvalue
##
      dLL dDF
            1 0.853247
## 1 0.02
## [1] "Testing H2 vs H0"
                  pvalue
##
     dLL dDF
## 1 3.81
            3 0.05453768
## [1] "Testing H3 vs H1"
     dLL dDF
                  pvalue
## 1 4.96
           4 0.04192149
## [1] "Testing H3 vs H2"
##
      dLL dDF
                 pvalue
## 1 1.16
            2 0.3124842
```

```
##A11
print(SOCS1)
## $HO
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
            L1
                     alpha
                                myshape
##
  12.47978310 0.04433325
                            3.56671359
##
## Log-likelihood: -895.85
## Best method: bobyqa
##
## $H1
##
## Call:
  bbmle::mle2(minuslogl = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
  Coefficients:
##
            T.1
                        L2
                                  alpha
                                            myshape
## 12.52829370 12.42740777
                            0.04064683 3.56873122
##
## Log-likelihood: -895.83
## Best method: bobyga
##
## $H2
  $H2$groupA
##
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
## Coefficients:
##
           T.1
                   alpha
                            myshape
```

```
## 13.0361320 0.1368053 3.4447276
##
## Log-likelihood: -457.09
## Best method: bobyqa
## $H2$groupB
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
  Coefficients:
##
                   alpha
                            myshape
  11.7697137 -0.1119764 3.7746229
##
##
## Log-likelihood: -434.94
## Best method: bobyqa
##
##
## $H3
##
  $H3$groupA
##
##
  Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
##
  Coefficients:
                      L2
##
                              alpha
                                       myshape
## 13.3177711 12.6583063 0.1026318 3.4670416
## Log-likelihood: -456.8
## Best method: bobyqa
##
## $H3$groupB
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
```

```
control = config$control)
##
##
## Coefficients:
##
           L1
                      L2
                              alpha
                                       myshape
## 11.2507889 12.1387258 -0.1140703 3.7955853
## Log-likelihood: -434.07
## Best method: bobyqa
bananaPlot(mod = SOCS1$H0,
             data = field,
             response = "SOCS1",
             group = "Sex") +
  scale_fill_manual(values = c("blue", "red")) +
  scale_color_manual(values = c("blue", "red")) +
 theme_bw()
```

Scale for 'fill' is already present. Adding another scale for 'fill', which
will replace the existing scale.



```
group = "Sex")
## [1] "Analysing data for response: TICAM1"
## [1] "Fit for the response: TICAM1"
## [1] "Fitting for all"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupA : F"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
```

model = "weibull"

```
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
     dLL dDF
                 pvalue
## 1 0.22
           1 0.5037225
## [1] "Testing H1 no alpha vs alpha"
     dLL dDF
                 pvalue
## 1 0.25
            1 0.4795001
## [1] "Testing H2 groupA no alpha vs alpha"
     dLL dDF
                pvalue
## 1 0.01
           1 0.9159583
## [1] "Testing H2 groupB no alpha vs alpha"
##
     dLL dDF
                 pvalue
## 1 0.57
            1 0.2875548
## [1] "Testing H3 groupA no alpha vs alpha"
##
     dLL dDF
                pvalue
## 1 0.06
           1 0.7298372
## [1] "Testing H3 groupB no alpha vs alpha"
     dLL dDF
                pvalue
## 1 0.51
           1 0.3121757
## [1] "Testing H1 vs H0"
     dLL dDF
                 pvalue
##
## 1 0.04
            1 0.7824371
## [1] "Testing H2 vs H0"
     dLL dDF
                pvalue
## 1 0.36
           3 0.8686206
## [1] "Testing H3 vs H1"
     dLL dDF
                pvalue
## 1 1.96
            4 0.4163236
## [1] "Testing H3 vs H2"
```

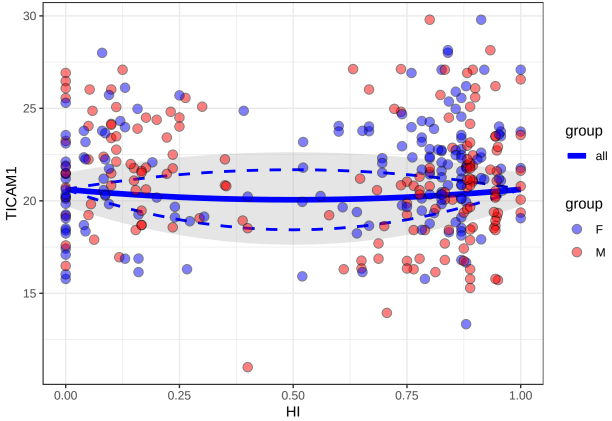
```
dLL dDF
                 pvalue
            2 0.1938556
## 1 1.64
##A11
print(TICAM1)
## $HO
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
  Coefficients:
##
            T.1
                     alpha
                               myshape
##
  20.61868464 0.05446208 5.00000000
##
## Log-likelihood: -898.42
## Best method: bobyga
##
## $H1
##
## Call:
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
  Coefficients:
##
            L1
                        L2
                                  alpha
                                            myshape
## 20.53603913 20.71213328 0.05835373 5.00000000
##
## Log-likelihood: -898.38
## Best method: bobyqa
##
## $H2
## $H2$groupA
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
```

##

