## Eosinofiele Oesofagitis dataset EDA

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#### **Dataset**

TODO neocate en ffed Nutritional intake was assessed in 40 Dutch adult EoE patients participating in the Supplemental Elemental Trial (SET) using 3-day food diaries. In this randomized controlled trial, diagnosed patients received either a four-food elimination diet alone (FFED) or FFED with addition of an amino acid-based formula (Neocate) for 6 weeks. Disease severity was assessed by peak eosinophil count/high power field (PEC) in esophageal biopsy specimens. Multiple linear regression analyses were performed to assess associations between the intake of nutrients and foods per 1000 kCal and PEC, both at baseline and after the 6 weeks diet, while controlling for baseline variables. What do vars mean datatypes, units, values What is the dependent class variable

#### Introduction

#### Research question

Can the peak eosinophil baseline in individuals with eosinophilic oesophagitis be predicted with classical machine learning using the measured nutrient intake data?

#### Missing data

Missing columns TODO There are quite a bit of missing attributes and missing values in this data set. Luckily there are plenty of attributes that do have data from where the machine learning algorithm can make its prediction.

37 empty instances 180 empty attributes Lots of columns with more then 70 % missing values

```
FALSE
FALSE Attaching package: 'janitor'
FALSE The following objects are masked from 'package:stats':
FALSE
FALSE
          chisq.test, fisher.test
FALSE Loading required package: lattice
FALSE Loading required package: MASS
FALSE
FALSE Attaching package: 'memisc'
FALSE The following objects are masked from 'package:stats':
FALSE
FALSE
          contr.sum, contr.treatment, contrasts
FALSE The following object is masked from 'package:base':
FALSE
FALSE
          as.array
```

```
FALSE
```

FALSE Attaching package: 'ggplot2'

FALSE The following object is masked from 'package:memisc':

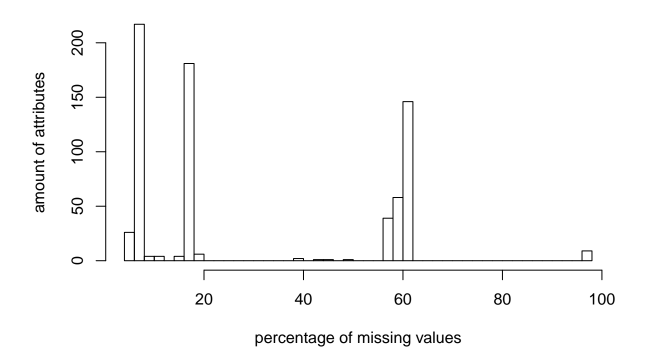
**FALSE** 

FALSE syms

FALSE Removing 37 empty rows of 79 rows total (46.8%).

FALSE Removing 180 empty columns of 879 columns total (Removed: Date, Eoscutoff, Lenght, Weight, BMI, A

### Amount of attributes with missing values



FALSE [1] "columns removed because too many missing values"

FALSE [1] 155

FALSE [1] "columns romved in total"

FALSE [1] 335

FALSE [1] 544

#### distrubution

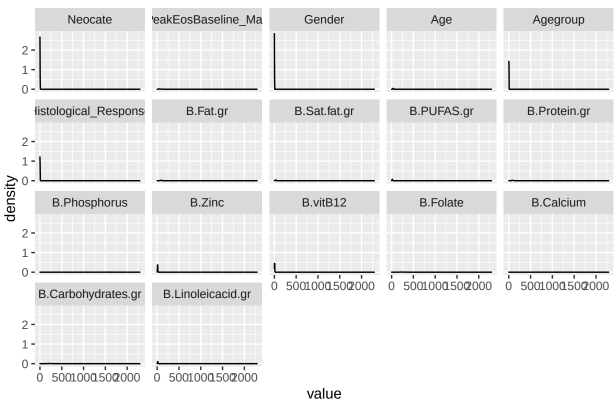
Most attributes seem normally distributed how ever their range differ quite significantly so a normalization step might be required other wise the range can bias the machine learning algorithm to specific attributes.

