# FEDERAL STATE AUTONOMOUS EDUCATIONAL INSTITUTE OF HIGHER EDUCATION ITMO UNIVERSITY

Report on learning practice # 2

Analysis of multivariate random variables

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## Non-parametric estimation of PDF for MRV

Figures 1-9 depict non-parametric estimations in form of histogram and using kernel density function for the chosen random variables.

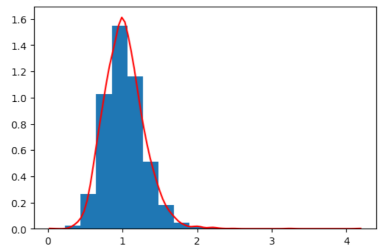


Figure 1 – Non-parametric estimation of PDF in form of histogram and using kernel density function for "Reproduction rate"

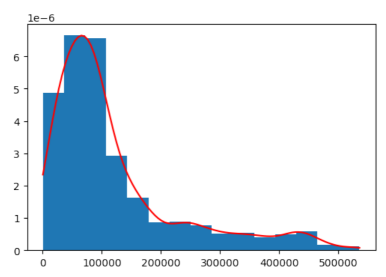


Figure 2 – Non-parametric estimation of PDF in form of histogram and using kernel density function for "Total cases per million"

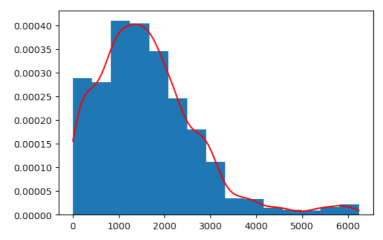


Figure 3 – Non-parametric estimation of PDF in form of histogram and using kernel density function for "Total deaths per million"

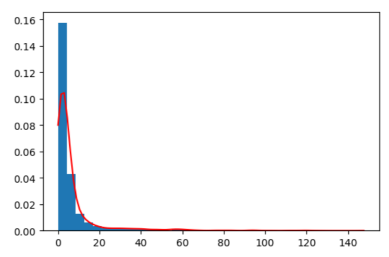


Figure 4 – Non-parametric estimation of PDF in form of histogram and using kernel density function for "New tests smoothed per thousand"

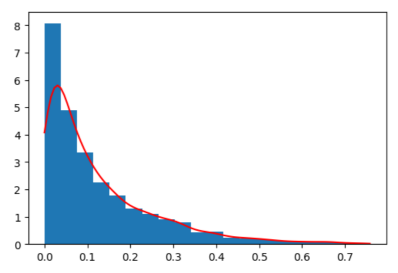


Figure 5 – Non-parametric estimation of PDF in form of histogram and using kernel density function for "Positive rate"

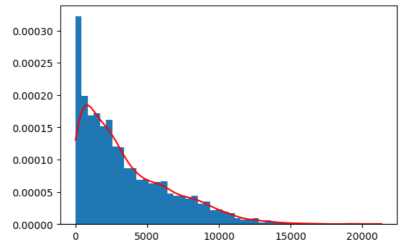


Figure 6 – Non-parametric estimation of PDF in form of histogram and using kernel density function for "New vaccinations smoothed per million"

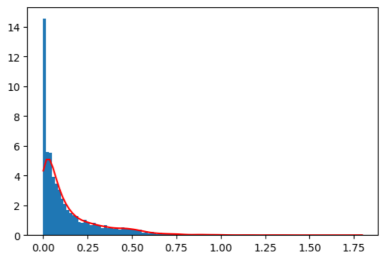


Figure 7 – Non-parametric estimation of PDF in form of histogram and using kernel density function for "New people vaccinated smoothed per hundred"

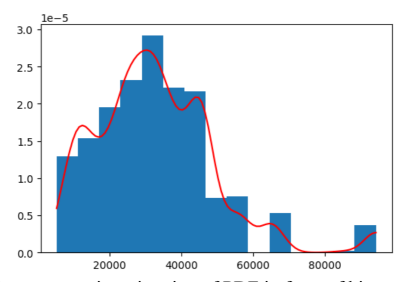


Figure 8 – Non-parametric estimation of PDF in form of histogram and using kernel density function for "GDP per capita"

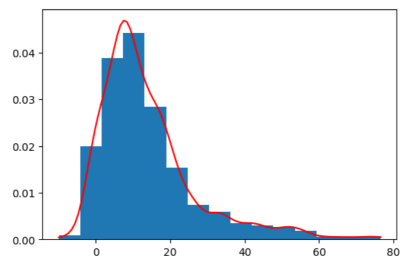


Figure 9 – Non-parametric estimation of PDF in form of histogram and using kernel density function for "Excess mortality cumulative"

Among given above-mentioned random variables, the first one is a target and the rest are predictors. Figures 10-17 show us joint probability density plots in form of histogram for variations between target and each predictor. Each image also contains points of different color. Colors represent data for different continents.

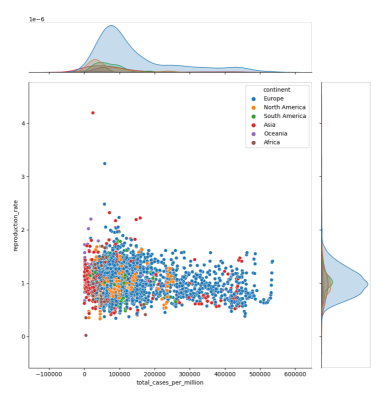


Figure 10 – Joint probability density plot in form of histogram between target "Reproduction rate" and predictor "Total cases per million"

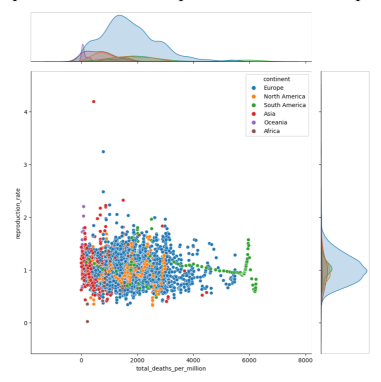


Figure 11 – Joint probability density plot in form of histogram between target "Reproduction rate" and predictor "Total deaths per million"

