Charactersing effect of anaemia on mortality in severe malaria

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Background

This looks at the severe malaria legacy dataset from MORU

Imputation of missing variables

Quite a lot of the important covariates are missing in the older studies. We use linear regression to estimate these unknown variables:

- Mising base deficit is imputed using bicarbonate (if available) else using respiratory rate
- Missing Blood urea nitrogen is imputed using creatinine

```
Impute base deficit from bicarbonate
```

```
BD_and_bicarbonate = !is.na(Leg_data$BD) & !is.na(Leg_data$bicarbonate)

print(paste('We have ', sum(BD_and_bicarbonate), 'observations for both bicarbonate and base deficit'))

## [1] "We have 5067 observations for both bicarbonate and base deficit"

mod_impute1 = lmer(BD ~ bicarbonate + (1 | studyID) + (1 | country), data= Leg_data[BD_and_bicarbonate, missing_BD = is.na(Leg_data$BD)

Available_Bicarbonate = !is.na(Leg_data$bicarbonate)

print(paste(sum(missing_BD & Available_Bicarbonate), 'observations will now be imputed'))

## [1] "309 observations will now be imputed"

# impute with model

Leg_data$BD[missing_BD & Available_Bicarbonate] = predict(mod_impute1,newdata=Leg_data[missing_BD & Ava

Impute base deficit from lactate

BD_and_lactate = !is.na(Leg_data$BD) & !is.na(Leg_data$lactate)

print(paste('We have ', sum(BD_and_lactate), 'observations for both lactate and base deficit'))
```