FALSE Removing 2 empty rows of 42 rows total (4.76%).

FALSE No empty columns to remove.

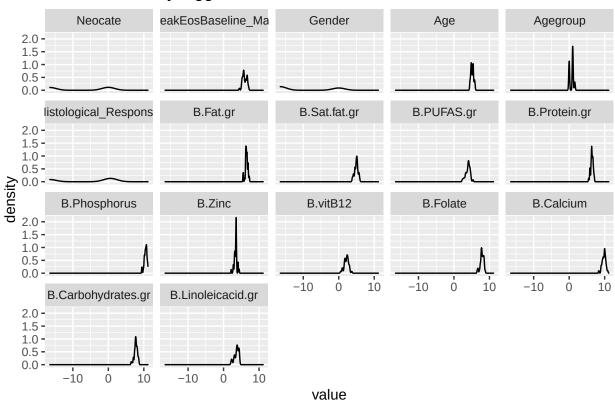
FALSE No id variables; using all as measure variables FALSE No id variables; using all as measure variables

### attributes density



FALSE No id variables; using all as measure variables

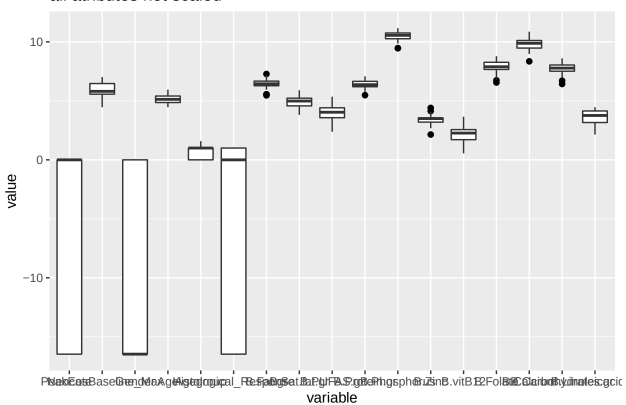
### attributes density logged



### Normalisation This is all attributes compared with no normalization, min-max and scaling normalization.

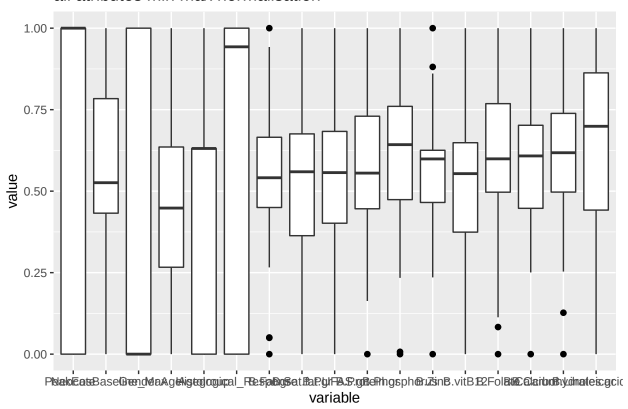
FALSE No id variables; using all as measure variables

### all atributes not scaled



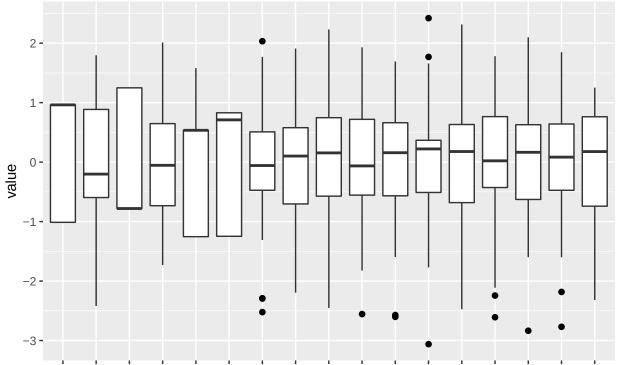
FALSE No id variables; using all as measure variables

### all atributes min max normalisation



FALSE No id variables; using all as measure variables

# all atributes scaling normalisation



### correlation matrix of the bones lenght and width