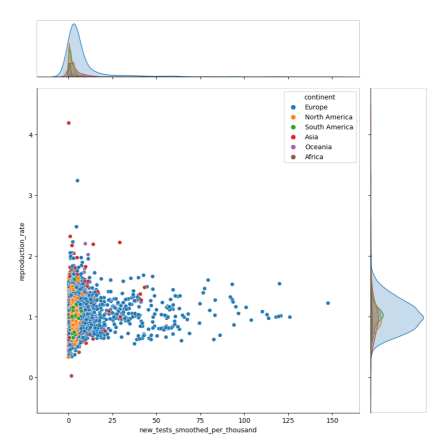


Figure 12 – Joint probability density plot in form of histogram between target "Reproduction rate" and predictor "New tests smoothed per thousand"

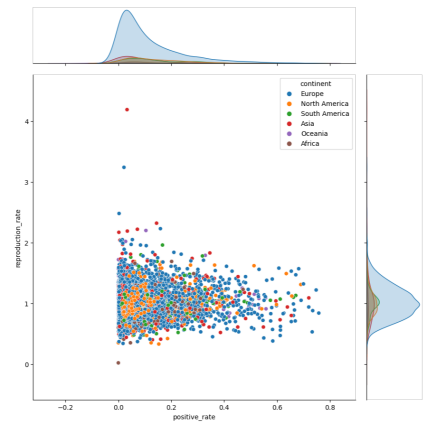


Figure 13 – Joint probability density plot in form of histogram between target "Reproduction rate" and predictor "Positive rate"

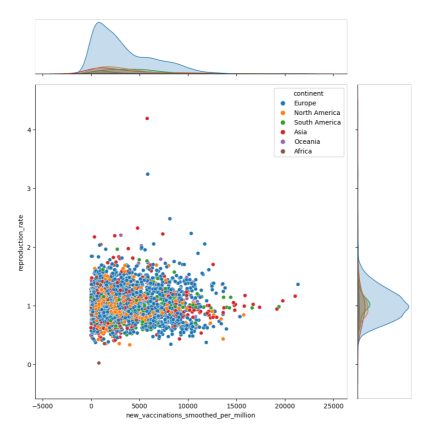


Figure 14 – Joint probability density plot in form of histogram between target "Reproduction rate" and predictor "New vaccinations smoothed per million"

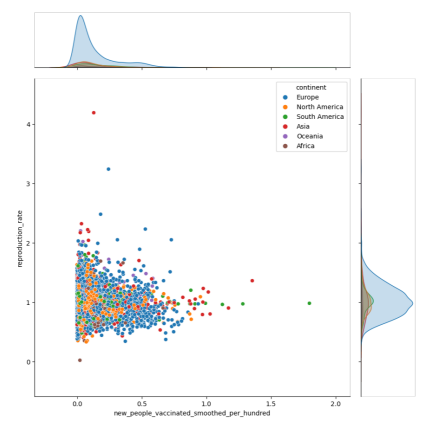


Figure 15 – Joint probability density plot in form of histogram between target "Reproduction rate" and predictor "New people vaccinated smoothed per hundred"

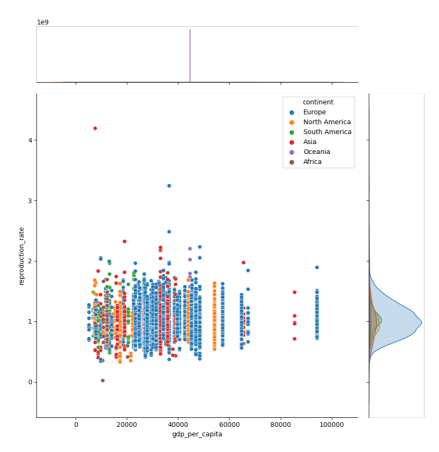


Figure 16 – Joint probability density plot in form of histogram between target "Reproduction rate" and predictor "GDP per capita"

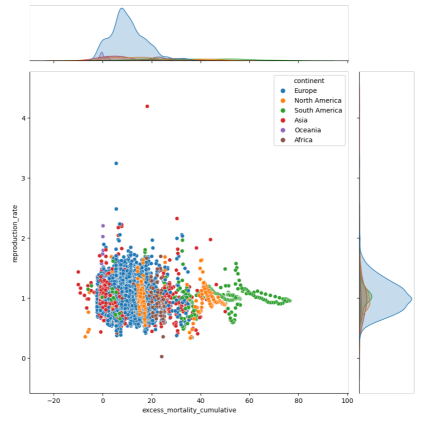


Figure 17 – Joint probability density plot in form of histogram between target "Reproduction rate" and predictor "Excess mortality cumulative"

Figures 18 - 25 show us joint probability density plots in form of kernel density function for variations between target and each predictor. Each image also contains lines of different color. Colors represent data for different continents.

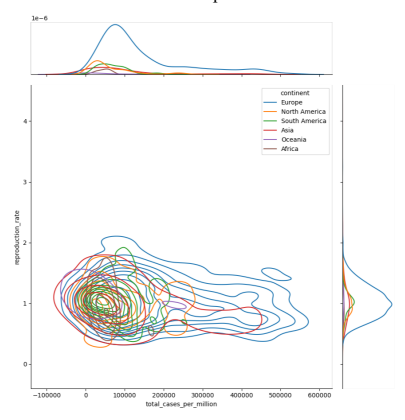


Figure 18 – Joint probability density plot in form of kernel density function between target "Reproduction rate" and predictor "Total cases per million"

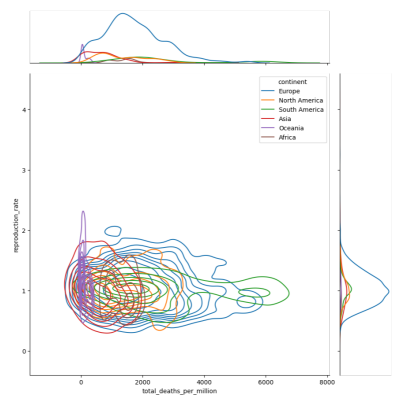


Figure 19 – Joint probability density plot in form of kernel density function between target "Reproduction rate" and predictor "Total deaths per million"