[1] "We have 632 observations for both lactate and base deficit"

```
if(length(unique(Leg_data$studyID[BD_and_lactate]))==1){
  mod_impute2 = lm(BD ~ lactate, data= Leg_data[BD_and_lactate,])
} else {
 mod_impute2 = lmer(BD ~ lactate + (1 | studyID), data= Leg_data[BD_and_lactate,])
}
missing_BD = is.na(Leg_data$BD)
Available_Lactate = !is.na(Leg_data$lactate)
print(paste(sum(missing BD & Available Lactate), 'observations will now be imputed'))
## [1] "722 observations will now be imputed"
# impute with model
Leg data$BD[missing BD & Available Lactate] = predict(mod impute2, newdata=Leg data[missing BD & Availab
Impute base deficit from respiratory rate
BD and rr = !is.na(Leg data$BD) & !is.na(Leg data$rr)
print(paste('We have ', sum(BD_and_rr), 'observations for both resp rate and base deficit'))
## [1] "We have 7572 observations for both resp rate and base deficit"
mod_impute3 = lmer(BD ~ rr + (1 | studyID), data= Leg_data[BD_and_rr,])
missing_BD = is.na(Leg_data$BD)
Available_rr = !is.na(Leg_data$rr)
print(paste(sum(missing_BD & Available_rr), 'observations will now be imputed'))
## [1] "1650 observations will now be imputed"
Leg_data$BD[missing_BD & Available_rr] = predict(mod_impute3,newdata=Leg_data[missing_BD & Available_rr
Impute blood urea nitrogen from creatinine:
BUN_and_cr = !is.na(Leg_data$BUN) & !is.na(Leg_data$creatinine)
print(paste('We have ', sum(BUN_and_cr), 'observations for both blood urea nitrogen and creatinine'))
## [1] "We have 1453 observations for both blood urea nitrogen and creatinine"
mod_impute4 = lmer(BUN ~ creatinine + (1 | studyID), data= Leg_data[BUN_and_cr,])
missing_BUN = is.na(Leg_data$BUN)
Available_cr = !is.na(Leg_data$creatinine)
print(paste(sum(missing_BUN & Available_cr), 'observations will now be imputed'))
## [1] "679 observations will now be imputed"
Leg_data$BUN[missing_BUN & Available_cr] = predict(mod_impute4,newdata=Leg_data[missing_BUN & Available
Resulting data we can now use: The contributions of the different studies:
vars_interest = c('outcome', 'HCT', 'LPAR_pct', 'BD', 'BUN', 'poedema',
                  'convulsions','coma','AgeInYear','drug_class')
complete_cases = apply(Leg_data[,vars_interest], 1, function(x) sum(is.na(x))) == 0
Complete_Leg_data = Leg_data[complete_cases,] # for the model fitting
Complete_Leg_data$studyID = as.factor(as.character(Complete_Leg_data$studyID))
# Whole dataset
table(Leg data$studyID)
##
##
            AAV
                          ΑQ
                                 AQGambia
                                                AQUAMAT Core Malaria
##
            370
                         560
                                      579
                                                   5494
                                                                1122
##
      SEAQUAMAT
```

```
##
           1461
# in the complete dataset (all variables recorded)
table(Complete_Leg_data$studyID)
##
##
            AAV
                          ΑQ
                                  AQGambia
                                                AQUAMAT Core Malaria
##
            214
                          150
                                       168
                                                   3666
      SEAQUAMAT
##
##
           1333
Complete_Leg_data$drug_AS = 0
Complete Leg data$drug AS[Complete Leg data$drug class=='artemisinin']=1
# remove infinite log parasitaemias
ind_keep = !(is.infinite(Complete_Leg_data$LPAR_pct) | is.nan(Complete_Leg_data$LPAR_pct))
Complete_Leg_data = Complete_Leg_data[ind_keep,]
Data summaries
Africa = c('The Gambia', 'Mozambique', 'Ghana', 'Kenya', 'Nigeria', 'Tanzania', 'Uganda', 'Rwanda', 'Congo')
Asia = c('Thailand','Vietnam','Bangladesh','Myanmar','India','Indonesia')
writeLines(paste('Children in Africa:',
                 sum(Complete_Leg_data$AgeInYear < 15 & Complete_Leg_data$country %in% Africa)))</pre>
## Children in Africa: 3779
writeLines(paste('Adults in Africa:',
                 sum(Complete_Leg_data$AgeInYear >= 15 & Complete_Leg_data$country %in% Africa)))
## Adults in Africa: 44
writeLines(paste('Children in Asia:',
                 sum(Complete_Leg_data$AgeInYear < 15 & Complete_Leg_data$country %in% Asia)))</pre>
## Children in Asia: 198
writeLines(paste('Adults in Asia:',
                 sum(Complete_Leg_data$AgeInYear >= 15 & Complete_Leg_data$country %in% Asia)))
## Adults in Asia: 2095
Exploratory analysis
```

```
for(s in unique(Complete_Leg_data$studyID)){
   print(paste(s, ', mortality of:', round(100*mean(Complete_Leg_data$outcome[Complete_Leg_data$studyID=
}

## [1] "Core Malaria , mortality of: 23 %"

## [1] "AQGambia , mortality of: 12 %"

## [1] "AAV , mortality of: 12 %"

## [1] "SEAQUAMAT , mortality of: 18 %"

## [1] "AQUAMAT , mortality of: 9 %"

## [1] "AQ , mortality of: 23 %"
```

