A Preprocessing analysis of clinical data of TCGA-KIRC patients

This project contains a pipeline of clinical analysis of the Cancer Genome Atlas Kidney Renal Clear Cell Carcinoma (TCGA-KIRC) data of patients from Genomic Data Commons Data Portal and cBioPortal.

In this section, we present a preprocessing analysis of clinical data.

1. Importing data

```
kirc_clin_raw <- read_delim("data/kirc_tcga_clinical_data.tsv", "\t",</pre>
                            escape_double = FALSE,
                            trim_ws = TRUE)
class(kirc_clin_raw)
## [1] "spec_tbl_df" "tbl_df"
                                   "tbl"
                                                  "data.frame"
dim(kirc_clin_raw)
## [1] 538 89
names(kirc_clin_raw)
##
    [1] "Study ID"
   [2] "Patient ID"
   [3] "Sample ID"
##
##
   [4] "Diagnosis Age"
   [5] "Neoplasm American Joint Committee on Cancer Clinical Distant Metastasis M Stage"
##
   [6] "American Joint Committee on Cancer Metastasis Stage Code"
   [7] "Neoplasm Disease Lymph Node Stage American Joint Committee on Cancer Code"
##
##
   [8] "Neoplasm Disease Stage American Joint Committee on Cancer Code"
  [9] "American Joint Committee on Cancer Publication Version Type"
## [10] "American Joint Committee on Cancer Tumor Stage Code"
## [11] "Cancer Type"
## [12] "Cancer Type Detailed"
## [13] "Neoplasm American Joint Committee on Cancer Clinical Group Stage"
## [14] "Neoplasm American Joint Committee on Cancer Clinical Regional Lymph Node N Stage"
## [15] "Neoplasm American Joint Committee on Cancer Clinical Primary Tumor T Stage"
## [16] "Days to Sample Collection."
## [17] "Last Alive Less Initial Pathologic Diagnosis Date Calculated Day Value"
## [18] "Days to Sample Procurement"
## [19] "Disease Free (Months)"
## [20] "Disease Free Status"
## [21] "Disease code"
## [22] "Performance Status"
## [23] "Erythrocyte sedimentation rate"
## [24] "Ethnicity Category"
## [25] "Lymphomatous Extranodal Site Involvement Indicator"
## [26] "Form completion date"
```

```
## [27] "Fraction Genome Altered"
## [28] "Neoplasm Histologic Grade"
## [29] "Hemoglobin level"
## [30] "Neoplasm Histologic Type Name"
## [31] "Neoadjuvant Therapy Type Administered Prior To Resection Text"
## [32] "Prior Cancer Diagnosis Occurence"
## [33] "ICD-10 Classification"
## [34] "International Classification of Diseases for Oncology, Third Edition ICD-0-3 Histology Code"
## [35] "International Classification of Diseases for Oncology, Third Edition ICD-0-3 Site Code"
## [36] "Idh level"
## [37] "Informed consent verified"
## [38] "Year Cancer Initial Diagnosis"
## [39] "Is FFPE"
## [40] "Karnofsky Performance Score"
## [41] "Primary Tumor Laterality"
## [42] "Longest Dimension"
## [43] "Primary Lymph Node Presentation Assessment Ind-3"
## [44] "Lymph nodes examined positive"
## [45] "Lymph Node(s) Examined Number"
## [46] "First Pathologic Diagnosis Biospecimen Acquisition Method Type"
## [47] "Mutation Count"
## [48] "New Neoplasm Event Post Initial Therapy Indicator"
## [49] "Oct embedded"
## [50] "Oncotree Code"
## [51] "Overall Survival (Months)"
## [52] "Overall Survival Status"
## [53] "Specimen Collection Method"
## [54] "Other Patient ID"
## [55] "Other Sample ID"
## [56] "Pathology Report File Name"
## [57] "Pathology report uuid"
## [58] "Performance Status Assessment Timepoint Category"
## [59] "Platelet count"
## [60] "Project code"
## [61] "Tissue Prospective Collection Indicator"
## [62] "Race Category"
## [63] "Did patient start adjuvant postoperative radiotherapy?"
## [64] "Tissue Retrospective Collection Indicator"
## [65] "Number of Samples Per Patient"
## [66] "Sample Initial Weight"
## [67] "Sample Type"
## [68] "Sample type id"
## [69] "Serum calcium level"
## [70] "Sex"
## [71] "Shortest Dimension"
## [72] "Tumor Tissue Site"
## [73] "Person Cigarette Smoking History Pack Year Value"
## [74] "Started Smoking Year"
## [75] "Stopped Smoking Year"
## [76] "Specimen Current Weight"
## [77] "Specimen Freezing Means"
```

[80] "Adjuvant Postoperative Targeted Therapy Administered Indicator"

[78] "Specimen Second Longest Dimension"

[79] "Stage Other"

```
## [84] "Patient Smoking History Category"
## [85] "Primary Therapy Outcome Success Type"
## [86] "Person Neoplasm Status"
## [87] "Vial number"
## [88] "Patient's Vital Status"
## [89] "WBC"
glimpse(kirc_clin_raw)
## Rows: 538
## Columns: 89
## $ `Study ID`
## $ `Patient ID`
## $ `Sample ID`
## $ `Diagnosis Age`
## $ `Neoplasm American Joint Committee on Cancer Clinical Distant Metastasis M Stage`
## $ `American Joint Committee on Cancer Metastasis Stage Code`
## $ `Neoplasm Disease Lymph Node Stage American Joint Committee on Cancer Code`
## $ `Neoplasm Disease Stage American Joint Committee on Cancer Code`
## $ `American Joint Committee on Cancer Publication Version Type`
## $ `American Joint Committee on Cancer Tumor Stage Code`
## $ `Cancer Type`
## $ `Cancer Type Detailed`
## $ `Neoplasm American Joint Committee on Cancer Clinical Group Stage`
## $ `Neoplasm American Joint Committee on Cancer Clinical Regional Lymph Node N Stage`
## $ `Neoplasm American Joint Committee on Cancer Clinical Primary Tumor T Stage`
## $ `Days to Sample Collection.`
## $ `Last Alive Less Initial Pathologic Diagnosis Date Calculated Day Value`
## $ `Days to Sample Procurement`
## $ `Disease Free (Months)`
## $ `Disease Free Status`
## $ `Disease code`
## $ `Performance Status`
## $ `Erythrocyte sedimentation rate`
## $ `Ethnicity Category`
## $ `Lymphomatous Extranodal Site Involvement Indicator`
## $ `Form completion date`
## $ `Fraction Genome Altered`
## $ `Neoplasm Histologic Grade`
## $ `Hemoglobin level`
## $ `Neoplasm Histologic Type Name`
## $ `Neoadjuvant Therapy Type Administered Prior To Resection Text`
## $ `Prior Cancer Diagnosis Occurence`
## $ `ICD-10 Classification`
## $ `International Classification of Diseases for Oncology, Third Edition ICD-0-3 Histology Code`
## $ `International Classification of Diseases for Oncology, Third Edition ICD-0-3 Site Code`
## $ `Idh level`
## $ `Informed consent verified`
## $ `Year Cancer Initial Diagnosis`
## $ `Is FFPE`
## $ `Karnofsky Performance Score`
## $ `Primary Tumor Laterality`
```