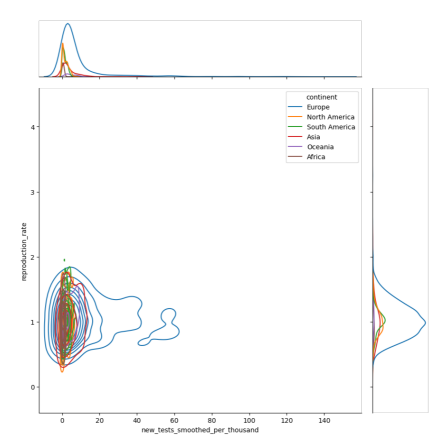


Figure 20 – Joint probability density plot in form of kernel density function between target "Reproduction rate" and predictor "New tests smoothed per thousand"

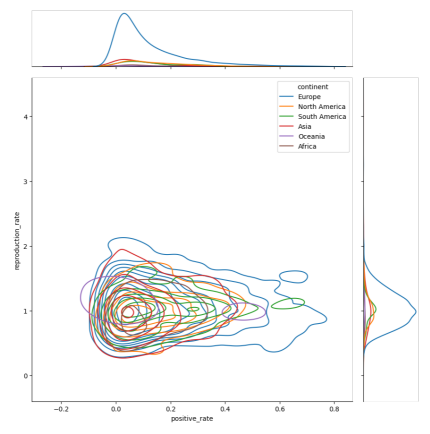


Figure 21 – Joint probability density plot in form of kernel density function between target "Reproduction rate" and predictor "Positive rate"

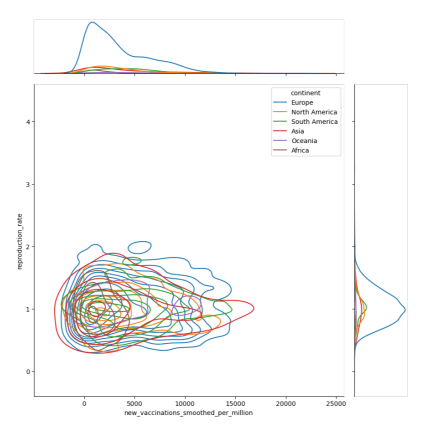


Figure 22 – Joint probability density plot in form of kernel density function between target "Reproduction rate" and predictor "New vaccinations smoothed per million"

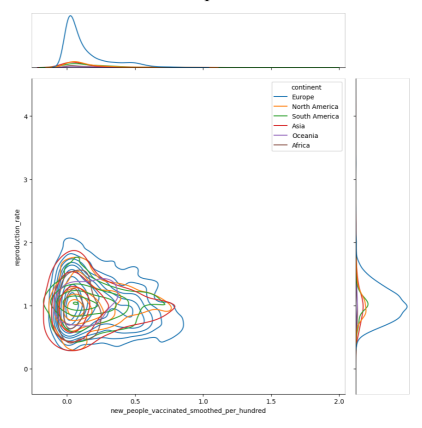


Figure 23 – Joint probability density plot in form of kernel density function between target "Reproduction rate" and predictor "New people vaccinated smoothed per hundred"

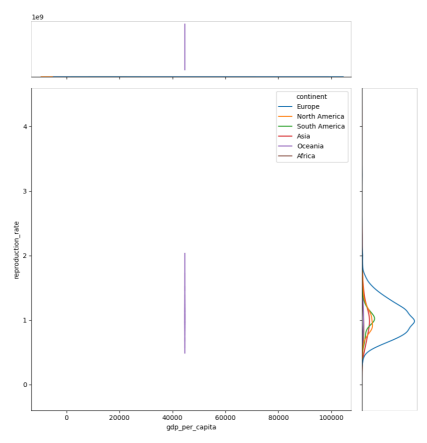


Figure 24 – Joint probability density plot in form of kernel density function between target "Reproduction rate" and predictor "GDR per capita"

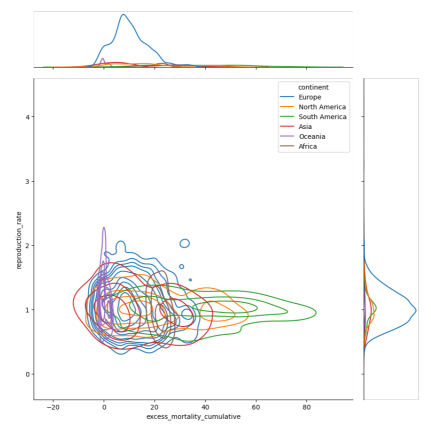


Figure 25 – Joint probability density plot in form of kernel density function between target "Reproduction rate" and predictor "Excess mortality cumulative"

The next Figure 26 depicts pairwise joint probability density histogram between five variables.

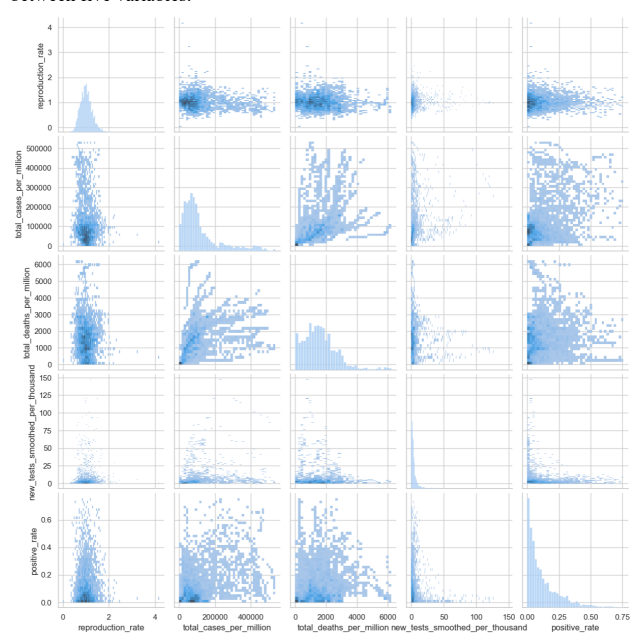


Figure 26 – Pairwise joint probability density between five random variables **Estimation of multivariate mean and variance** 