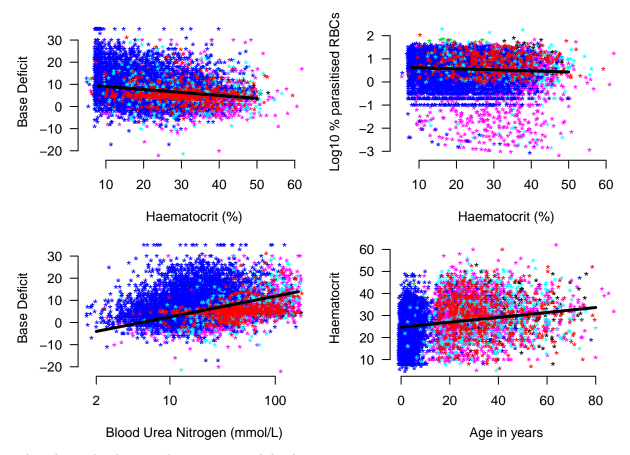
```
for(s in unique(Complete_Leg_data$studyID)){
  print(paste0(s, ', ages:', round(quantile(Complete_Leg_data$AgeInYear[Complete_Leg_data$studyID==s],
}
## [1] "Core Malaria, ages:1 Core Malaria, ages:27 Core Malaria, ages:75"
## [1] "AQGambia, ages:1 AQGambia, ages:4 AQGambia, ages:9"
## [1] "AAV, ages:15 AAV, ages:34 AAV, ages:77"
## [1] "SEAQUAMAT, ages:2 SEAQUAMAT, ages:25 SEAQUAMAT, ages:87"
## [1] "AQUAMAT, ages:0 AQUAMAT, ages:2 AQUAMAT, ages:78"
## [1] "AQ, ages:15 AQ, ages:30 AQ, ages:74"
for(s in unique(Complete_Leg_data$studyID)){
  print(s)
  print(table(Complete_Leg_data$drug[Complete_Leg_data$studyID==s]))
}
   [1] "Core Malaria"
##
##
                   Artemether
                                             Chloroquine Lumefantrine
    Amodiaquine
                                Artesunate
##
                           11
                                        368
                                                        2
##
     Mefloquine
                          NAC
                                    Quinine
##
                            6
                                        262
   [1] "AQGambia"
##
##
##
    Amodiaquine
                   Artemether
                                 Artesunate
                                             Chloroquine Lumefantrine
##
                           82
                          NAC
##
     Mefloquine
                                    Quinine
##
                            0
                                         86
   [1] "AAV"
##
##
                                 Artesunate
##
                   Artemether
                                             Chloroquine Lumefantrine
    Amodiaquine
##
                          102
                                        112
##
                          NAC
     Mefloquine
                                    Quinine
                            0
##
                                          0
##
   [1] "SEAQUAMAT"
##
##
    Amodiaquine
                   Artemether
                                 Artesunate
                                             Chloroquine Lumefantrine
##
              0
                            0
                                        645
                                                        0
##
     Mefloquine
                          NAC
                                    Quinine
##
              0
                            0
                                        628
##
   [1] "AQUAMAT"
##
##
    Amodiaquine
                   Artemether
                                 Artesunate
                                             Chloroquine Lumefantrine
##
              0
                            0
                                       1837
                                                        0
     Mefloquine
##
                          NAC
                                    Quinine
##
              0
                            0
                                       1818
   [1] "AQ"
##
##
##
    Amodiaquine
                   Artemether
                                 Artesunate
                                             Chloroquine Lumefantrine
                                                        0
                                                                      0
##
                           73
                                          0
                          NAC
##
     Mefloquine
                                    Quinine
                            0
                                         77
##
```

