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```
library("gapminder")
library("ggplot2")
library("dplyr")
##
library("gam")
library("MASS")
data.gapminder <- data.frame(gapminder)</pre>
train.data <- data.gapminder %>% filter(year>=1952,year<=1992)</pre>
test.data <- data.gapminder %>% filter(year>=1993,year<=2007)
lm.fit<-lm(lifeExp~gdpPercap,train.data)</pre>
summary.lm<-summary(lm.fit)</pre>
summary(lm.fit)
##
## Call:
## lm(formula = lifeExp ~ gdpPercap, data = train.data)
##
## Residuals:
       Min
                10 Median
##
                                 3Q
                                        Max
## -76.914 -8.048 2.143
                             8.889 18.419
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) 5.284e+01 3.620e-01 145.98
                                               <2e-16 ***
                                               <2e-16 ***
## gdpPercap 7.233e-04 3.301e-05
                                      21.91
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 10.66 on 1276 degrees of freedom
## Multiple R-squared: 0.2734, Adjusted R-squared: 0.2728
## F-statistic: 480.1 on 1 and 1276 DF, p-value: < 2.2e-16
```

Summary of the model:

Formula:

In our case we have used **lifeExp** as a response variable and **gdpPercap** as predictor variables.

Residual Summary:

- One of the assumptions for linear models is that the errors follow a **normal distribution**, and as a consequence the residuals should as well.
- The residuals summary tells us about the **symmetry** of the residual distribution.
- As per the assumptions of the linear model our median should be close to 0 as the expected value of the residuals is 0.In our model the **median is not close to zero**.
- Further, the **first Quartile** and **third Quartile** should be close to each other in **magnitude**, as they would be under a symmetric zero mean distribution.
- The maximum and minimum residual should also have similar magnitude.
- However in our case **first quartile** and **third quartile** have similar magnitude but **minimum** and **maximum** residuals are far apart from each other in magnitude **violating symmetry** assumption.

Coefficients

Estimates:

- The estimate of the **intercept** gives us the expected response variable value when all the features are **zero**. In our case intercept gives us the **life expectancy** when the **gdpPercapita** is **zero**. i.e 5.284e+01 will be the life expectancy when the gdp per Capita is zero.
- If we increase the **gdpPercap** by **one unit**, there will be an increase of 7.233e-04 in response variable(lifeExpectancy).
- Positive estimate implies increase/decrease in predictor increases/decreases the response value and negative estimate implies increase/decrease in predictor decreases/increases the response value.
- If the estimated value of any variable is 0, it means that the corresponding predictor variable is not significant(not showing any impact in predicting the response variable value) to the model.

Standard Error:

- The **standard error** tells us the standard error of our estimate, which allows us to **construct marginal confidence intervals** for the estimate of that particular variable.
- If **S.E**(β_i) is the standard error and β_i is the estimated coefficient for a variable i, then we can construct confidence interval by $\beta_i \pm Z_{\alpha/2} * S.E(\beta_i)$.

t-value:

- The coefficient **t-value** tells us about how far our estimated parameter is from a hypothesized 0 value, scaled by the standard deviation of the estimate.
- We want it to be far away from zero as we could reject the null hypothesis and we could declare that a relationship between our predictor and response exists.
- t-values are also used to compute p-values.

Pr(>|t|):

- Under the t distribution with n-p-1 degrees of freedom, this tells us the **probability** of observing a value at least as extreme as our β_i , where n is no. of observations and p is no of predictor variables.
- If this probability is sufficiently low, we can reject the null hypothesis that this coefficient is 0 and declare that there exists a relationship between corresponding variable and the response variable.
- A **p-value of 0.05** or less is a good cut-off point.
- In our case the **intercept**, **gdpPercap** are having **p value very less than 0.05** and hence considered as significant to our model.
- The 'signif. Codes' associated with each estimate tells us the significance level of the respective variables. Three stars (or asterisks) represent a highly significant p-value and "" blank represents a least significant.

Residual Standard Error:

- It gives us the sum of squares of all the residuals.
- While fitting the model, the linear regression algorithm will try to adjust the beta's (coefficients) such that the RSE is as minimum as possible.

F statistic:

- The model with zero predictor variables is also called "Intercept Only Model".
- F Test for overall significance compares an **intercept only regression model** with our current model.
- It tries to check if the addition of these variables together into the model is significant enough or not .
- We can think of hypothesis test for this as:
- H0: The fit of the **intercept only model** and our **current model** is the same. i.e. Additional variables do not provide value taken together
- Ha: The fit of the **intercept only model** is significantly less compared to our **current model**. i.e. Additional variables do make the **model significantly better**.
- P Value of F Statistic 204.3 on 8 and 1021 DF, is < **2.2e-16** is extremely small, i.e smaller than 0.05 so we can reject **null hypothesis** and say that overall addition of variables is significantly improving the model.

Multiple R-squared, Adjusted R-squared:

- R² is a statistical measure that tells how well our model has fitted the data.
- It tells us the proportion of variance that our model was able to explain out of the total variance in the actual data.
- Mathematically, R² is given as:

$$R^2 = 1 - \frac{\sum_{i} (y_i - \hat{y}_i)^2}{\sum_{i} (y_i - \tilde{y})^2}$$

- It always lies between 0 and 1. A number near to 0 represents that the model did not explain the variance in the response variable well and a number close to 1 represents that the model was able to explain the total variance in the data.
- However, it's hard to define what level of (R²) is appropriate to claim the model fits well. It varies with the application and the domain studied. For example in an airline industry to fit a model for block time optimization with an R² of 0.4-0.5 is very good. Where as in rocket/satellite launching systems, an R² of 0.97-0.98 is also not a good fit
- In multiple regression, the (R²) will increase as more no of variables are included in the model. That's why the adjusted (R²) is the preferred measure as it adjusts for the number of variables considered.

• Expression for adjusted R² is given by :

$$R^{2} = 1 - \frac{\sum_{i}(y_{i} - \hat{y}_{i})^{2}/(n-p)}{\sum_{i}(y_{i} - \hat{y})^{2}/(n-1)}$$

where n is the number of samples and p is number of the predictor variables.

Testing of hypothesis:

- We can check the significance of any variable by looking at its p value.
- P-value basically tells us the probability of the predictor variable coefficient being zero.
- If the p-value is very less(generally checked if it is less than 0.05), we can say that there is very less chance of the coefficient being zero of the concerned variable and hence the variable is said to be significant(having impact in predicting response/output variable).

Linear Model:

In our case if we observe the summary of our linear fit, the p value of gdpPercap is <2e-16. It shows that the chance of coefficient of gdpPercap being zero is almost close to zero. Hence our hypothesis "lifeExp is dependent on gdpPercap" is correct.