<chr

<chr

<chr

<dbl

<chr

<chr

<chr

<chr

<chr

<chr

<chr

<chr

<lg1

<lg1

<lg1

<dbl

<dbl

<lg1

<dbl

<chr

<chr

<dbl

<chr

<chr

<lg1

<chr

<dbl

<chr

<dbl

<chr

<dbl

<chr

[81] "Time between clamping and freezing"
[82] "Time between excision and freezing"

[83] "Tissue Source Site"

```
## $ `Longest Dimension`
## $ `Primary Lymph Node Presentation Assessment Ind-3`
## $ `Lymph nodes examined positive`
## $ `Lymph Node(s) Examined Number`
## $ `First Pathologic Diagnosis Biospecimen Acquisition Method Type`
## $ `Mutation Count`
## $ `New Neoplasm Event Post Initial Therapy Indicator`
## $ `Oct embedded`
## $ `Oncotree Code`
## $ `Overall Survival (Months)`
## $ `Overall Survival Status`
## $ `Specimen Collection Method`
## $ `Other Patient ID`
## $ `Other Sample ID`
## $ `Pathology Report File Name`
## $ `Pathology report uuid`
## $ `Performance Status Assessment Timepoint Category`
## $ `Platelet count`
## $ `Project code`
## $ `Tissue Prospective Collection Indicator`
## $ `Race Category`
## $ `Did patient start adjuvant postoperative radiotherapy?`
## $ `Tissue Retrospective Collection Indicator`
## $ `Number of Samples Per Patient`
## $ `Sample Initial Weight`
## $ `Sample Type`
## $ `Sample type id`
## $ `Serum calcium level`
## $ Sex
## $ `Shortest Dimension`
## $ `Tumor Tissue Site`
## $ `Person Cigarette Smoking History Pack Year Value`
## $ `Started Smoking Year`
## $ `Stopped Smoking Year`
## $ `Specimen Current Weight`
## $ `Specimen Freezing Means`
## $ `Specimen Second Longest Dimension`
## $ `Stage Other`
## $ `Adjuvant Postoperative Targeted Therapy Administered Indicator`
## $ `Time between clamping and freezing`
## $ `Time between excision and freezing`
## $ `Tissue Source Site`
## $ `Patient Smoking History Category`
## $ `Primary Therapy Outcome Success Type`
## $ 'Person Neoplasm Status'
## $ `Vial number`
## $ 'Patient's Vital Status'
## $ WBC
```

<dbl

<chr

<dbl

<dbl

<lg1

<dbl

<chr

<lg1

<chr

<dbl

<chr

<lg1

<chr

<dbl

<dbl

<chr

<dbl

<chr

<chr

<dbl

<chr

<dbl

<dbl

<dbl

<lg1

<lg1

<dbl

<lg1

<chr

<lg1

<lg1

<chr

<dbl

<chr

<chr

<chr

<chr

<chr

Table 1: Data summary

skim(kirc_clin_raw)

Name	kirc_clin_raw
Number of rows	538

Table 1: Data summary

Number of columns	89
Column type frequency:	
character	54
logical	13
numeric	22
Group variables	None

Variable type: character

skim_variable	n_missing	complete_rate
Study ID	0	1.00
Patient ID	0	1.00
Sample ID	0	1.00
Neoplasm American Joint Committee on Cancer Clinical Distant Metastasis M Stage	504	0.06
American Joint Committee on Cancer Metastasis Stage Code	2	1.00
Neoplasm Disease Lymph Node Stage American Joint Committee on Cancer Code	0	1.00
Neoplasm Disease Stage American Joint Committee on Cancer Code	3	0.99
American Joint Committee on Cancer Publication Version Type	367	0.32
American Joint Committee on Cancer Tumor Stage Code	0	1.00
Cancer Type	0	1.00
Cancer Type Detailed	0	1.00
Disease Free Status	99	0.82
Disease code	534	0.01
Erythrocyte sedimentation rate	527	0.02
Ethnicity Category	152	0.72
Form completion date	0	1.00
Neoplasm Histologic Grade	3	0.99
Hemoglobin level	83	0.85
Neoplasm Histologic Type Name	0	1.00
Neoadjuvant Therapy Type Administered Prior To Resection Text	0	1.00
Prior Cancer Diagnosis Occurence	0	1.00
ICD-10 Classification	0	1.00
International Classification of Diseases for Oncology, Third Edition ICD-O-3 Histology Code	0	1.00
International Classification of Diseases for Oncology, Third Edition ICD-O-3 Site Code	0	1.00
Idh level	451	0.16
Informed consent verified	0	1.00
Is FFPE	1	1.00
Primary Tumor Laterality	0	1.00
Primary Lymph Node Presentation Assessment Ind-3	7	0.99
New Neoplasm Event Post Initial Therapy Indicator Oncotree Code	503	0.07
Overall Survival Status	0	1.00 1.00
Other Patient ID	_	1.00
	0	
Other Sample ID Pathology Report File Name	$1 \\ 1$	1.00 1.00
Pathology report uuid	1	1.00
Performance Status Assessment Timepoint Category	428	0.20
Platelet count	93	0.20
Project code	534	0.01
1 Tojoco codo	554	0.01

skim_variable	$n_{missing}$	$complete_rate$
Tissue Prospective Collection Indicator	20	0.96
Race Category	7	0.99
Did patient start adjuvant postoperative radiotherapy?	506	0.06
Tissue Retrospective Collection Indicator	18	0.97
Sample Type	0	1.00
Serum calcium level	172	0.68
Sex	0	1.00
Tumor Tissue Site	0	1.00
Adjuvant Postoperative Targeted Therapy Administered Indicator	506	0.06
Tissue Source Site	0	1.00
Primary Therapy Outcome Success Type	507	0.06
Person Neoplasm Status	35	0.93
Vial number	1	1.00
Patient's Vital Status	3	0.99
WBC	96	0.82

Variable type: logical

skim_variable	n_missing	complete_rate	m
Neoplasm American Joint Committee on Cancer Clinical Group Stage	538	0.00	N
Neoplasm American Joint Committee on Cancer Clinical Regional Lymph Node N Stage	538	0.00	Ν
Neoplasm American Joint Committee on Cancer Clinical Primary Tumor T Stage	538	0.00	Ν
Days to Sample Procurement	538	0.00	Ν
Lymphomatous Extranodal Site Involvement Indicator	538	0.00	Ν
First Pathologic Diagnosis Biospecimen Acquisition Method Type	538	0.00	N
Oct embedded	503	0.07	(
Specimen Collection Method	538	0.00	N
Specimen Current Weight	538	0.00	N
Specimen Freezing Means	538	0.00	N
Stage Other	538	0.00	N
Time between clamping and freezing	538	0.00	N
Time between excision and freezing	538	0.00	N