Let's look at the key predictive variables. We use a random effects term to model differences between studies.

```
## Linear mixed model fit by REML ['lmerMod']
```

```
## Formula: BD ~ HCT + (1 | studyID/country)
##
     Data: Complete_Leg_data
##
## REML criterion at convergence: 40261.9
## Scaled residuals:
             10 Median
                               30
                                       Max
## -4.4421 -0.6612 -0.1488 0.5224 4.7209
##
## Random effects:
## Groups
                    Name
                               Variance Std.Dev.
## country:studyID (Intercept) 2.6525 1.6286
## studyID
                    (Intercept) 0.8373 0.9151
## Residual
                                41.8947 6.4726
## Number of obs: 6116, groups: country:studyID, 18; studyID, 6
##
## Fixed effects:
               Estimate Std. Error t value
## (Intercept) 10.339058
                          0.653393
              -0.133548
                          0.009699 - 13.77
##
## Correlation of Fixed Effects:
##
       (Intr)
## HCT -0.394
## Linear mixed model fit by REML ['lmerMod']
## Formula: LPAR_pct ~ HCT + (1 | studyID/country)
##
     Data: Complete_Leg_data
##
## REML criterion at convergence: 13822.9
## Scaled residuals:
##
              1Q Median
      Min
                               3Q
                                       Max
## -4.7144 -0.5555 0.1598 0.7265 2.4355
##
## Random effects:
## Groups
                               Variance Std.Dev.
                    Name
## country:studyID (Intercept) 0.00946 0.09726
                    (Intercept) 0.07496 0.27379
## studyID
                               0.55564 0.74541
## Residual
## Number of obs: 6116, groups: country:studyID, 18; studyID, 6
##
## Fixed effects:
               Estimate Std. Error t value
## (Intercept) 0.659944
                          0.121244
                                    5.443
## HCT
              -0.004579
                          0.001116 -4.105
##
## Correlation of Fixed Effects:
       (Intr)
## HCT -0.251
## Linear mixed model fit by REML ['lmerMod']
## Formula: BD ~ log10(BUN) + (1 | studyID/country)
##
     Data: Complete_Leg_data
##
```

```
## REML criterion at convergence: 39236.2
##
## Scaled residuals:
      Min 1Q Median
                             ЗQ
                                     Max
## -5.6063 -0.6369 -0.1041 0.5191 5.0754
##
## Random effects:
## Groups
                               Variance Std.Dev.
                   Name
## country:studyID (Intercept) 2.876 1.696
## studyID
              (Intercept) 6.858
                                      2.619
## Residual
                              35.405
                                      5.950
## Number of obs: 6116, groups: country:studyID, 18; studyID, 6
## Fixed effects:
##
              Estimate Std. Error t value
## (Intercept) -6.8409
                       1.2574
                                  -5.44
## log10(BUN)
                9.3530
                          0.2559
                                   36.55
##
## Correlation of Fixed Effects:
             (Intr)
## log10(BUN) -0.293
## Linear mixed model fit by REML ['lmerMod']
## Formula: HCT ~ AgeInYear + (1 | studyID/country)
##
     Data: Complete_Leg_data
##
## REML criterion at convergence: 43534.9
##
## Scaled residuals:
      Min
              1Q Median
## -3.1004 -0.7399 -0.0515 0.6927 3.5627
## Random effects:
## Groups
                   Name
                               Variance Std.Dev.
## country:studyID (Intercept) 5.722
                                       2.392
## studyID
              (Intercept) 7.322
                                       2.706
                                      8.454
## Residual
                              71.467
## Number of obs: 6116, groups: country:studyID, 18; studyID, 6
##
## Fixed effects:
              Estimate Std. Error t value
## (Intercept) 24.69246
                          1.36141 18.137
## AgeInYear
             0.11159
                          0.01159
                                  9.626
##
## Correlation of Fixed Effects:
            (Intr)
## AgeInYear -0.185
```

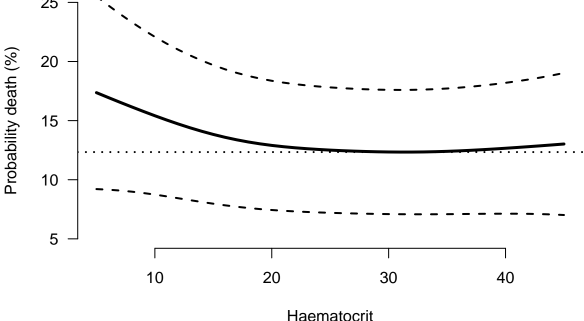


The relationship between haematocrit and death:

par(las=1, bty='n')

```
modHCT=gam(outcome ~ s(HCT) + s(studyID, bs='re') + s(country, bs='re'),data = Complete_Leg_data, famile
summary(modHCT)
##
## Family: binomial
## Link function: logit
##
## Formula:
  outcome ~ s(HCT) + s(studyID, bs = "re") + s(country, bs = "re")
##
##
  Parametric coefficients:
##
##
               Estimate Std. Error z value Pr(>|z|)
  (Intercept) -1.8865
                                     -7.87 3.54e-15 ***
                            0.2397
##
##
                  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
##
## Approximate significance of smooth terms:
                 edf Ref.df Chi.sq p-value
##
               2.304 2.922
                              5.482 0.15478
## s(HCT)
## s(studyID) 3.611 5.000 314.182 0.00361 **
## s(country) 10.766 14.000 162.027 0.01464 *
##
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Complete_Leg_data\$country=as.factor(Complete_Leg_data\$country)



Predictive value of anaemia on death adjusting for confounders

Before fitting the more complex GAM models we explore the standard glm (logistic regression) models.

