Classification Model on Antiretroviral Therapy Reaction and Failure Developed on the Unique Records of the Akwa Ibom HIV Database

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Introduction

This undertaking is part of the capstone in the Professional Certificate Program in Data Science of Harvard Online. The corresponding R Markdown and R files are in the GitHub of nsmacaspac.

Unique Records of the Akwa Ibom HIV Database

In a previous study on patient response to antiretroviral therapy, Ekpenyong, Etebong, and Jackson (2019, 3) used a database of patients who received treatment for HIV from thirteen health centers in Akwa Ibom, Nigeria, between 2015 and 2018. Two years later, they published the processed dataset (Ekpenyong et al. 2021b, Appendix) with minor oversight in the accompanying article, which were easily reconciled through the 2019 study and were appropriately referenced throughout this project. The processed dataset was composed of an Individual Treatment Change Episodes table with a column for each antiretroviral drug administered and a concatenated Unique Records table with the drugs combined into a column for each antiretroviral therapy of three drugs administered. For the purpose of this project, we utilize only the Unique Records table.

The Unique Records table was imported with the corresponding read_xlsx function in the language R. The dataset was composed of 1,056 patient records, each with 15 columns: patient identification, sex, baseline CD4 count, follow-up CD4 count, baseline RNA load, follow-up RNA load, baseline weight, followup weight, drug combination, and patient response and drug reaction classifications 1 to 5 (fig. 1). The immunological marker CD4 counts were given in cells per cubic millimeter (Ekpenyong et al. 2021a, 8). The viral RNA loads were expressed in times 10² copies (Ekpenyong, Etebong, and Jackson 2019, 10). The weights ranged from 4.7 to 125 kg on account of the presence of pediatric patients (Ekpenyong, Etebong, and Jackson 2019, 2). The three-drug combinations of antiretroviral therapy were a complementary mix of nucleoside reverse transcriptase inhibitors tenofovir (TDF), lamivudine (3TC) and zidovudine (AZT), and non-nucleoside reverse transcriptase inhibitors efavirenz (EFV) and nivarapine (NVP) given in the first 6 months of treatment (Ekpenyong et al. 2021a, 8). Patient response and drug reaction were quantified and classified in the 2019 study using the advanced method of interval type-2 fuzzy logic system. The drug reaction classification used a binary system to indicate very high interaction (C1), high interaction (C2), low interaction (C3), very low interaction (C4), and no interaction (C5; Ekpenyong, Etebong, and Jackson 2019, 11). Very high and high interactions signified treatment failure as well (Ekpenyong, Etebong, and Jackson 2019, 10).

Tidy Dataset

The dataset was rendered into tidy format to prepare it for preprocessing. The fifteen columns were renamed consistently with their aforementioned descriptions, with vhi_tf corresponding to very high interac-

Unique Records										[Target Classes]]
PID	SEX	BCD4	FCD4	BRNA	FRNA	BWt(kg)	FWt(kg)	DRUGCOMB	PR	C1	C2	C3	C4	C5
1	F	148	106	3	1.3	42	43	TDF+3TC+EFV	53.56	0	0	1	0	0
2	F	145	378	2.5	1.3	57	60	AZT+3TC+NVP	55.33	0	0	0	1	0
3	M	78	131	4.1	1.7	70	75	AZT+3TC+NVP	50.00	0	1	0	0	0
4	M	295	574	4.4	1.9	64	66	AZT+3TC+NVP	50.00	0	0	1	0	0
5	F	397	792	1.9	1.3	52	55	AZT+3TC+NVP	76.00	0	0	0	0	1

Figure 1: First rows of the Unique Records table.

tion treatment failure and ni corresponding to no interaction. Missing values were not detected.

```
colnames(dataset) <- c("id", "sex", "bcd4", "fcd4", "brna", "frna", "bweight", "fweight", "therapy", "r</pre>
head(dataset, n = 5)
     id sex bcd4 fcd4 brna frna bweight fweight
                                                       therapy response vhi_tf hi_tf
          F
             148
                  106
                        3.0
                             1.3
                                       42
                                               43 TDF+3TC+EFV 53.56199
      2
          F
                        2.5
                             1.3
                                       57
                                                60 AZT+3TC+NVP 55.33422
                                                                              0
             145
                   378
                                                                                     0
      3
          Μ
              78
                   131
                        4.1
                             1.7
                                       70
                                               75 AZT+3TC+NVP 50.00000
                                                                              0
                                                                                     1
                                       64
                                               66 AZT+3TC+NVP 50.00000
                                                                                     0
     4
          Μ
             295
                   574
                        4.4
                             1.9
                                                                              0
     5
          F
             397
                   792
                        1.9
                             1.3
                                       52
                                               55 AZT+3TC+NVP 76.00000
                                                                                     0
     li vli ni
## 1
     1
          0
             0
## 2
     0
          1
             0
## 3
     0
          0
             0
## 4
      1
          0
## 5 0
          0
```

The brna and frna columns were multiplied by 10^2 to simplify the unit from times 10^2 copies to just copies. This aligns them with the unit used for viral RNA load in the WHO definition of HIV (World Health Organization, n.d.).