Variable type: numeric

skim_variable	n_missing	complete_rate	mean	sd
Diagnosis Age	0	1.00	60.58	12.14
Days to Sample Collection.	503	0.07	545.26	566.08
Last Alive Less Initial Pathologic Diagnosis Date Calculated Day Value	0	1.00	0.00	0.00
Disease Free (Months)	99	0.82	40.30	31.64
Performance Status	399	0.26	0.24	0.48
Fraction Genome Altered	9	0.98	0.17	0.17
Year Cancer Initial Diagnosis	0	1.00	2006.02	2.76
Karnofsky Performance Score	490	0.09	88.33	20.56
Longest Dimension	36	0.93	1.66	0.66
Lymph nodes examined positive	407	0.24	0.34	1.44
Lymph Node(s) Examined Number	404	0.25	6.30	5.97
Mutation Count	87	0.84	73.85	127.76
Overall Survival (Months)	0	1.00	44.33	32.26
Number of Samples Per Patient	0	1.00	1.00	0.06

skim_variable	n_missing	$complete_rate$	mean	sd
Sample Initial Weight	503	0.07	296.29	366.21
Sample type id	0	1.00	1.01	0.17
Shortest Dimension	36	0.93	0.38	0.21
Person Cigarette Smoking History Pack Year Value	516	0.04	28.55	15.77
Started Smoking Year	525	0.02	1978.38	17.35
Stopped Smoking Year	525	0.02	1994.77	15.12
Specimen Second Longest Dimension	36	0.93	0.94	0.31
Patient Smoking History Category	450	0.16	1.91	1.19

```
#View(kirc_clin_raw)
```

2. Cleaning data

Select variables based on NA count (> 50% complete is a good choice!).

```
NA_fifty <- dim(kirc_clin_raw)[1]/2

NA_sum <- colSums(is.na(kirc_clin_raw))
NA_sum <- as.data.frame(NA_sum)
NA_sum <- tibble::rownames_to_column(NA_sum, "variables")
NA_sum <- NA_sum %>%
    filter(NA_sum < NA_fifty)

kirc_clean <- kirc_clin_raw %>%
    select(one_of(NA_sum$variables))
```

Remove duplicate observations:

```
kirc_clean0 <- kirc_clean %>%
    distinct_at('Patient ID', .keep_all = TRUE)
```

Remove nuneric variables with unique observations:

```
skim(kirc_clean0)
```

Table 5: Data summary

Name Number of rows	kirc_clean0 537
Number of columns	55
Column type frequency:	
character	43
numeric	12
Group variables	None

Variable type: character

skim_variable	$n_{missing}$	$complete_rate$
Study ID	0	1.00

skim_variable	n_missing	complete_rate
Patient ID	0	1.00
Sample ID	0	1.00
American Joint Committee on Cancer Metastasis Stage Code	2	1.00
Neoplasm Disease Lymph Node Stage American Joint Committee on Cancer Code	0	1.00
Neoplasm Disease Stage American Joint Committee on Cancer Code	3	0.99
American Joint Committee on Cancer Tumor Stage Code	0	1.00
Cancer Type	0	1.00
Cancer Type Detailed	0	1.00
Disease Free Status	99	0.82
Ethnicity Category	152	0.72
Form completion date	0	1.00
Neoplasm Histologic Grade	3	0.99
Hemoglobin level	83	0.85
Neoplasm Histologic Type Name	0	1.00
Neoadjuvant Therapy Type Administered Prior To Resection Text	0	1.00
Prior Cancer Diagnosis Occurence	0	1.00
ICD-10 Classification	0	1.00
International Classification of Diseases for Oncology, Third Edition ICD-O-3 Histology Code	0	1.00
International Classification of Diseases for Oncology, Third Edition ICD-O-3 Site Code	0	1.00
Informed consent verified	0	1.00
Is FFPE	0	1.00
Primary Tumor Laterality	0	1.00
Primary Lymph Node Presentation Assessment Ind-3	7	0.99
Oncotree Code	0	1.00
Overall Survival Status	0	1.00
Other Patient ID	0	1.00
Other Sample ID	0	1.00
Pathology Report File Name	0	1.00
Pathology report uuid	0	1.00
Platelet count	93	0.83
Tissue Prospective Collection Indicator	20	0.96
Race Category	7	0.99
Tissue Retrospective Collection Indicator	18	0.97
Sample Type	0	1.00
Serum calcium level	172	0.68
Sex	0	1.00
Tumor Tissue Site	0	1.00
Tissue Source Site	0	1.00
Person Neoplasm Status	35	0.93
Vial number	0	1.00
Patient's Vital Status	3	0.99
WBC	96	0.82

Variable type: numeric

skim_variable	n_missing	$complete_rate$	mean	sd
Diagnosis Age	0	1.00	60.59	12.15
Last Alive Less Initial Pathologic Diagnosis Date Calculated Day Value	0	1.00	0.00	0.00
Disease Free (Months)	99	0.82	40.24	31.66
Fraction Genome Altered	9	0.98	0.17	0.17
Year Cancer Initial Diagnosis	0	1.00	2006.02	2.76

skim_variable	n_missing	$complete_rate$	mean	sd
Longest Dimension	35	0.93	1.66	0.66
Mutation Count	86	0.84	73.85	127.76
Overall Survival (Months)	0	1.00	44.26	32.25
Number of Samples Per Patient	0	1.00	1.00	0.04
Sample type id	0	1.00	1.00	0.00
Shortest Dimension	35	0.93	0.38	0.21
Specimen Second Longest Dimension	35	0.93	0.94	0.31

Table 8: Data summary

Name	kirc_clean1
Number of rows	537
Number of columns	52
Column type frequency: character	43
numeric	9
Group variables	None

Variable type: character

$n_{missing}$	$complete_rate$
0	1.00
0	1.00
0	1.00
2	1.00
0	1.00
3	0.99
0	1.00
0	1.00
0	1.00
99	0.82
152	0.72
0	1.00
3	0.99
83	0.85
0	1.00
0	1.00
0	1.00
0	1.00
0	1.00
0	1.00
	0 0 0 2 0 3 0 0 0 99 152 0 3 83 0 0 0

skim_variable	n_missing	complete_rate
Informed consent verified	0	1.00
Is FFPE	0	1.00
Primary Tumor Laterality	0	1.00
Primary Lymph Node Presentation Assessment Ind-3	7	0.99
Oncotree Code	0	1.00
Overall Survival Status	0	1.00
Other Patient ID	0	1.00
Other Sample ID	0	1.00
Pathology Report File Name	0	1.00
Pathology report uuid	0	1.00
Platelet count	93	0.83
Tissue Prospective Collection Indicator	20	0.96
Race Category	7	0.99
Tissue Retrospective Collection Indicator	18	0.97
Sample Type	0	1.00
Serum calcium level	172	0.68
Sex	0	1.00
Tumor Tissue Site	0	1.00
Tissue Source Site	0	1.00
Person Neoplasm Status	35	0.93
Vial number	0	1.00
Patient's Vital Status	3	0.99
WBC	96	0.82