```
## Coefficients:
##
            T.1
                     alpha
                               myshape
## 20.33826707 -0.01232009
                            5.00000000
##
## Log-likelihood: -444.13
## Best method: bobyga
## $H2$groupB
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
  Coefficients:
##
##
                   alpha
                            myshape
##
  20.9029777 0.1210809 5.0000000
##
## Log-likelihood: -453.93
## Best method: bobyga
##
## $H3
##
   $H3$groupA
##
## Call:
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
  Coefficients:
##
                        L2
                                  alpha
                                            myshape
##
  19.80926933 21.15065608 0.04130248 5.00000000
## Log-likelihood: -443.09
## Best method: bobyqa
##
## $H3$groupB
##
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
```

```
upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
## Coefficients:
                      L2
##
           T.1
                              alpha
                                       myshape
## 21.4417557 20.4513218 0.1164115 5.0000000
##
## Log-likelihood: -453.32
## Best method: bobyqa
bananaPlot(mod = TICAM1$H0,
             data = field,
             response = "TICAM1",
             group = "Sex") +
  scale_fill_manual(values = c("blue", "red")) +
  scale_color_manual(values = c("blue", "red")) +
 theme_bw()
```

Scale for 'fill' is already present. Adding another scale for 'fill', which ## will replace the existing scale.



```
field$Sex <- as.factor(field$Sex)
speparam <- parasiteLoad::getParamBounds("weibull", data = field, response = "TNF")</pre>
```

```
TNF <- parasiteLoad::analyse(data = field,</pre>
                        response = "TNF",
                        model = "weibull",
                        group = "Sex")
## [1] "Analysing data for response: TNF"
## [1] "Fit for the response: TNF"
## [1] "Fitting for all"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupA : F"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
```

```
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting for groupB : M"
## [1] "Fitting model basic without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model basic with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced without alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Fitting model advanced with alpha"
## Warning in bbmle::mle2(response ~ dweibull(shape = myshape, scale =
## MeanLoad(L1, : some parameters are on the boundary: variance-covariance
## calculations based on Hessian may be unreliable
## [1] "Did converge"
## [1] "Testing HO no alpha vs alpha"
     dLL dDF
                pvalue
## 1 0.13
           1 0.6150504
## [1] "Testing H1 no alpha vs alpha"
               pvalue
## dLL dDF
## 1 0.1
          1 0.6559595
## [1] "Testing H2 groupA no alpha vs alpha"
     dLL dDF
              pvalue
           1 0.876648
## 1 0.01
## [1] "Testing H2 groupB no alpha vs alpha"
##
     dLL dDF
                pvalue
## 1 0.38
           1 0.3810045
## [1] "Testing H3 groupA no alpha vs alpha"
               pvalue
## dLL dDF
## 1 0
           1 0.9271205
## [1] "Testing H3 groupB no alpha vs alpha"
     dLL dDF
                pvalue
           1 0.4206925
## 1 0.32
## [1] "Testing H1 vs H0"
##
     dLL dDF
                pvalue
## 1 0.03
           1 0.8035592
## [1] "Testing H2 vs H0"
     dLL dDF
                pvalue
## 1 0.27
           3 0.9097406
```