The vhi_tf, hi_tf, li, vli, and ni columns were verified to have only one value per row. Hence, the binary system was relabeled as vhi_tf to ni using the case_when function and merged under a newly defined dreaction column. This brought down the number of columns to eleven.

```
dataset1 <- dataset |>
  mutate(brna = brna*10^2) |>
  mutate(frna = frna*10^2) |> # simplifies the unit from times 10^2 copies to just copies
  mutate(dreaction = case_when(vhi_tf == 1 ~ "vhi_tf",
                               hi_tf == 1 ~ "hi_tf",
                               li == 1 ~ "li",
                               vli == 1 ~ "vli"
                               ni == 1 ~ "ni")) |> # relabels drug reactions as vhi_tf to ni and merges
  select(-vhi tf, -hi tf, -li, -vli, -ni)
head(dataset1, n = 5)
     id sex bcd4 fcd4 brna frna bweight fweight
                                                      therapy response dreaction
     1
            148
                  106
                       300
                            130
                                      42
                                              43 TDF+3TC+EFV 53.56199
                                                                               1i
      2
          F
             145
                  378
                       250
                             130
                                      57
                                              60 AZT+3TC+NVP 55.33422
                                                                              vli
      3
                                      70
          Μ
              78
                  131
                        410
                             170
                                              75 AZT+3TC+NVP 50.00000
                                                                           hi_tf
      4
             295
                  574
                       440
                             190
                                      64
                                              66 AZT+3TC+NVP 50.00000
                                                                               1i
          Μ
                             130
                                      52
                                              55 AZT+3TC+NVP 76.00000
             397
                  792
                       190
                                                                               ni
```

Preprocessed Dataset

The sex, therapy, and dreaction variables of the dataset were changed from character to numeric class using the factor and as numeric functions to allow for numerical data examination. All eleven variables showed good data variability.

??? DISTRIBUTION

The correlation coefficients between variables were visualized using the corrplot function (fig. 2; Wei and Simko 2021). Patient response and drug reaction highly correlated with each other as expected as they refer to similar information. Hence, only drug reaction was retained as an outcome for the purpose of this project. On the other hand, CD4 count was negatively correlated with drug reaction whereas RNA load was positively correlated with it. This means that the bcd4, fcd4, brna and frna variables contained information relevant to drug reaction and were kept as predictors.

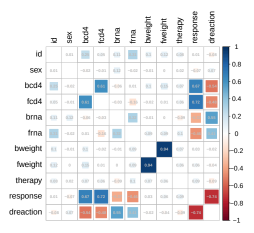


Figure 2: Matrix of the correlation coefficients between variables.

The final dataset was streamlined to contain only these four predictors and one outcome for faster classification modeling.

```
dataset2 <- dataset1 |>
  select(bcd4, fcd4, brna, frna, dreaction) |> # keeps CD4 count and RNA load as predictors and drug re
  mutate(dreaction = factor(dreaction, c("ni", "vli", "li", "hi_tf", "vhi_tf")))
head(dataset2, n = 5)
     bcd4 fcd4 brna frna dreaction
##
      148
           106
                 300
                      130
                                 1i
                 250
                                 vli
      145
           378
                      130
   3
       78
           131
                 410
                      170
                              hi_tf
      295
           574
                 440
                      190
                                 1i
      397
           792
                190
                      130
                                 ni
```

Predictors

CD4 Count

RNA Load

Classification Models

• why this partition

k-Nearest Neighbor Model

· why this model

Recursive Partitioning and Regression Trees Model

Rborist Model

Quadratic Discriminant Analysis Model

Classification Model

Conclusion

-meaningful decisions on antiretroviral therapy administration

References

Ekpenyong, Moses E., Mercy E. Edoho, Ifiok J. Udo, Philip I. Etebong, Nseobong P. Uto, Tenderwealth C. Jackson, and Nkem M. Obiakor. 2021a. "A Transfer Learning Approach to Drug Resistance Classification in Mixed HIV Dataset." *Informatics in Medicine Unlocked* 24: 100568. https://doi.org/10.1016/j.imu.2021. 100568.

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Irizarry, Rafael A. 2002. Introduction to Data Science: Data Analysis and Prediction Algorithms with R. http://rafalab.dfci.harvard.edu/dsbook/.

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