Variable type: numeric

skim_variable	n_missing	$complete_rate$	mean	sd	p0	p25	p50	
Diagnosis Age	0	1.00	60.59	12.15	26.00	52.00	61.00	7
Disease Free (Months)	99	0.82	40.24	31.66	-11.79	13.43	36.20	6
Fraction Genome Altered	9	0.98	0.17	0.17	0.00	0.06	0.12	
Year Cancer Initial Diagnosis	0	1.00	2006.02	2.76	1998.00	2004.00	2006.00	200'
Longest Dimension	35	0.93	1.66	0.66	0.40	1.20	1.50	}
Mutation Count	86	0.84	73.85	127.76	1.00	34.00	48.00	6
Overall Survival (Months)	0	1.00	44.26	32.25	0.00	18.10	38.96	6
Shortest Dimension	35	0.93	0.38	0.21	0.10	0.20	0.30	1
Specimen Second Longest Dimension	35	0.93	0.94	0.31	0.30	0.70	0.90	

Remove character variables with unique observations:

Table 11: Data summary

Name	kirc_clean2
Number of rows	537

Table 11: Data summary

Number of columns	41
Column type frequency:	
character	32
numeric	9
Group variables	None

Variable type: character

skim_variable	n_missing	complete_rate
Patient ID	0	1.00
Sample ID	0	1.00
American Joint Committee on Cancer Metastasis Stage Code	2	1.00
Neoplasm Disease Lymph Node Stage American Joint Committee on Cancer Code	0	1.00
Neoplasm Disease Stage American Joint Committee on Cancer Code	3	0.99
American Joint Committee on Cancer Tumor Stage Code	0	1.00
Disease Free Status	99	0.82
Ethnicity Category	152	0.72
Form completion date	0	1.00
Neoplasm Histologic Grade	3	0.99
Hemoglobin level	83	0.85
Neoadjuvant Therapy Type Administered Prior To Resection Text	0	1.00
Prior Cancer Diagnosis Occurence	0	1.00
International Classification of Diseases for Oncology, Third Edition ICD-O-3 Histology Code	0	1.00
Primary Tumor Laterality	0	1.00
Primary Lymph Node Presentation Assessment Ind-3	7	0.99
Overall Survival Status	0	1.00
Other Patient ID	0	1.00
Other Sample ID	0	1.00
Pathology Report File Name	0	1.00
Pathology report uuid	0	1.00
Platelet count	93	0.83
Tissue Prospective Collection Indicator	20	0.96
Race Category	7	0.99
Tissue Retrospective Collection Indicator	18	0.97
Serum calcium level	172	0.68
Sex	0	1.00
Tissue Source Site	0	1.00
Person Neoplasm Status	35	0.93
Vial number	0	1.00
Patient's Vital Status	3	0.99
WBC	96	0.82

Variable type: numeric

skim_variable	n_missing	complete_rate	mean	sd	p0	p25	p50	
Diagnosis Age	0	1.00	60.59	12.15	26.00	52.00	61.00	7
Disease Free (Months)	99	0.82	40.24	31.66	-11.79	13.43	36.20	60

skim_variable	$n_missing$	$complete_rate$	mean	sd	p0	p25	p50	
Fraction Genome Altered	9	0.98	0.17	0.17	0.00	0.06	0.12	
Year Cancer Initial Diagnosis	0	1.00	2006.02	2.76	1998.00	2004.00	2006.00	200
Longest Dimension	35	0.93	1.66	0.66	0.40	1.20	1.50	
Mutation Count	86	0.84	73.85	127.76	1.00	34.00	48.00	6
Overall Survival (Months)	0	1.00	44.26	32.25	0.00	18.10	38.96	6
Shortest Dimension	35	0.93	0.38	0.21	0.10	0.20	0.30	
Specimen Second Longest Dimension	35	0.93	0.94	0.31	0.30	0.70	0.90	

Remove character variables with similar information - check each one!

```
table(kirc clean2$`Overall Survival Status`, exclude = NULL)
##
## DECEASED
             LIVING
##
        177
                 360
table(kirc_clean2$`Patient's Vital Status`, exclude = NULL)
## Alive Dead <NA>
     360
          174
kirc_clean3 <- kirc_clean2 %>%
     select(!c('Sample ID', 'Other Patient ID', 'Other Sample ID', 'Pathology Report File Name', 'Patho
# removing other variables not directly related to patient - check each one!
kirc clean4 <- kirc clean3 %>%
     select(!c('Form completion date','International Classification of Diseases for Oncology, Third Edi
```

3. Changing variables names

Using snake style

```
kirc clean4 <- kirc clean4 %>%
     rename(patient_id = 'Patient ID',
            age = 'Diagnosis Age',
            metastasis_stg = 'American Joint Committee on Cancer Metastasis Stage Code',
            neoplasm_ln_stg = 'Neoplasm Disease Lymph Node Stage American Joint Committee on Cancer Cod
            neoplasm_stg = 'Neoplasm Disease Stage American Joint Committee on Cancer Code',
            tumor_stg = 'American Joint Committee on Cancer Tumor Stage Code',
            disease_free_mth = 'Disease Free (Months)',
            disease_free_stt = 'Disease Free Status',
            ethnicity = 'Ethnicity Category',
            frac_genome_alter = 'Fraction Genome Altered',
            histology_grd = 'Neoplasm Histologic Grade',
            hemoglobin = 'Hemoglobin level',
            neoadj_therapy = 'Neoadjuvant Therapy Type Administered Prior To Resection Text',
            prior_cancer = 'Prior Cancer Diagnosis Occurence',
            year_diagnose = 'Year Cancer Initial Diagnosis',
            tumor_lateral = 'Primary Tumor Laterality',
            long_dim = 'Longest Dimension',
            primer_ln_ind3 = 'Primary Lymph Node Presentation Assessment Ind-3',
            mutation_cnt = 'Mutation Count',
```

```
over_surv_mth = 'Overall Survival (Months)',
over_surv_stt = 'Overall Survival Status',
platelet = 'Platelet count',
tissue_prospect = 'Tissue Prospective Collection Indicator',
race = 'Race Category',
tissue_retrospect = 'Tissue Retrospective Collection Indicator',
serum_ca = 'Serum calcium level',
sex = 'Sex',
short_dim = 'Shortest Dimension',
second_long_dim = 'Specimen Second Longest Dimension',
tissue_site = 'Tissue Source Site',
person_neoplasm_stt = 'Person Neoplasm Status',
wbc = 'WBC')
```

4. Taming data

Use lubridate for dates

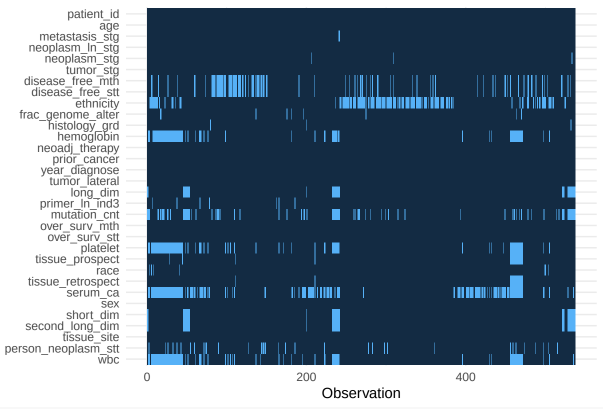
```
kirc_clean4 <- kirc_clean4 %>%
    mutate_if(is.character, as.factor) %>%
    mutate(patient_id = as.character(patient_id))
```

5. Checking NA patterns

Check distincts types of NAs: MCAR, MAR, MNAR

```
kirc_clean4 %>%
  missing_plot()
```

