## Pima Indian

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### Variables Used:

Pregnancy: Number of times pregnant. Glucose: Oral Glucose Tolerance Test.

Blood Pressure: Diastolic blood pressure (mm Hg) Skin Thickness: Triceps skin fold thickness (mm)

Insulin: 2-Hour serum insulin (mu U/ml)

BMI: Body mass index (weight in kg/(height in m)^2)
DiabetesPedigreeFunction: Diabetes pedigree function.

Age: Years of Age.

Outcome: 0 = Does Not Have Diabetes, 1 = Does Have Diabetes

## Preparing the Data for Use:

- 1. Adjusting Outcome Integer into a Factor.
- 2. Replacing 0's with NA values within the Insulin, BMI, and SkinThickness fields.
- 3. Creating a 'GlucoseGroup' variable based on labels set by the American Diabetes Association (https://diabetes.org/diabetes/a1c/diagnosis)
- 4. Creating a new variable 'WeightGroup' based on BMI value. Using bins set my the American Cancer Society (https://www.cancer.org/healthy/cancer-causes/diet-physical-activity/body-weight-and-cancer-risk/adult-bmi.html).

```
glimpse(PI)
```

```
## Rows: 768
## Columns: 9
## $ Pregnancies
                              <int> 6, 1, 8, 1, 0, 5, 3, 10, 2, 8, 4, 10, 10, 1, ~
                              <int> 148, 85, 183, 89, 137, 116, 78, 115, 197, 125~
## $ Glucose
## $ BloodPressure
                              <int> 72, 66, 64, 66, 40, 74, 50, 0, 70, 96, 92, 74~
## $ SkinThickness
                              <int> 35, 29, 0, 23, 35, 0, 32, 0, 45, 0, 0, 0, 0, ~
## $ Insulin
                              <int> 0, 0, 0, 94, 168, 0, 88, 0, 543, 0, 0, 0, 0, ~
## $ BMI
                              <dbl> 33.6, 26.6, 23.3, 28.1, 43.1, 25.6, 31.0, 35.~
## $ DiabetesPedigreeFunction <dbl> 0.627, 0.351, 0.672, 0.167, 2.288, 0.201, 0.2~
                              <int> 50, 31, 32, 21, 33, 30, 26, 29, 53, 54, 30, 3~
## $ Age
## $ Outcome
                              <int> 1, 0, 1, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 1, 1, ~
```

```
## Rows: 768
## Columns: 11
## $ Pregnancies
                              <int> 6, 1, 8, 1, 0, 5, 3, 10, 2, 8, 4, 10, 10, 1, ~
                              <int> 148, 85, 183, 89, 137, 116, 78, 115, 197, 125~
## $ Glucose
## $ GlucoseGroup
                              <fct> Diabetes, Normal, Diabetes, Normal, Diabetes,~
                              <dbl> 33.6, 26.6, 23.3, 28.1, 43.1, 25.6, 31.0, 35.~
## $ BMI
                              <fct> Obese, Overweight, NormalWeight, Overweight, ~
## $ WeightGroup
## $ BloodPressure
                              <int> 72, 66, 64, 66, 40, 74, 50, NA, 70, 96, 92, 7~
## $ SkinThickness
                              <int> 35, 29, NA, 23, 35, NA, 32, NA, 45, NA, NA, N~
## $ Insulin
                              <int> NA, NA, NA, 94, 168, NA, 88, NA, 543, NA, NA,~
## $ DiabetesPedigreeFunction <dbl> 0.627, 0.351, 0.672, 0.167, 2.288, 0.201, 0.2~
                              <int> 50, 31, 32, 21, 33, 30, 26, 29, 53, 54, 30, 3~
## $ Age
## $ Outcome
                              <fct> 1, 0, 1, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 1, 1, ~
```

Critique: This dataset only contains the diastolic blood pressure levels. In order to get a better picture of artery health, systolic blood pressure levels should have been included.

All 0 values were changed to NA values as they don't close to typical measurements in BMI, BloodPressure, Skin Thickness, or Insulin levels in any population. I assume they are missing values, instead.