```
glm.fit <- glm(lifeExp~gdpPercap,data=train.data)</pre>
summary.glm <- summary(glm.fit)</pre>
summary(glm.fit)
##
## Call:
## glm(formula = lifeExp ~ gdpPercap, data = train.data)
##
## Deviance Residuals:
      Min
                10 Median
                                  3Q
##
                                          Max
## -76.914 -8.048 2.143 8.889
                                       18.419
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 5.284e+01 3.620e-01 145.98
                                             <2e-16 ***
## gdpPercap 7.233e-04 3.301e-05
                                     21.91
                                             <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

```
## (Dispersion parameter for gaussian family taken to be 113.7217)

##
##
## Null deviance: 199710 on 1277 degrees of freedom
## Residual deviance: 145109 on 1276 degrees of freedom
## AIC: 9680.5
##
## Number of Fisher Scoring iterations: 2
```

Summary:

- The summary of the glm model also has similar terms like deviance residuals, coefficients (which has estimate, std. Error, t value, p value), significant codes.
- **Deviance** is a measure of goodness of fit of a generalized linear model.
- The **Saturated Model** is a model that assumes each data point has its own parameters (which means you have n parameters to estimate.)
- The **Null Model** assumes the exact "opposite", in that is assumes one parameter for all of the data points, which means you only estimate 1 parameter i.e, intercept..
- The **Proposed Model** assumes you can explain your data points with p parameters + an intercept term, so you have p+1 parameters.
- The **null deviance** shows how well the response variable is predicted by a model that includes only the intercept (grand mean).
- Akaike's information criterion (AIC) compares the quality of a set of statistical models to each other.
- The AIC will take each model and rank them from best to worst. The "best" model will be the one that neither under-fits nor over-fits.
- Akaike's Information Criterion is usually calculated with software. The basic formula is defined as:

AIC = -2(log-likelihood) + 2K

Where:

K is the number of model parameters (the number of variables in the model plus the intercept). Log-likelihood is a measure of model fit. The higher the number, the better the fit. This is usually obtained from statistical output.