Missing values map



missing_glimpse(kirc_clean4)

##		label	var_type	n	missing_n	missing_percent
##	patient_id	patient_id	<chr></chr>		0	0.0
##	age	age	<dbl></dbl>	537	0	0.0
##	metastasis_stg	metastasis_stg	<fct></fct>	535	2	0.4
##	neoplasm_ln_stg	neoplasm_ln_stg	<fct></fct>	537	0	0.0
##	neoplasm_stg	neoplasm_stg	<fct></fct>	534	3	0.6
##	tumor_stg	tumor_stg	<fct></fct>	537	0	0.0
##	disease_free_mth	disease_free_mth	<dbl></dbl>	438	99	18.4
##	disease_free_stt	disease_free_stt	<fct></fct>	438	99	18.4
##	ethnicity	ethnicity	<fct></fct>	385	152	28.3
##	<pre>frac_genome_alter</pre>	<pre>frac_genome_alter</pre>	<dbl></dbl>	528	9	1.7
##	histology_grd	histology_grd	<fct></fct>	534	3	0.6
##	hemoglobin	hemoglobin	<fct></fct>	454	83	15.5
##	neoadj_therapy	neoadj_therapy	<fct></fct>	537	0	0.0
##	<pre>prior_cancer</pre>	<pre>prior_cancer</pre>	<fct></fct>	537	0	0.0
##	year_diagnose	year_diagnose	<dbl></dbl>	537	0	0.0
##	tumor_lateral	tumor_lateral	<fct></fct>	537	0	0.0
##	long_dim	long_dim	<dbl></dbl>	502	35	6.5
##	primer_ln_ind3	<pre>primer_ln_ind3</pre>	<fct></fct>	530	7	1.3
##	mutation_cnt	mutation_cnt	<dbl></dbl>	451	86	16.0
##	over_surv_mth	over_surv_mth	<dbl></dbl>	537	0	0.0
##	over_surv_stt	over_surv_stt	<fct></fct>	537	0	0.0
##	platelet	platelet	<fct></fct>	444	93	17.3
##	tissue_prospect	tissue_prospect	<fct></fct>	517	20	3.7
##	race	race	<fct></fct>	530	7	1.3

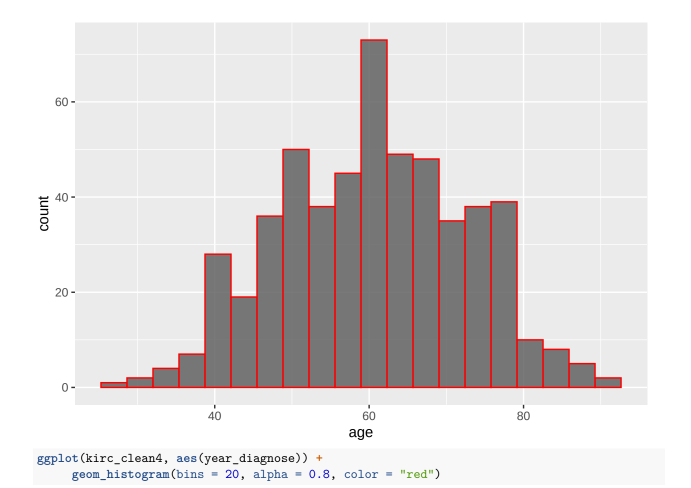
```
## tissue retrospect
                          tissue_retrospect
                                                <fct> 519
                                                                  18
                                                                                  3.4
                                                <fct> 365
                                                                 172
                                                                                 32.0
## serum_ca
                                   serum_ca
## sex
                                         sex
                                                <fct> 537
                                                                   0
                                                                                  0.0
## short_dim
                                                <dbl> 502
                                                                  35
                                                                                  6.5
                                  short_dim
## second_long_dim
                            second_long_dim
                                                <dbl> 502
                                                                  35
                                                                                  6.5
## tissue site
                                tissue site
                                                <fct> 537
                                                                   0
                                                                                  0.0
## person_neoplasm_stt person_neoplasm_stt
                                                <fct> 502
                                                                  35
                                                                                  6.5
## wbc
                                                <fct> 441
                                                                                 17.9
                                                                  96
```

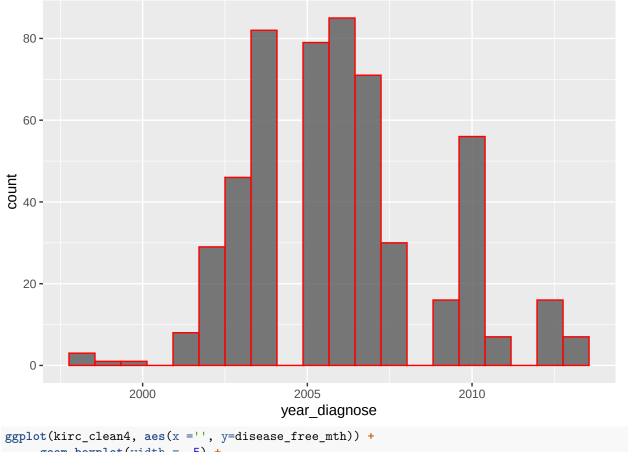
6. Checking numeric variables

Check data distribution, plausible ranges, outliers; Thinking about deleting outliers from dataset? Need to evaluate carefully each one!

```
kirc_clean4 %>%
    select_if(is.numeric) %>%
    summary()
```

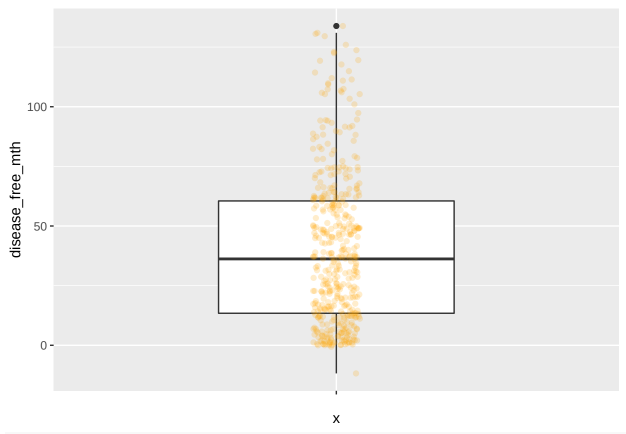
```
##
         age
                     disease_free_mth frac_genome_alter year_diagnose
##
                            :-11.79
           :26.00
                     Min.
                                       Min.
                                              :0.00000
                                                         Min.
                                                                 :1998
   Min.
    1st Qu.:52.00
                     1st Qu.: 13.43
                                       1st Qu.:0.06295
                                                          1st Qu.:2004
   Median :61.00
                     Median : 36.20
                                      Median :0.12065
                                                         Median:2006
##
                           : 40.24
   Mean
           :60.59
                                              :0.17016
                                                         Mean
##
                     Mean
                                       Mean
                                                                 :2006
##
    3rd Qu.:70.00
                     3rd Qu.: 60.51
                                       3rd Qu.:0.20885
                                                          3rd Qu.:2007
##
   Max.
           :90.00
                     Max.
                            :133.84
                                      Max.
                                              :0.94770
                                                         Max.
                                                                 :2013
##
                     NA's
                            :99
                                       NA's
                                              :9
##
       long_dim
                                       over_surv_mth
                                                            short_dim
                     mutation_cnt
##
   Min.
           :0.400
                     Min.
                                1.00
                                       Min.
                                               : 0.00
                                                         Min.
                                                                 :0.1000
##
    1st Qu.:1.200
                     1st Qu.:
                               34.00
                                       1st Qu.: 18.10
                                                          1st Qu.:0.2000
##
   Median :1.500
                     Median :
                               48.00
                                       Median : 38.96
                                                         Median :0.3000
##
   Mean
           :1.662
                            : 73.85
                                               : 44.26
                     Mean
                                       Mean
                                                         Mean
                                                                 :0.3759
##
    3rd Qu.:2.000
                     3rd Qu.:
                               65.50
                                        3rd Qu.: 63.21
                                                          3rd Qu.:0.5000
           :4.000
                            :1392.00
                                                                 :1.0000
##
  {\tt Max.}
                     Max.
                                       Max.
                                               :149.05
                                                          Max.
##
   NA's
           :35
                     NA's
                            :86
                                                          NA's
                                                                 :35
##
   second_long_dim
   Min.
           :0.3000
   1st Qu.:0.7000
##
##
   Median :0.9000
## Mean
           :0.9368
   3rd Qu.:1.1000
##
  {\tt Max.}
           :2.0000
##
   NA's
           :35
ggplot(kirc_clean4, aes(age)) +
     geom_histogram(bins = 20, alpha = 0.8, color = "red")
```





```
ggplot(kirc_clean4, aes(x = '', y=disease_free_mth)) +
    geom_boxplot(width = .5) +
    geom_jitter(width = 0.05, alpha = 0.2, color = "orange")
```