X	E(X)	Var(X)
Reproduction rate	1.023385	$7.484235 \cdot 10^{-2}$
Total cases per million	119878.364158	$1.217849 \cdot 10^{10}$
Total deaths per million	1617.485002	$1.206946 \cdot 10^6$
New tests smoothed per	6.443801	$1.595720 \cdot 10^2$
thousand		
Positive rate	0.123713	$1.702659 \cdot 10^{-2}$
New vaccinations	3508.352299	$1.004960 \cdot 10^7$
smoothed per million		

New people vaccinated	0.139452	$3.113892 \cdot 10^{-2}$	
smoothed per hundred			
GDP per capita	32880.455968	$2.839982 \cdot 10^{8}$	
Excess mortality	13.950828	$1.721878 \cdot 10^2$	
cumulative			

Table 1 – Estimation of multivariate mathematical expectation and variance for the chosen variables

# Non-parametric estimation of conditional distributions, mathematical expectations and variances

Let condition C be Reproduction rate = 1.0. So, Figure 27 depicts distribution of variables under this condition. And Table 2 contains conditional mathematical expectation and conditional variance.

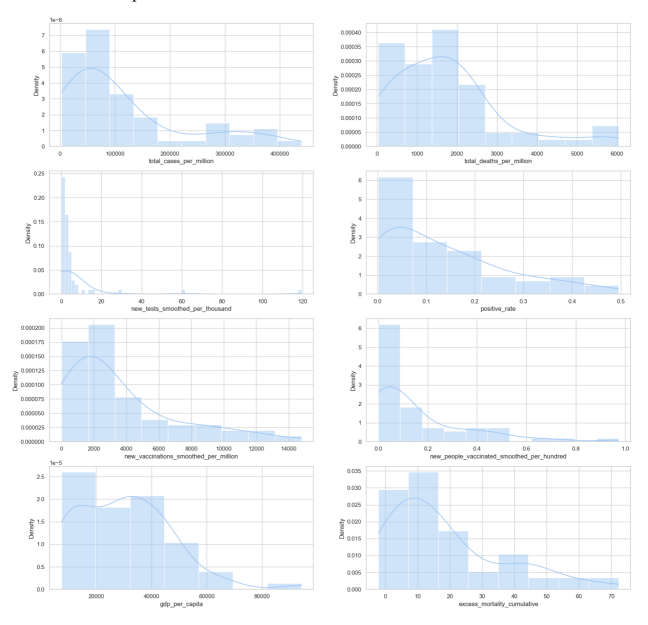


Figure 27 – Conditional distributions of variables

X	E(X C)	Var(X C)
Total cases per million	113775.970403	$1.250462 \cdot 10^{10}$
Total deaths per million	1743.478855	$1.921855 \cdot 10^6$
New tests smoothed per	6.184790	$1.875900 \cdot 10^2$
thousand		
Positive rate	0.135581	$1.613816 \cdot 10^{-2}$
New vaccinations	3569.225806	$1.184156 \cdot 10^7$
smoothed per million		
New people vaccinated	0.153016	$4.130244 \cdot 10^{-2}$
smoothed per hundred		
GDP per capita	30048.161435	$3.001686 \cdot 10^8$
Excess mortality	18.273065	$3.198207 \cdot 10^2$
cumulative		

Table 2 – Conditional mathematical expectation and conditional variance for predictors

# Estimation of pair correlation coefficients, confidence intervals for them and significance levels

Figure 28 shows pair correlation coefficients represented as a heatmap diagram. Table 3 contains information about confidence intervals for them and significance level.

X	Y	Coefficient	Conf. interval	Signif. level
Total cases per million	Total deaths per million	0.3592	(0.3427, 0.4092)	0.000000
New vaccinations smoothed per million	Excess mortality cumulative	-0.0630	(-0.0963, -0.0298)	0.000202
New people vaccinated smoothed per hundred	Excess mortality cumulative	-0.0031	(-0.0363,0.0302)	0.856135
GDP per capita	Excess mortality cumulative	-0.5742	(-0.6871, -0.6206)	0.000000
Total cases per million	New vaccinations smoothed per million	-0.3183	(-0.3629, -0.2965)	0.000000

Table 3 – Pair correlation coefficients confidence intervals and significance levels

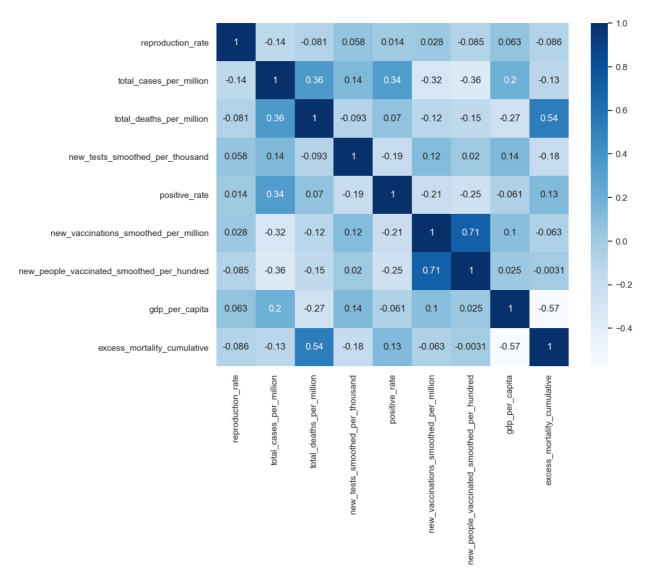


Figure 28 – Pair correlation coefficients

Let's highlight the most significant correlations in our data:

- new\_people\_vaccinated\_smoothed\_per\_hundred & new\_vaccinations\_smoothed\_per\_million: 0.71 (strong positive correlation);
- gdp\_per\_capita & excess\_mortality\_cumulative: -0.57 (average negative correlation);
- excess\_mortality\_cumulative & total\_deaths\_per\_million: 0.54 (average positive correlation);
- total deaths per million & total cases per million: 0.36 (average positive correlation)
- new\_people\_vaccinated\_smoothed\_per\_hundred & total\_cases\_per\_million: -0.36 (average negative correlation);
- positive rate & total cases per million: 0.34 (average positive correlation);
- new\_vaccinations\_smoothed\_per\_million & total\_cases\_per\_million: -0.32 (average negative correlation);
- total deaths per million & gdp per capita: -0.27 (average negative correlation);

The remaining correlations have a small dependence and their contribution is insignificant.