# **Exploratory Data Analysis**

- First, looking at statistic summaries with skim()— keeping the data with NA values. The reason for skim() over boxplots is because after glancing at the data, I can only assume that standard deviations may be small for some variables. This prompts me to look the spread of the data numerically, rather than visually, in order to be precise.
- Second, looking at distributions for the samples that had Diabetes.
- Then, I look at distributions for the samples that did not have Diabetes.
- After creating distributions using facet\_wrap() and adjusting for a free x-scale, I find that there are either normal or rightskewed distributions only:
  - Right-Skewed: This is true for variables Age, Insulin, Pregnancies, and Pedigree Function.

```
skimr::skim(PI)
```

Data summary

Name	PI
Number of rows	768
Number of columns	11
Column type frequency:	
factor	3
numeric	8
Group variables	None

### Variable type: factor

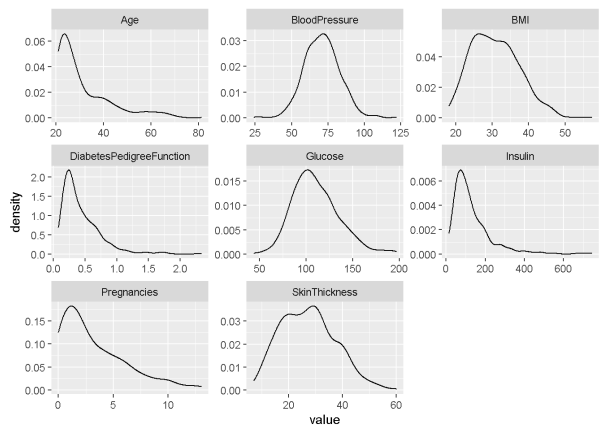
skim_variable	n_missing	complete_rate	ordered	n_unique	top_counts
GlucoseGroup	5	0.99	FALSE	3	Dia: 311, Pre: 260, Nor: 192
WeightGroup	11	0.99	FALSE	4	Obe: 472, Ove: 179, Nor: 102, Und: 4
Outcome	0	1.00	FALSE	2	0: 500, 1: 268

### Variable type: numeric

skim_variable	n_missing	complete_rate	mean	sd	p0	p25	p50	p75	p100	hist
Pregnancies	0	1.00	3.85	3.37	0.00	1.00	3.00	6.00	17.00	<b>—</b> —
Glucose	5	0.99	121.69	30.54	44.00	99.00	117.00	141.00	199.00	
BMI	11	0.99	32.46	6.92	18.20	27.50	32.30	36.60	67.10	
BloodPressure	35	0.95	72.41	12.38	24.00	64.00	72.00	80.00	122.00	
SkinThickness	227	0.70	29.15	10.48	7.00	22.00	29.00	36.00	99.00	
Insulin	374	0.51	155.55	118.78	14.00	76.25	125.00	190.00	846.00	
DiabetesPedigreeFunction	0	1.00	0.47	0.33	0.08	0.24	0.37	0.63	2.42	<b>—</b> ——
Age	0	1.00	33.24	11.76	21.00	24.00	29.00	41.00	81.00	<b>—</b> ——

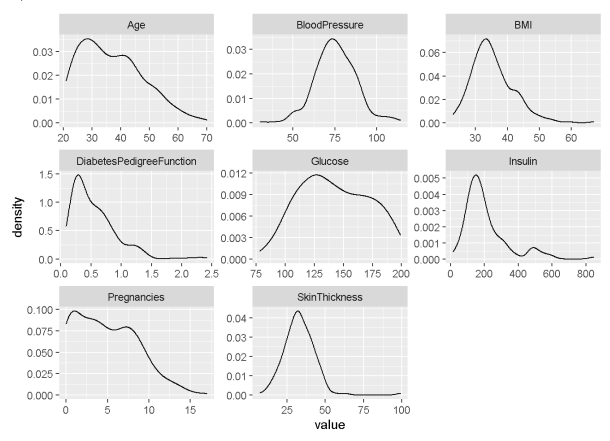
```
PI %>% filter(Outcome == 0) %>%
  purrr::keep(is.numeric) %>%
  gather() %>%
  ggplot(aes(value)) +
   facet_wrap(~ key, scales = "free") +
   geom_density()
```

## Warning: Removed 406 rows containing non-finite values (stat\_density).



```
PI %>% filter(Outcome == 1) %>%
  purrr::keep(is.numeric) %>%
  gather() %>%
  ggplot(aes(value)) +
   facet_wrap(~ key, scales = "free") +
   geom_density()
```