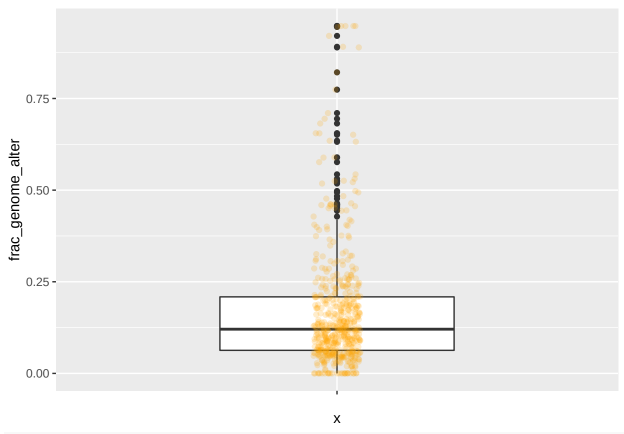
- ## Warning: Removed 99 rows containing non-finite values (stat_boxplot).
- ## Warning: Removed 99 rows containing missing values (geom_point).



boxplot.stats(kirc_clean4\$disease_free_mth)

```
## $stats
## [1] -11.79 13.40 36.20 60.55 130.98
##
## $n
## [1] 438
##
## $conf
## [1] 32.6404 39.7596
##
## $out
## [1] 133.84
# filter(disease_free_mth >= 0)
ggplot(kirc_clean4, aes(x ='', y=frac_genome_alter)) +
     geom_boxplot(width = .5) +
     geom_jitter(width = 0.05, alpha = 0.2, color = "orange")
## Warning: Removed 9 rows containing non-finite values (stat_boxplot).
```

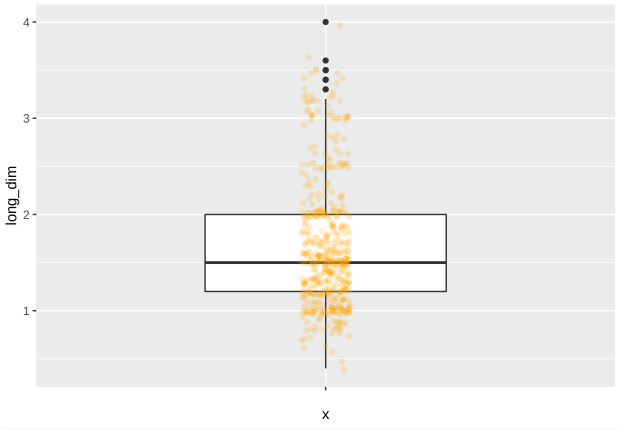
Warning: Removed 9 rows containing missing values (geom_point).



boxplot.stats(kirc_clean4\$frac_genome_alter)

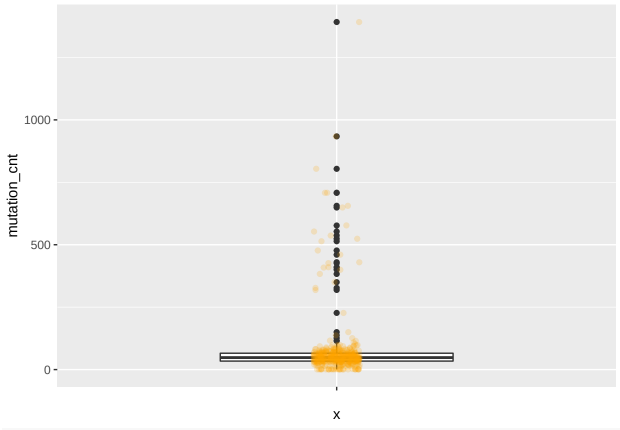
```
## [1] 0.00000 0.06290 0.12065 0.20920 0.42800
##
## $n
## [1] 528
##
## $conf
## [1] 0.1105903 0.1307097
##
## $out
## [1] 0.8213 0.6552 0.4608 0.9477 0.5888 0.9208 0.7741 0.4837 0.9477 0.4610
## [11] 0.6549 0.6511 0.5180 0.8910 0.8893 0.9477 0.5246 0.4568 0.4937 0.9477
## [21] 0.4438 0.6947 0.5218 0.4768 0.4593 0.4447 0.9452 0.6347 0.5311 0.4562
## [31] 0.4617 0.5256 0.6318 0.5430 0.4506 0.5764 0.7102 0.4641 0.5894 0.4976
## [41] 0.4513 0.6818
ggplot(kirc_clean4, aes(x ='', y=long_dim)) +
     geom_boxplot(width = .5) +
     geom_jitter(width = 0.05, alpha = 0.2, color = "orange")
```

Warning: Removed 35 rows containing missing values (geom_point).