#### Task formulation for regression, multivariate correlation

Training a model for predicting "Reproduction rate" based on "Total cases per million", "Total deaths per million", "New tests smoothed per thousand", "Positive rate", "New vaccinations smoothed per million", "New people vaccinated smoothed per hundred", "GDP per capita", "Excess mortality cumulative" variables.

### Regression model, multicollinearity and regularization

We have built regression model. Table 4 consist of metrics obtained after fitting ordinal regression model.

Metric	Value	
Mean Absolute Error	0.192338	
Mean Squared Error	0.653560	
R2 Score	0.063770	
Mean Absolute Percentage Error	20.379919	

Table 4 – Ordinal regression model metrics

After that we have applied Lasso regularization to model (with  $\alpha = 0.1$ ). Table 5 consist of metrics obtained after fitting Lasso-regression model.

Metric	Value	
Mean Absolute Error	0.193504	
Mean Squared Error	0.067059	
R2 Score	0.039364	

Table 5 – Lasso-regression model metrics

Finally, a LARS regression was used. Table 6 shows its metrics.

Metric	Value	
Mean Absolute Error	0.192338	
Mean Squared Error	0.065356	
R2 Score	0.063770	

Table 6 – LARS model metrics

Also  $2^{nd}$  degree polynomial regression models was fitted.

#### Quality analysis for regression model

Figure 29 illustrates real values and values predicted by linear model for "Reproduction rate" variable. Figure 30 contains Quantile-Quantile plot for this model. Distribution of residuals for linear model is depicted at Figure 31. Table 7 contains confident intervals for regression coefficients for linear regression.

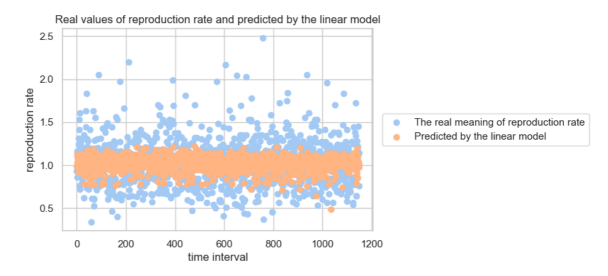


Figure 29 – Real and predicted points by linear model

Figure 32 illustrates real values and values predicted by polynomial model for "Reproduction rate" variable. Figure 33 contains Quantile-Quantile plot for this model.

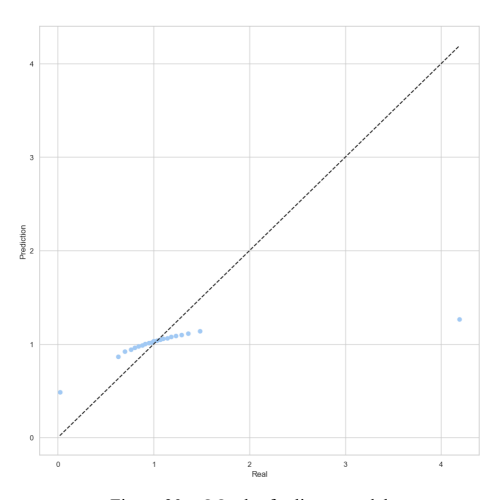


Figure 30 – QQ-plot for linear model

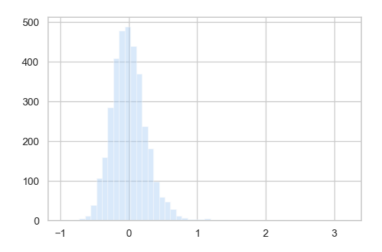


Figure 31 – Distribution of residuals for linear model

Let's apply the Anderson test to check the distribution of residuals for normality:

Statistic: 14.264

significance level: critical values

15.0: 0.575, data does not look normal (reject H0) 10.0: 0.655, data does not look normal (reject H0) 5.0: 0.786, data does not look normal (reject H0) 2.5: 0.917, data does not look normal (reject H0) 1.0: 1.091, data does not look normal (reject H0)

We also apply Kolmogorov-Smirnov Test and Cramer-Von Mises Test to check the distribution of residuals for normality:

Kolmogorov-Smirnov Test: 0.00001 Cramer-Von Mises Test: 0.00000

So, residuals are not distributed normally.

X	Interval left	Interval right
Total cases per million	$-8.315253 \cdot 10^{-7}$	$-3.065669 \cdot 10^{-7}$
Total deaths per million	$7.420195 \cdot 10^{-5}$	$1.260864 \cdot 10^{-4}$
New tests smoothed per thousand	$4.017558 \cdot 10^{-3}$	$7.307548 \cdot 10^{-3}$
Positive rate	$5.893598 \cdot 10^{-1}$	$9.321734 \cdot 10^{-1}$
New vaccinations smoothed per million	$1.731372 \cdot 10^{-5}$	$3.541832 \cdot 10^{-5}$
New people vaccinated smoothed per	$-2.714395 \cdot 10^{-1}$	$6.117958 \cdot 10^{-2}$
hundred		
GDP per capita	$1.425285 \cdot 10^{-5}$	$1.633377 \cdot 10^{-5}$
Excess mortality cumulative	$8.211649 \cdot 10^{-3}$	$1.238924 \cdot 10^{-2}$