```
##
      Data: Complete_Leg_data
##
##
        AIC
                 BIC
                       logLik deviance df.resid
##
     3459.7
              3540.3 -1717.8
                                3435.7
                                           6104
##
## Scaled residuals:
       Min
                10 Median
                                30
## -3.9034 -0.3324 -0.1914 -0.1076 15.4072
##
## Random effects:
## Groups Name
                        Variance Std.Dev.
## country (Intercept) 1.501e-01 3.875e-01
## studyID (Intercept) 1.919e-09 4.381e-05
## Number of obs: 6116, groups: country, 15; studyID, 6
##
## Fixed effects:
##
                 Estimate Std. Error z value Pr(>|z|)
               -7.000057
                            0.306929 -22.807 < 2e-16 ***
## (Intercept)
                            0.005284
                                       3.111 0.001863 **
## HCT
                 0.016441
## LPAR pct
                -0.001281
                            0.060471 -0.021 0.983095
## AgeInYear
                 0.013715
                           0.003840
                                      3.571 0.000355 ***
                 1.338046
                            0.100906 13.260 < 2e-16 ***
## coma
## convulsions1 0.513532
                            0.116864
                                       4.394 1.11e-05 ***
## poedema1
                 0.543720
                            0.385373
                                       1.411 0.158276
## log10(BUN)
                 1.778368
                            0.166012 10.712 < 2e-16 ***
## BD
                 0.121719
                            0.007183 16.944 < 2e-16 ***
## drug_AS
                -0.343604
                            0.090337 -3.804 0.000143 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
##
               (Intr) HCT
                             LPAR_p AgInYr coma
                                                  cnvls1 poedm1 110(BU BD
## HCT
               -0.483
               -0.041 0.030
## LPAR_pct
                0.039 -0.178
                              0.001
## AgeInYear
## coma
               -0.163 -0.027 0.075 0.000
## convulsins1 -0.133 -0.073 0.017 0.108 -0.220
## poedema1
               -0.004 -0.005 -0.006 -0.048 0.027
                                                   0.000
## log10(BUN)
             -0.700 0.064 -0.047 -0.245 -0.014
                                                   0.103 0.006
## BD
               -0.148   0.199   -0.180   0.135   -0.024   0.024   -0.008   -0.262
               -0.091 -0.012 -0.024 -0.022 0.007 0.004 -0.025 -0.044 -0.020
## drug AS
## convergence code: 0
## Model failed to converge with max|grad| = 0.00101495 (tol = 0.001, component 1)
Now let's make counterfactual predictions of anaemia on death for the patients in the database.
myquantiles = c(0.25,0.5,0.75) # this is 50% predictive interval
overall_median_mortality = median(100*predict(mod_full_GLM, type='response'))
par(las=1, bty='n')
x_hcts = seq(4,45, by=1)
probs_lin = array(dim = c(3, length(x_hcts)))
for(i in 1:length(x_hcts)){
  mydata = Complete_Leg_data
 mydata$HCT=x_hcts[i]
```

```
ys = 100*predict(mod_full_GLM, newdata = mydata, re.form=NA, type='response')
probs_lin[,i] = quantile(ys, probs=myquantiles)
}
```

The way to interpret this 'counterfactual' plot is as follows: suppose that every individual in the dataset was assigned (as in a intervention) a specific haematocrit X, what would the resulting per patient probability of death be. Here we summarise these probabilities by the predicted mean probability of death and 80% predictive intervals.