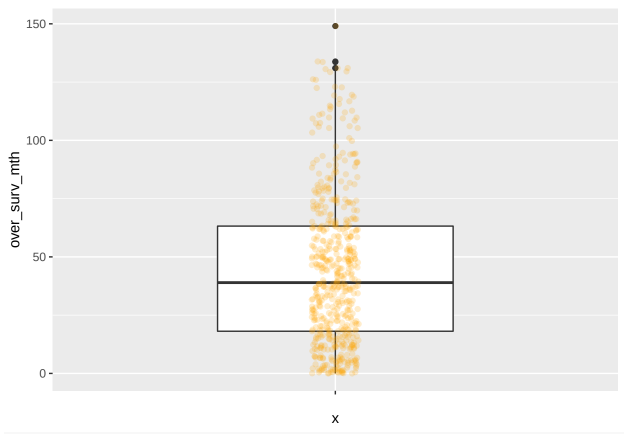
boxplot.stats(kirc_clean4\$long_dim)

 $\hbox{\tt \#\# Warning: Removed 86 rows containing missing values (geom_point).}$



boxplot.stats(kirc_clean4\$mutation_cnt)

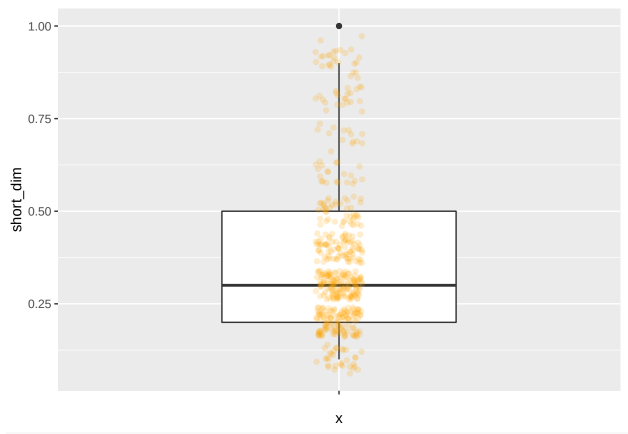
```
## $stats
## [1]
        1.0 34.0 48.0 65.5 109.0
## $n
## [1] 451
##
## $conf
## [1] 45.65642 50.34358
##
## $out
## [1]
       514 656 577 537 477 150 137 708 1392 460 327
                                                                     383
                                                            934 409
## [16] 319 524
                426 227 553 400 350 410 430 708 649
                                                            126
                                                                116
ggplot(kirc_clean4, aes(x ='', y=over_surv_mth)) +
    geom_boxplot(width = .5) +
    geom_jitter(width = 0.05, alpha = 0.2, color = "orange")
```



boxplot.stats(kirc_clean4\$over_surv_mth)

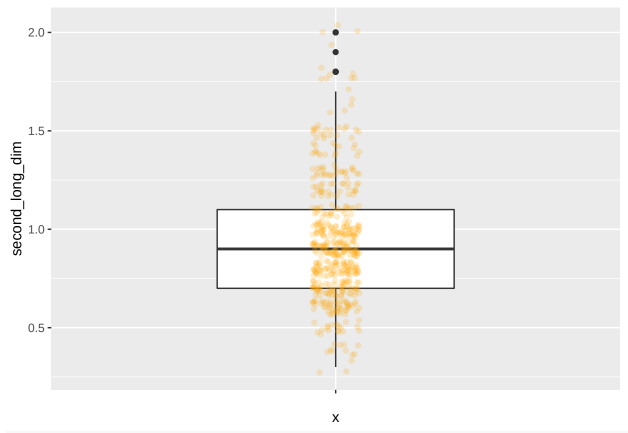
```
## $stats
## [1]
        0.00 18.10 38.96 63.21 130.55
##
## $n
## [1] 537
##
## $conf
## [1] 35.88431 42.03569
##
## $out
## [1] 133.84 149.05 131.04 130.98 133.61
ggplot(kirc_clean4, aes(x ='', y=short_dim)) +
     geom_boxplot(width = .5) +
     geom_jitter(width = 0.05, alpha = 0.2, color = "orange")
## Warning: Removed 35 rows containing non-finite values (stat_boxplot).
```

Warning: Removed 35 rows containing missing values (geom_point).



boxplot.stats(kirc_clean4\$short_dim)

Warning: Removed 35 rows containing missing values (geom_point).



boxplot.stats(kirc_clean4\$second_long_dim)

```
## $stats
## [1] 0.3 0.7 0.9 1.1 1.7
##
## $n
## [1] 502
##
## $conf
## [1] 0.8717925 0.9282075
##
## $out
## [1] 1.8 2.0 1.8 1.9 1.8 2.0 2.0 1.8 1.8 1.8 1.8
```

7. Checking categorical variables

Check frequency, lables and levels

```
kirc_clean4 %>%

select_if(is.factor) %>%

summary()
```

```
## metastasis_stg neoplasm_ln_stg
                                                      tumor_stg
                                       neoplasm_stg
## MO :426
                   NO:240
                                    Stage I :269
                                                    T1a
                                                           :142
  M1 : 79
                   N1: 17
                                    Stage II : 57
                                                           :122
##
                                                    T3a
   MX
       : 30
                   NX:280
                                    Stage III:125
                                                    T<sub>1</sub>b
                                                           :111
## NA's: 2
                                    Stage IV: 83
                                                    T2
                                                           : 55
```

```
NA's : 3 T3b
##
                                                        : 53
##
                                                 Т1
                                                        : 22
                                                 (Other): 32
##
##
                                               ethnicity
              disease_free_stt
                                                           histology_grd
##
   DiseaseFree
                      :311
                               HISPANIC OR LATINO
                                                    : 26
                                                           G1 : 14
   Recurred/Progressed:127
                               NOT HISPANIC OR LATINO:359
                                                           G2 :230
##
   NA's
                  : 99
                               NA's
                                                    :152
                                                           G3 : 207
                                                           G4 : 78
##
##
                                                           GX : 5
##
                                                           NA's: 3
##
##
      hemoglobin neoadj_therapy
   Elevated: 5
                  No:519
##
                  Yes: 18
##
   Low
           :263
##
   Normal:186
##
   NA's
           : 83
##
##
##
##
                                            prior cancer tumor lateral
##
  No
                                                  :459
                                                        Bilateral: 1
##
                                                  : 72
                                                         Left
                                                                :253
  Yes, History of Prior Malignancy
                                                  : 2
                                                         Right
                                                                  :283
##
   Yes, History of Synchronous/Bilateral Malignancy: 4
##
##
##
##
   primer_ln_ind3 over_surv_stt
                                    platelet
                                               tissue_prospect
  NO :395
                  DECEASED: 177
                                 Elevated: 38
                                               NO :465
  YES :135
                  LIVING :360
                                        : 46
                                               YES: 52
                                Low
   NA's: 7
                                               NA's: 20
##
                                 Normal:360
##
                                 NA's
                                        : 93
##
##
##
##
                                  tissue_retrospect
                          race
                                                        serum_ca
                                                                       sex
  ASIAN
                           : 8
                                  NO : 53
                                                    Elevated: 10
                                                                 Female:191
##
  BLACK OR AFRICAN AMERICAN: 56
                                  YES:466
                                                    Low
                                                            :204
                                                                  Male :345
## WHITE
                                                                  MALE : 1
                            :466
                                  NA's: 18
                                                    Normal:151
##
  NA's
                                                    NA's
                            : 7
                                                            :172
##
##
##
##
    tissue_site person_neoplasm_stt
                                          wbc
##
          :142
                 TUMOR FREE:361
                                    Elevated:164
          :107
  B0
                 WITH TUMOR:141
##
                                    Low
                                            : 9
  CJ
          : 71
                 NA's
                      : 35
                                    Normal :268
##
##
  A3
          : 52
                                    NA's
                                            : 96
          : 40
##
  CZ
## B8
          : 33
  (Other): 92
# agregating levels
kirc_clean5 <- kirc_clean4 %>%
```

```
mutate(tumor_stg = fct_collapse(tumor_stg,
                             T1 = c('T1', 'T1a', 'T1b'),
                             T2 = c('T2', 'T2a', 'T2b'),
                             T3 = c('T3', 'T3a', 'T3b', 'T3c')))
kirc_clean5 <- kirc_clean4 %>%
     mutate(prior_cancer = fct_collapse(prior_cancer,
               Yes = c('Yes', 'Yes, History of Prior Malignancy', 'Yes, History of Synchronous/Bilatera
kirc_clean5 <- kirc_clean4 %>%
     mutate(sex = fct_collapse(sex, Male = c('MALE', 'Male')))
kirc clean5 <- kirc clean4 %>%
     mutate(tissue_site = fct_collapse(tissue_site,
                         A = c('A3', 'AK', 'AS'),
                         B = c('B0', 'B2', 'B4', 'B8', 'BP'),
                         C = c('CJ', 'CW', 'CZ'),
                         G = c('G6', 'GK'),
                         M = c('MM', 'MW'))
# droping levels
kirc_clean5 <- kirc_clean4 %>%
     mutate(race = fct_recode(race, NULL = 'ASIAN'))
kirc clean5 <- kirc clean4 %>%
  mutate(tissue_site = fct_recode(tissue_site, NULL = '3Z', NULL='6D', NULL='DV', NULL='EU', NULL='G', )
# kirc_clean5 <- kirc_clean4 %>%
     mutate(race = fct_drop(race, only = 'ASIAN'))
# recoding levels
# OBS: It can be donne latter, for regression analysis
# kirc_clean5 <- kirc_clean4 %>%
       mutate(sex = fct_recode(sex, '1'='Male', '2'='Female'))
#
# kirc_clean5 <- kirc_clean4 %>%
      mutate(sex = if_else(sex %in% c('Male', 'Female'), 1, 0))
# table(kirc_clean5$metastasis_stg, exclude = NULL)
# table(kirc_clean5$neoplasm_ln_stg, exclude = NULL)
# table(kirc_clean5$neoplasm_stg, exclude = NULL)
# table(kirc_clean5$tumor_stg, exclude = NULL)
# table(kirc_clean45disease_free_stt, exclude = NULL)
# table(kirc_clean5$ethnicity, exclude = NULL)
# table(kirc_clean5$histology_grd, exclude = NULL)
# table(kirc_clean5$hemoglobin, exclude = NULL)
# table(kirc_clean5$neoadj_therapy, exclude = NULL)
# table(kirc_clean5$prior_cancer, exclude = NULL)
# table(kirc clean5$tumor lateral, exclude = NULL)
# table(kirc_clean5$primer_ln_ind3, exclude = NULL)
# table(kirc_clean5platelet, exclude = NULL)
# table(kirc_clean5$tissue_prospect, exclude = NULL)
```

```
# table(kirc_clean5$race, exclude = NULL)
# table(kirc_clean5$tissue_retrospect, exclude = NULL)
# table(kirc_clean5$serum_ca, exclude = NULL)
# table(kirc_clean5$sex, exclude = NULL)
# table(kirc_clean5$tissue_site, exclude = NULL)
# table(kirc_clean5$person_neoplasm_stt, exclude = NULL)
# table(kirc_clean5$person_neoplasm_stt, exclude = NULL)
```