Table 7 – Confidence interval of regression coefficients

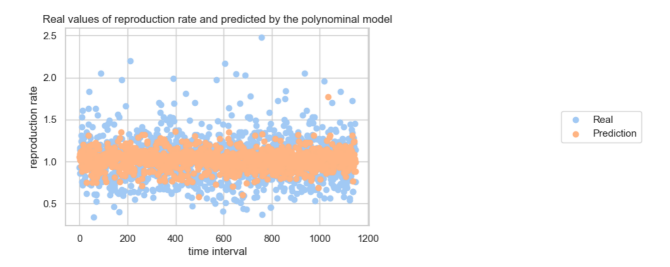


Figure 32 – Real and predicted points by polynomial model

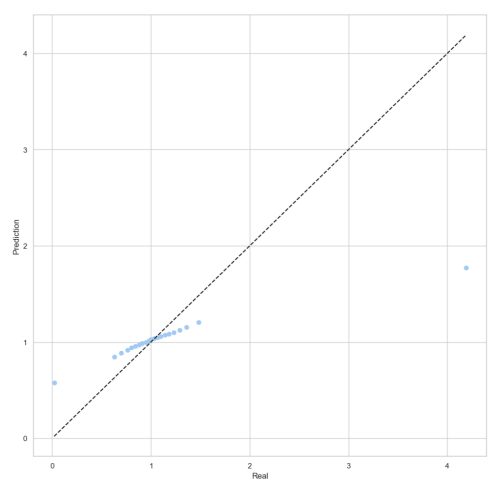


Figure 33 – QQ-plot for polynomial model

### **Appendix**

```
#!/usr/bin/env python
# coding: utf-8
# In[1]:
import matplotlib as mpl
import matplotlib.pyplot as plt
import numpy as np
import pandas as pd
import scipy.stats
import seaborn as sns
from sklearn.model selection import train test split
from sklearn.linear model import Lars
from sklearn.linear model import LinearRegression
from sklearn.linear model import ElasticNet
from sklearn.metrics import mean absolute error
from sklearn.metrics import mean squared error
from sklearn.metrics import r2 score
from sklearn.linear model import LassoLarsIC
# In[2]:
pd.set option('display.max columns', 70)
# In[3]:
source df = pd.read csv('owid-covid-data.csv')
source df
# ## 1. Make a non-parametric estimation of PDF in form of histogram and
using kernel density function for MRV (or probability law in case of
discrete MRV).
# Dataset content:
# - total cases per million - Total confirmed cases of COVID-19 per
1,000,000 people
# - total deaths per million - Total deaths attributed to COVID-19 per
1,000,000 people
# - reproduction rate - Real-time estimate of the effective reproduction
rate (R) of COVID-19
# - new tests smoothed per thousand - New tests for COVID-19 (7-day
smoothed) per 1,000 people
# - positive rate - The share of COVID-19 tests that are positive, given
as a rolling 7-day average (this is the inverse of tests per case)
# - new vaccinations smoothed per million - New COVID-19 vaccination doses
administered (7-day smoothed) per 1,000,000 people in the total population
# - new people vaccinated smoothed per hundred - Daily number of people
receiving their first vaccine dose (7-day smoothed) per 100 people in the
total population
# - gdp per capita - Share of the population that is 70 years and older in
2015
```

```
# - excess_mortality_cumulative - Percentage difference between the
cumulative number of deaths since 1 January 2020 and the cumulative
projected deaths for the same period based on previous years
# In[4]:
source df.continent.unique()
# In[5]:
columns = ['continent',
           'reproduction rate',
           'total_cases_per_million',
            'total deaths per million',
            'new tests smoothed per thousand',
            'positive rate',
            'new vaccinations smoothed per million',
            'new people vaccinated smoothed per hundred',
            'gdp per capita',
            'excess mortality cumulative']
df = source_df[columns].copy().dropna()
df.index = np.arange(0, len(df))
df
# In[6]:
target = columns[1]
predictors = columns[2:]
categorial = columns[0]
# In[7]:
df.describe()
# In[8]:
from scipy.stats import kde
def kernel density estimation(x, bins):
    density = kde.gaussian_kde(x)
    xgrid = np.linspace(x.min(), x.max(), 100)
    mpl.rcParams['figure.dpi'] = 100
    plt.hist(x, bins=bins, density = True)
    plt.plot(xgrid, density(xgrid), 'r-')
# In[9]:
cases = df['total cases per million']
kernel density estimation (cases, 15)
```

```
# In[10]:
deaths = df['total deaths per million']
kernel density estimation (deaths, 15)
# In[11]:
reproduction = df['reproduction rate']
kernel density estimation (reproduction, 20)
# In[12]:
tests = df['new_tests_smoothed_per_thousand']
kernel_density_estimation(tests, 35)
# In[13]:
positive = df['positive_rate']
kernel density estimation (positive, 20)
# In[14]:
vaccinations = df['new_vaccinations_smoothed_per_million']
kernel density estimation (vaccinations, 50)
# In[15]:
vac people = df['new people vaccinated smoothed per hundred']
kernel density estimation (vac people, 110)
# In[16]:
gdp = df['gdp per capita']
kernel density estimation (gdp, 15)
# In[17]:
emc = df['excess mortality cumulative']
kernel density estimation (emc, 15)
# In[18]:
```

```
for predictor in predictors:
    sns.jointplot(data=df, x=predictor, y=target, hue=categorial,
height=9)
   plt.plot()
# In[19]:
for predictor in predictors:
    sns.jointplot(data=df, x=predictor, y=target, kind='kde',
hue=categorial, height=9)
    plt.plot()
# In[20]:
# sns.pairplot(data=df[predictors + [target]], kind='kde',
plot kws=\{'levels':[0.1,0.2,0.3,0.4,0.5,0.6,0.7,0.8,0.9]\}
# In[21]:
sns.set_theme(style='whitegrid', palette='pastel')
ax = sns.pairplot(df[['reproduction rate',
           'total_cases_per_million',
            'total_deaths_per_million',
            'new_tests_smoothed_per_thousand',
            'positive rate']], kind='hist', diag kind='hist')
plt.show()
# In[22]:
# ax = sns.pairplot(df[['reproduction rate',
             'total cases per million',
              'total deaths per million',
#
              'new tests smoothed per thousand',
              'positive_rate']], diag_kind='kde')
# ax.map_lower(sns.kdeplot, levels=5, color='.1')
# plt.show()
# ## 2. Estimation of Multivariate Mathematical Expectation and Variance
# In[53]:
df[columns].mean()
# In[54]:
df[columns].var()
```

```
# ## 3. Non-parametric estimation of conditional distributions,
mathematical expectations and variances
# In[25]:
df conditional = df[df.reproduction rate == 1]
df conditional
# In[26]:
df_conditional[['total cases per million',
            'total deaths per million',
            'new tests smoothed per thousand',
            'positive rate',
            'new vaccinations smoothed per million',