```
## [1] "Testing H3 vs H1"
                 pvalue
##
      dLL dDF
## 1 1.16
            4 0.6784149
## [1] "Testing H3 vs H2"
      dLL dDF
                 pvalue
## 1 0.92
            2 0.3998039
##A11
print(TNF)
## $HO
##
## Call:
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
  Coefficients:
##
##
                     alpha
                               myshape
## 20.96793671 0.03843686 5.00000000
##
## Log-likelihood: -897.06
## Best method: bobyqa
##
## $H1
##
## Call:
##
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
  Coefficients:
                        L2
##
                                  alpha
                                            myshape
  21.04243399 20.88091647 0.03477237 5.00000000
##
## Log-likelihood: -897.03
## Best method: bobyqa
##
## $H2
## $H2$groupA
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibul1(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
```

##

```
myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
            T.1
                     alpha
                               myshape
  20.69112183 -0.01680606 5.00000000
##
## Log-likelihood: -441.49
## Best method: bobyqa
## $H2$groupB
##
## Call:
##
  bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
       scale = MeanLoad(L1, L1, alpha, HI)/gamma(1 + (1/myshape))),
       start = start, method = config$method, optimizer = config$optimizer,
##
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], alpha = paramBounds[["alphaLB"]],
##
           myshape = paramBounds[["myshapeLB"]]), upper = c(L1 = paramBounds[["L1UB"]],
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
  Coefficients:
##
            L1
                     alpha
                               myshape
## 21.25337880 0.09463402 5.00000000
## Log-likelihood: -455.3
  Best method: bobyqa
##
##
##
## $H3
  $H3$groupA
##
## Call:
   bbmle::mle2(minuslog1 = response ~ dweibull(shape = myshape,
##
##
       scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
       start = start, method = config$method, optimizer = config$optimizer,
##
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
##
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
##
       control = config$control)
##
##
   Coefficients:
##
                        L2
                                  alpha
                                            myshape
##
  20.42062926 21.11497687 0.01036138 5.00000000
##
## Log-likelihood: -441.22
## Best method: bobyqa
## $H3$groupB
##
## Call:
## bbmle::mle2(minuslog1 = response ~ dweibul1(shape = myshape,
```

```
scale = MeanLoad(L1, L2, alpha, HI)/gamma(1 + (1/myshape))),
##
##
       start = start, method = config$method, optimizer = config$optimizer,
       data = data, lower = c(L1 = paramBounds[["L1LB"]], L2 = paramBounds[["L2LB"]],
##
##
           alpha = paramBounds[["alphaLB"]], myshape = paramBounds[["myshapeLB"]]),
       upper = c(L1 = paramBounds[["L1UB"]], L2 = paramBounds[["L2UB"]],
##
##
           alpha = paramBounds[["alphaUB"]], myshape = paramBounds[["myshapeUB"]]),
##
       control = config$control)
##
##
  Coefficients:
##
            L1
                        L2
                                  alpha
                                            myshape
  21.81181965 20.77037818
                            0.08825218
                                        5.00000000
##
## Log-likelihood: -454.65
## Best method: bobyqa
bananaPlot(mod = TNF$HO,
             data = field,
             response = "TNF",
             group = "Sex") +
  scale_fill_manual(values = c("blue", "red")) +
  scale_color_manual(values = c("blue", "red")) +
 theme_bw()
```

Scale for 'fill' is already present. Adding another scale for 'fill', which
will replace the existing scale.