```
plot(x_hcts,probs_lin[2,], xlim=c(4,45), ylab='Predicted probability of death',
     xlab='Haematocrit (%)', ylim=c(0,20), lty=1, lwd=3, type='1')
lines(x_hcts, probs_lin[1,], lty=2, lwd=2)
lines(x hcts, probs lin[3,], lty=2, lwd=2)
abline(h=overall_median_mortality, lwd=3, col='blue',lty=2)
legend('topleft', col=c('black', 'black', 'blue'), lwd=3, lty=c(1,2,2),
       legend = c('Median counterfactual value', '25th and 75th counterfactual quantiles', 'Retrodicted :
      20
                    Median counterfactual value
Predicted probability of death
                    25th and 75th counterfactual quantiles
                    Retrodicted median value
      2
      9
      2
      0
                        10
                                         20
                                                          30
                                                                            40
```

More complex GAM model

The GAM model allows for non-linear relationships between certain variables and the outcome.

Here we fit as non-linear the effect of age and haematocrit on mortality. We add a random effect term for the studyID We should also be doing this for the study site...

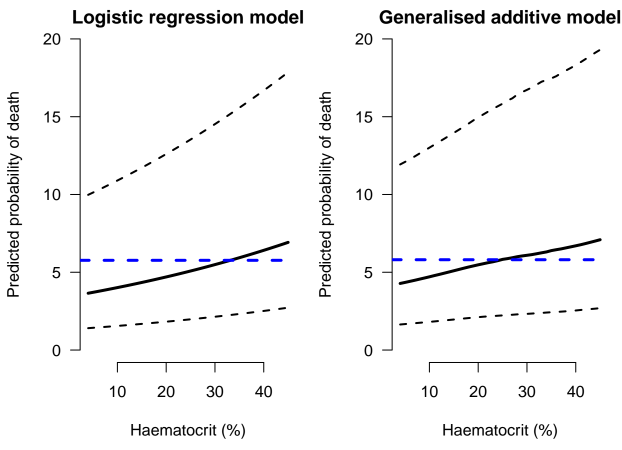
Haematocrit (%)

```
##
## Formula:
## outcome ~ s(HCT, AgeInYear) + LPAR pct + coma + convulsions +
      poedema + log10(BUN) + BD + drug_AS + s(studyID, bs = "re") +
##
##
      s(country, bs = "re")
##
## Parametric coefficients:
##
                Estimate Std. Error z value Pr(>|z|)
## (Intercept) -6.327596 0.270364 -23.404 < 2e-16 ***
## LPAR_pct
                0.001766
                         0.060455
                                     0.029 0.976696
## coma
                1.330827
                          0.100888 13.191 < 2e-16 ***
                                     4.557 5.2e-06 ***
## convulsions1 0.534704
                          0.117346
## poedema1
                0.547871
                          0.384139
                                     1.426 0.153801
## log10(BUN)
                1.701401
                          0.170552 9.976 < 2e-16 ***
## BD
                0.123330
                          0.007331 16.824 < 2e-16 ***
## drug_AS
               -0.343908
                           0.090360 -3.806 0.000141 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Approximate significance of smooth terms:
##
                        edf Ref.df Chi.sq p-value
## s(HCT, AgeInYear) 5.487067 7.664 33.634 3.60e-05 ***
                   0.004292 5.000 0.003
## s(studyID)
                                            0.496
                   9.942725 14.000 74.592 2.79e-15 ***
## s(country)
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## R-sq.(adj) =
               0.27
                        Deviance explained = 29.1%
## UBRE = -0.43785 Scale est. = 1
                                         n = 6116
```

Now we compute the corresponding counterfactual probabilities of death for the dataset for all values of the haematocrit:

```
overall_median_mortalityGAM = median(100*predict(mod_full_GAM, type='response'))
par(las=1, bty='n')
probs_gam = array(dim = c(3, length(x_hcts)))
for(i in 1:length(x_hcts)){
   mydata = Complete_Leg_data
   mydata$HCT=x_hcts[i]
   ys = 100*predict(mod_full_GAM, newdata = mydata, type='response')
   probs_gam[,i] = quantile(ys, probs=myquantiles)
}
```

We see that the effect of haematocrit on mortality is non-linear under this model: below 20 is protective, above 20 plateaus out:



Model comparison

```
Which model is better fit in terms of AIC
```