            'new people vaccinated smoothed per hundred',
            'gdp per capita',
            'excess mortality cumulative']].mean()
# In[27]:
df conditional[['total cases per million',
            'total deaths per million',
            'new tests smoothed per thousand',
            'positive rate',
            'new vaccinations smoothed per million',
            'new people vaccinated smoothed per hundred',
            'gdp per capita',
            'excess mortality cumulative']].var()
# In[28]:
figure, ax = plt.subplots(4, 2, figsize=(20, 20))
sns.set theme(style='whitegrid', palette='pastel')
total cases per million = sns.histplot(df.total cases per million,
ax=ax[0, 0], kde=True, stat='density')
total cases per million.set(xlabel='total cases per million')
total deaths per million = sns.histplot(df.total deaths per million,
ax=ax[0, 1], kde=True, stat='density')
total deaths per million.set(xlabel='total deaths per million')
new tests smoothed per thousand =
sns.histplot(df.new tests smoothed per thousand, ax=ax[1, 0], kde=True,
stat='density')
new tests smoothed per thousand.set(xlabel='new tests smoothed per thousan
d')
positive rate = sns.histplot(df.positive rate, ax=ax[1, 1], kde=True,
stat='density')
positive rate.set(xlabel='positive rate')
```

```
new vaccinations smoothed per million =
sns.histplot(df.new vaccinations smoothed per million, ax=ax[2, 0],
kde=True, stat='density')
new vaccinations smoothed per million.set(xlabel='new vaccinations smoothe
d per million')
new people vaccinated smoothed per hundred =
sns.histplot(df.new people vaccinated smoothed per hundred, ax=ax[2, 1],
kde=True, stat='density')
new people vaccinated smoothed per hundred.set(xlabel='new people vaccinat
ed smoothed per hundred')
gdp per capita = sns.histplot(df.gdp per capita, ax=ax[3, 0], kde=True,
stat='density')
gdp_per_capita.set(xlabel='gdp per capita')
excess mortality cumulative = sns.histplot(df.excess mortality cumulative,
ax=ax[3, 1], kde=True, stat='density')
excess mortality cumulative.set(xlabel='excess mortality cumulative')
plt.show()
# In[29]:
figure, ax = plt.subplots(4, 2, figsize=(20, 20))
sns.set theme(style='whitegrid', palette='pastel')
total cases per million =
sns.histplot(df conditional.total cases per million, ax=ax[0, 0],
kde=True, stat= density')
total cases per million.set(xlabel='total cases per million')
total deaths per million =
sns.histplot(df conditional.total deaths per million, ax=ax[0, 1],
kde=True, stat='density')
total deaths per million.set(xlabel='total deaths per million')
new tests smoothed per thousand =
sns.histplot(df conditional.new tests smoothed per thousand, ax=ax[1, 0],
kde=True, stat='density')
new_tests_smoothed_per_thousand.set(xlabel='new_tests_smoothed_per_thousan
d')
positive rate = sns.histplot(df conditional.positive rate, ax=ax[1, 1],
kde=True, stat='density')
positive rate.set(xlabel='positive rate')
new vaccinations smoothed per million =
sns.histplot(df_conditional.new_vaccinations_smoothed_per_million,
ax=ax[2, 0], kde=True, stat='density')
new vaccinations smoothed per million.set(xlabel='new vaccinations smoothe
d_per_million')
new people vaccinated smoothed per hundred =
sns.histplot(df_conditional.new_people_vaccinated_smoothed_per_hundred,
ax=ax[2, 1], kde=True, stat='density')
new_people_vaccinated_smoothed_per_hundred.set(xlabel='new people vaccinat
ed smoothed per hundred')
```

```
gdp per capita = sns.histplot(df conditional.gdp per capita, ax=ax[3, 0],
kde=True, stat='density')
gdp per capita.set(xlabel='gdp per capita')
excess mortality cumulative =
sns.histplot(df conditional.excess mortality cumulative, ax=ax[3, 1],
kde=True, stat='density')
excess mortality cumulative.set(xlabel='excess mortality cumulative')
plt.show()
# ## 4. Estimation of pair correlation coefficients, confidence intervals
for them and significance levels
# In[55]:
figure, ax = plt.subplots(1, 1, figsize=(10, 8))
sns.set theme(style='whitegrid', palette='pastel')
sns.heatmap(df[columns].corr(), cmap='Blues', annot=True)
plt.show()
# In[31]:
def return_estimation(array, target, alpha=0.05):
    r, p value = scipy.stats.pearsonr(array, target)
    r to z = np.arctanh(r) # Matches Fisher transform
    se = 1 / np.sqrt(array.size - 3) # Corresponding standard deviation
    z = scipy.stats.norm.ppf(1 - alpha / 2)
    lo z, hi z = r to z-z*se, r to z+z*se
\# lo_z, hi_z = np.tanh(lo_z), np.tanh(hi_z)
    print('Correlation Coefficient: {:.4f}'.format(r))
    print('Confidence Interval for the correlation coefficient: ({:.4f},
\{:.4f\})'.format(lo z, hi z))
    print('Significance Level: {:.6f}'.format(p_value))
# In[32]:
return estimation(df.total cases per million, df.total deaths per million)
# In[33]:
return estimation(df.new vaccinations smoothed per million,
df.excess mortality cumulative)
# In[34]:
```

```
return estimation(df.new people vaccinated smoothed per hundred,
df.excess mortality cumulative)
# In[35]:
return estimation(df.gdp per capita, df.excess mortality cumulative)
# In[36]:
return estimation(df.total cases per million,
df.new vaccinations smoothed per million)
# ## 5. Task formulation for regression. Estimate multivariate correlation
(target - predictors)
# Training a model for predicting reproduction rate based on
total_cases_per_million, total deaths per million,
new tests smoothed per thousand, positive rate,
new vaccinations smoothed per million,
new people vaccinated smoothed per hundred, gdp per capita,
excess mortality cumulative variables.