## Warning: Removed 246 rows containing non-finite values (stat\_density).



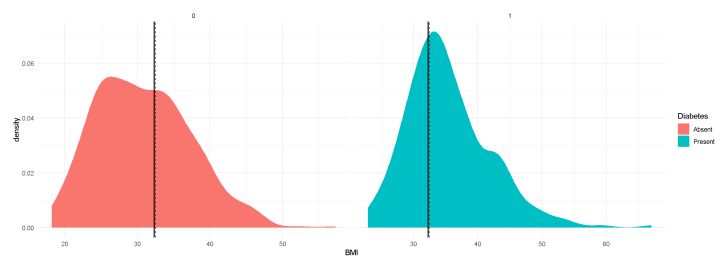
If we look at the skim()'s BMI histogram, it seems left-skewed which I can believe if we have more samples in which the person was negative for diabetes. Using skim()'s second output, I learn that there are 500 rows for those that did not have diabetes which definitely is more than 268 for those that did. Still, since BMI is a strong predictor of diabetes, I plot the distribution individually— assigning color red for the subset that have diabetes and for presentation purposes.

After doing so, I find that BMI statistics are actually normally distributed despite the right-skewed histogram found in skimr. I find that the mean is ~33 for both those with and without diabetes—indicating that the general population *is* very prone to the health condition. This can be confirmed if you perform research on the Arizona Pima Indians & their extraordinarily high rate of kidney disease and failure as a leading cause of death.

Insulin, on the other hand, *is* right-skewed. This means the samples tend to have higher Insulin values than lower Insulin values.

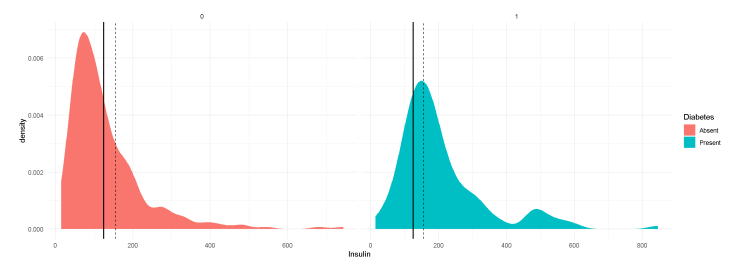
```
PI_BMI <- PI[!is.na(PI$BMI),]

ggplot(PI_BMI, aes(x=BMI,fill=Outcome)) + theme_minimal() +
  geom_density(color=NA) + facet_wrap(vars(Outcome), scale="free_x") +
  geom_vline(aes(xintercept = median(BMI)), size = .8, color = "black") +
  geom_vline(aes(xintercept = mean(BMI)), size=.5, linetype="dashed") +
  scale_fill_discrete(name = "Diabetes", labels = c("Absent", "Present"))</pre>
```



```
PI_Insulin <- PI[!is.na(PI$Insulin),]

ggplot(PI_Insulin, aes(x=Insulin,fill=Outcome)) + theme_minimal() +
  geom_density(color=NA) + facet_wrap(vars(Outcome), scale="free_x") +
  geom_vline(aes(xintercept = median(Insulin)), size = .8, color = "black") +
  geom_vline(aes(xintercept = mean(Insulin)), size=.5, linetype="dashed") +
  scale_fill_discrete(name = "Diabetes", labels = c("Absent", "Present"))</pre>
```



Creating two vectors in which I eliminate NA values for each field, independently. This way, I avoid eliminating data from the neighboring columns and portray an accurate distribution.

## Exploring same statistics for all samples that had every attribute present.

This new dataframe only has 392 rows rather than 768. This is around half of the dataframe at ~ 51%.

- · Noticing that every field's average dropped slightly.
  - · BMI still large.
  - Average Glucose levels are normal.
  - Average Blood Pressure is normal but I still see a high max at p100. This is expected, however, as the number of those diagnosed with diabetes now make up ~33% of the total samples.
  - Average Insulin levels for those with all attributes recorded are higher than what is considered normal by 16 mg/dL.