8. Saving dataset

```
write_csv(kirc_clean5, path = "data/kirc_clinic.csv")
rm(kirc_clean4, kirc_clean3, kirc_clean2, kirc_clean1, kirc_clean0, kirc_clean)
```

Further analysis

- A correlation analysis with t-test and ANOVA checking significant distinction between variables acording their vital status.
- A logistic regression analysis of each clinical variable weight.

sessionInfo()

```
## R version 3.6.3 (2020-02-29)
## Platform: x86_64-pc-linux-gnu (64-bit)
## Running under: Ubuntu 18.04.4 LTS
##
## Matrix products: default
          /usr/lib/x86_64-linux-gnu/openblas/libblas.so.3
## LAPACK: /usr/lib/x86_64-linux-gnu/libopenblasp-r0.2.20.so
## locale:
## [1] LC_CTYPE=pt_BR.UTF-8
                                   LC_NUMERIC=C
## [3] LC_TIME=pt_BR.UTF-8
                                   LC_COLLATE=en_US.UTF-8
   [5] LC_MONETARY=pt_BR.UTF-8
                                   LC_MESSAGES=en_US.UTF-8
##
  [7] LC_PAPER=pt_BR.UTF-8
                                   LC_NAME=C
  [9] LC ADDRESS=C
                                   LC TELEPHONE=C
## [11] LC_MEASUREMENT=pt_BR.UTF-8 LC_IDENTIFICATION=C
## attached base packages:
## [1] stats
                graphics grDevices utils
                                               datasets methods
                                                                   base
##
## other attached packages:
## [1] finalfit_1.0.1 skimr_2.1.1
                                        forcats_0.5.0
                                                        stringr_1.4.0
## [5] dplyr_0.8.5
                       purrr_0.3.4
                                        readr_1.3.1
                                                        tidyr_1.0.3
## [9] tibble_3.0.1
                        ggplot2_3.3.0
                                        tidyverse_1.3.0
## loaded via a namespace (and not attached):
                        lubridate_1.7.8 lattice_0.20-41 assertthat_0.2.1
## [1] Rcpp_1.0.4.6
## [5] digest_0.6.25
                         utf8_1.1.4
                                          R6_2.4.1
                                                           cellranger_1.1.0
## [9] repr_1.1.0
                        backports_1.1.6 reprex_0.3.0
                                                           evaluate_0.14
## [13] httr_1.4.1
                        highr_0.8
                                                           rlang_0.4.6
                                          pillar_1.4.4
                        rstudioapi_0.11 Matrix_1.2-18
## [17] readxl_1.3.1
                                                           rmarkdown_2.1
```

```
## [21] labeling_0.3
                         splines_3.6.3
                                          munsell_0.5.0
                                                           broom_0.5.6
## [25] compiler_3.6.3
                         modelr_0.1.7
                                          xfun_0.13
                                                           pkgconfig_2.0.3
## [29] base64enc_0.1-3
                         htmltools_0.4.0
                                          tidyselect_1.1.0 fansi_0.4.1
                                                           grid_3.6.3
## [33] crayon_1.3.4
                         dbplyr_1.4.3
                                          withr_2.2.0
## [37] nlme_3.1-147
                         jsonlite_1.6.1
                                          gtable_0.3.0
                                                           lifecycle_0.2.0
## [41] DBI_1.1.0
                         magrittr_1.5
                                          scales_1.1.1
                                                           cli_2.0.2
## [45] stringi_1.4.6
                         farver_2.0.3
                                          fs_1.4.1
                                                           mice_3.8.0
## [49] xml2_1.3.2
                                                           vctrs_0.3.0
                         ellipsis_0.3.0
                                          generics_0.0.2
                         tools_3.6.3
## [53] boot_1.3-25
                                          glue_1.4.0
                                                           hms_0.5.3
## [57] survival_3.1-12
                                          colorspace_1.4-1 rvest_0.3.5
                         yaml_2.2.1
                         haven_2.2.0
## [61] knitr_1.28
```