# In[56]:
figure, ax = plt.subplots(1, 1, figsize=(10, 8))
sns.set_theme(style='whitegrid', palette='pastel')
sns.heatmap(df[columns].corr(), cmap='Blues', annot=True)
plt.show()
# ## 6. Build regression model and make an analysis of multicollinearity
and regularization (if needed)
# In[38]:
# Highlight predictors
X = df[['total cases per million',
            'total deaths per million',
            'new tests smoothed per thousand',
            'positive rate',
            'new vaccinations smoothed per million',
            'new_people_vaccinated smoothed per hundred',
            'gdp per capita',
            'excess mortality cumulative']]
# Allocate the target variable
y = df[['reproduction rate']]
# Division into training and test samples
X train, X test, y_train, y_test = train_test_split(X, y, test_size=0.33,
random state=42)
X = []
for i in range(len(y test)):
```

```
x.append(i)
# Create a linear regression model
reg = LinearRegression(normalize=True)
# Train a linear regression model
reg.fit(X train, y train)
# Forecast on a test sample
y_pred = reg.predict(X test)
params = np.append(reg.intercept , reg.coef )
# In[39]:
# Calculate regression metrics
mae = mean absolute error(y test, y pred)
mse = mean_squared_error(y_test, y_pred)
r2 = r2_score(y_test, y_pred)
print('Mean absolute error = ', mae)
print('Mean squared error = ', mse)
print('R2 Score = ', r2)
# In[40]:
def mean absolute percentage_error(y_true, y_pred):
    return np.mean(np.abs((y true - y pred) / y true)) * 100
mape = mean_absolute_percentage_error(y_test, y_pred)
print('Mean absolute percentage error = ', mape)
# In[41]:
from sklearn import linear model
clf = linear model.Lasso(alpha=0.1)
clf.fit(X_train, y_train)
print(clf.coef )
# In[42]:
y pred lasso = clf.predict(X test)
mae lasso = mean absolute error(y test, y pred lasso)
mse lasso = mean squared error(y test, y pred lasso)
r2_lasso = r2_score(y_test, y_pred_lasso)
print('Mean Absolute Error with lasso =', mae_lasso)
print('Mean Squared Error with lasso =', mse_lasso)
print('R2 Score with lasso =', r2_lasso)
# In[43]:
lars = Lars()
lars.fit(X_train, y_train)
mae = mean_absolute_error(y_test, lars.predict(X_test))
mse = mean squared error(y test, lars.predict(X test))
```

```
r2 = r2_score(y_test, lars.predict(X_test))
print('Mean Absolute Error with lars =', mae)
print('Mean Squared Error with lars =', mse)
print('R2 Score with lars =', r2)
# ## 7. Analyze the quality of regression model (distribution of
residuals, determination coefficient)
# In[44]:
# Graph of real and predicted values
plt.scatter(x, y test, label = u'The real meaning of reproduction rate')
plt.scatter(x, y_pred, label = u'Predicted by the linear model')
plt.title(u'Real values of reproduction rate and predicted by the linear
model')
plt.legend(loc="center right", borderaxespad=0.1, bbox to anchor=(1.7,
0.5)
plt.xlabel(u'time interval')
plt.ylabel(u'reproduction rate')
# In[45]:
y pred all = np.array(reg.predict(X))
# In[46]:
# Plotting a quantile biplot based on real and predicted values
percs = np.linspace(0, 100, 21)
qn first = np.percentile(y, percs)
qn second = np.percentile(y_pred_all, percs)
plt.figure(figsize=(12, 12))
min qn = np.min([qn first.min(), qn second.min()])
max qn = np.max([qn first.max(), qn second.max()])
x = np.linspace(min qn, max qn)
plt.plot(qn_first, qn_second, ls="", marker="o", markersize=6)
plt.plot(x, x, color="k", ls="--")
plt.xlabel(u'Real')
plt.ylabel(u'Prediction')
# In[47]:
# Building and training of the 2nd degree polynomial regression
from sklearn.preprocessing import PolynomialFeatures
poly = PolynomialFeatures(2)
X train new = poly.fit transform(X train)
poly = PolynomialFeatures(2)
X test new = poly.fit transform(X test)
reg = LinearRegression(normalize=True)
reg.fit(X train new, y train)
y pred poly = reg.predict(X test new)
```

```
X new = poly.fit transform(X)
y pred poly all = reg.predict(X new)
# In[48]:
# Plotting a quantile biplot based on real and predicted values
percs = np.linspace(0, 100, 21)
qn first = np.percentile(y, percs)
qn second = np.percentile(y pred poly all , percs)
plt.figure(figsize=(12, 12))
min qn = np.min([qn first.min(), qn second.min()])
max qn = np.max([qn first.max(), qn second.max()])
x = np.linspace(min qn, max qn)
plt.plot(qn first, qn second, ls="", marker="o", markersize=6)
plt.plot(x, x, color="k", ls="--")
plt.xlabel(u'Real')
plt.ylabel(u'Prediction')
# In[58]:
mae poly = mean absolute error(y test, y pred poly)
mse poly = mean squared error(y test, y pred poly)
r2 poly = r2_score(y_test, y_pred_poly)
print('Mean absolute error with Polynomial model = ', mae poly)
print('Mean squared error with Polynomial model = ', mse poly)
print('R2 Score with Polynomial model = ', r2)
# In[49]:
# Visualization of real and predicted values with polynominal model
for i in range(len(y test)):
    x.append(i)
plt.scatter(x, y_test, label = u'Real')
plt.scatter(x, y pred poly, label = u'Prediction')
plt.title(u'Real values of reproduction rate and predicted by the
polynominal model')
plt.legend(loc="center right", borderaxespad=0.1, bbox to anchor=(1.9,
0.5))
plt.xlabel(u'time interval')
plt.ylabel('reproduction rate')
# In[50]:
# Plotting the distribution of residuals
y1 = np.array(y)
y2 = np.array(y_pred_all)
y diff = y1[:, 0] - y2[:, 0]
sns.distplot(y diff, kde=False)
```

```
# In[51]:

# Confidence interval of regression coef
import numpy as np, statsmodels.api as sm
mod = sm.OLS(y_train, X_train)
res = mod.fit()
print (res.conf_int(0.01))
```