For those with all attributes recorded, using the quartile ranges, I can determine that at least 75% of the women in the samples will: + Have had at least one pregnancy. + Be 23 years of age or older. + Have glucose levels of 99 or higher. (The average is higher than what's considered normal by 16 miligrams / decilitre). + Have a BMI of 28.4 or higher. + Have blood pressure of 62 or more (Nothing out of the ordinary).

skimr::skim(PI %>% drop\_na())

#### Data summary

Name	PI %>% drop_na()
Number of rows	392
Number of columns	11
Column type frequency:	
factor	3
numeric	8
Group variables	None

### Variable type: factor

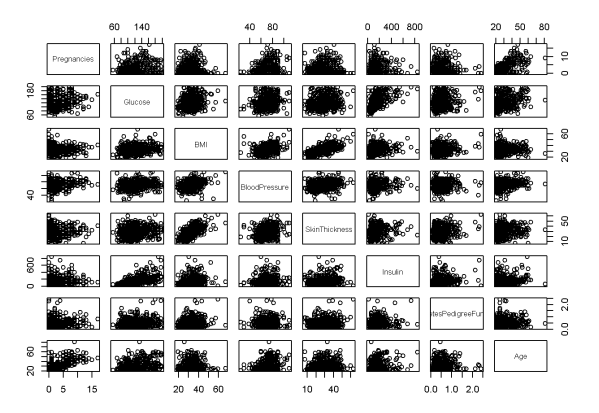
skim_variable	n_missing	complete_rate	ordered	n_unique	top_counts
GlucoseGroup	0	1	FALSE	3	Dia: 168, Pre: 122, Nor: 102
WeightGroup	0	1	FALSE	4	Obe: 262, Ove: 85, Nor: 44, Und: 1
Outcome	0	1	FALSE	2	0: 262, 1: 130

### Variable type: numeric

skim_variable	n_missing	complete_rate	mean	sd	p0	p25	p50	p75	p100	hist
Pregnancies	0	1	3.30	3.21	0.00	1.00	2.00	5.00	17.00	
Glucose	0	1	122.63	30.86	56.00	99.00	119.00	143.00	198.00	_===
BMI	0	1	33.09	7.03	18.20	28.40	33.20	37.10	67.10	
BloodPressure	0	1	70.66	12.50	24.00	62.00	70.00	78.00	110.00	
SkinThickness	0	1	29.15	10.52	7.00	21.00	29.00	37.00	63.00	
Insulin	0	1	156.06	118.84	14.00	76.75	125.50	190.00	846.00	<b>I</b>
DiabetesPedigreeFunction	0	1	0.52	0.35	0.09	0.27	0.45	0.69	2.42	
Age	0	1	30.86	10.20	21.00	23.00	27.00	36.00	81.00	

Checking correlations between variables to avoid multi-collinearity for future predictions. Noticing that the variables that have the highest correlation are: 1) Pregnancy and Age 2) Skin Thickness and BMI 3) Insulin and Glucose

```
numeric_columns <- select_if(PI, is.numeric) %>% tidyr::drop_na()
plot(numeric_columns)
```



### Comparing the mean values for women who have diabetes against those who did not:

```
num_0 <- PI %>% filter(Outcome==0) %>% drop_na() %>% select(Pregnancies, Glucose, BMI, BloodPressure, SkinThic
kness, Insulin, DiabetesPedigreeFunction, Age) %>%
    colMeans() %>% as.data.frame() %>% tibble::rownames_to_column() %>%
    rename("0" = ".")

num_1 <- PI %>% filter(Outcome==1) %>% drop_na() %>% select(Pregnancies, Glucose, BMI, BloodPressure, SkinThic
kness, Insulin, DiabetesPedigreeFunction, Age) %>%
    colMeans() %>% as.data.frame() %>% tibble::rownames_to_column() %>%
    rename("1" = ".")

stat_means <- inner_join(num_0,num_1, by="rowname") %>% rename("Variable" = "rowname")
stat_means
```

```
Variable
##
                                                   1
## 1
                 Pregnancies
                                           4.4692308
                               2.7213740
## 2
                     Glucose 111.4312977 145.1923077
## 3
                          BMI 31.7507634 35.7776923
               BloodPressure 68.9694656
                                          74.0769231
## 4
## 5
               SkinThickness 27.2519084 32.9615385
## 6
                     Insulin 130.8549618 206.8461538
## 7 DiabetesPedigreeFunction
                               0.4721679
                                           0.6255846
## 8
                         Age 28.3473282 35.9384615
```