Generalised Linear model:

By observing the summary of the generalised linear model fit the p value of gdpPercap is less than 2e-16 which says that the chance of coefficient of gdpPercap becoming zero is very less.**Hence our hypothesis "lifeExp is dependent on gdpPercap" is correct.**

```
loess.fit <- loess(lifeExp~gdpPercap,data=train.data)</pre>
summary.loess <- summary(loess.fit)</pre>
summary(loess.fit)
## Call:
## loess(formula = lifeExp ~ gdpPercap, data = train.data)
## Number of Observations: 1278
## Equivalent Number of Parameters: 5.5
## Residual Standard Error: 7.249
## Trace of smoother matrix: 6.03 (exact)
##
## Control settings:
    span : 0.75
##
##
    degree
##
   family : gaussian
##
    surface : interpolate
                                cell = 0.2
    normalize: TRUE
##
## parametric: FALSE
## drop.square: FALSE
```

Summary:

- Summary has no of observations, residual standard error(sum of squares of residuals), smoother matrix has the weights of the observations that are used in finding the smooth curve locally. Different control settings used to fit the loess regression locally.
- Loess regression doesn't produce coefficients,p-values so there's no way to test the hypothesis.
- The weights to observations change as x changes, so while there is an underlying simple equation that could be used to find the fitted value at any specified $x=x0^*$.
- That equation will be different for every such value of x you evaluate it. Loess computes local coefficients based on k-nearest neighbours (within a window).
- It generates more than one coefficient for every observation.
- So we cannot analyse the coefficients of gdpPercap and hence cannot tell the dependancy of lifeExp on gdpPercap.

```
gam.fit <- gam(lifeExp~gdpPercap,data=train.data)</pre>
## Warning in model.matrix.default(mt, mf, contrasts): non-list contrasts
argument
## ignored
summary.gam <- summary(gam.fit)</pre>
summary(gam.fit)
##
## Call: gam(formula = lifeExp ~ gdpPercap, data = train.data)
## Deviance Residuals:
##
       Min
                10 Median
                                3Q
                                       Max
## -76.914 -8.048
                     2.143
                             8.889 18.419
##
## (Dispersion Parameter for gaussian family taken to be 113.7217)
##
##
       Null Deviance: 199709.7 on 1277 degrees of freedom
## Residual Deviance: 145108.9 on 1276 degrees of freedom
## AIC: 9680.544
##
## Number of Local Scoring Iterations: 2
## Anova for Parametric Effects
               Df Sum Sq Mean Sq F value
                           54601 480.13 < 2.2e-16 ***
                1 54601
## gdpPercap
## Residuals 1276 145109
                             114
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Summary:

summary of gam fit has deviance residuals, null deviance, redidual deviance, AIC, no of iterations for the model to converge, coefficients which are similar to the above discussed terms in lm,glm models.

generalized additive model:

We can observe from the summary of the generalised additive model fit that the p value of gdpPercap is less than 2e-16 which says that the chance of coefficient of gdpPercap becoming zero is very less.

Hence our hypothesis "lifeExp is dependent on gdpPercap" is correct according to generalised additive model.

```
rlm.fit <- rlm(lifeExp~gdpPercap,data=train.data)</pre>
summary.rlm <- summary(rlm.fit)</pre>
summary(rlm.fit)
##
## Call: rlm(formula = lifeExp ~ gdpPercap, data = train.data)
## Residuals:
         Min
##
                    10
                          Median
                                         30
                                                  Max
## -129.2430 -7.1031
                          0.4585
                                     7.2206
                                              18.6475
##
## Coefficients:
##
               Value
                        Std. Error t value
## (Intercept) 50.4612
                          0.3171
                                    159.1503
## gdpPercap
                 0.0012
                          0.0000
                                     41.6799
##
## Residual standard error: 10.69 on 1276 degrees of freedom
#first argument is t value, second argument is degrees of freedom, lower.tail
# logical; if TRUE (default), probabilities are P[X \le x], otherwise, P[X > x]
pValue <- 2*pt(summary.rlm$coefficients[6],summary.rlm$df[2],lower.tail =</pre>
FALSE)
cat("pValue of gdpPerCap is :",pValue)
## pValue of gdpPerCap is : 2.399971e-240
```

Summary:

summary of rlm model is same as lm model as rlm is modification to lm model which gives less weights to the outliers.

robust linear model:

As the summary of the rlm fit does not include p values, we can find p values from the t value using pt(t-value,degreesOfFreedom) function.

We can observe p value of gdpPercap is 2.399971e-240 which says that the chance of coefficient of gdpPercap becoming zero is very less.

Hence our hypothesis "lifeExp is dependent on gdpPercap" is correct according to generalised additive model.

Question2: Use the data from summary and plot a histogram to show how well does your model fits the data.

```
lm.residuals <- data.frame(residuals = residuals(lm.fit))
ggplot(data=lm.residuals,mapping=aes(x=residuals))+
    geom_histogram(binwidth=1,aes(y = ..density..), alpha =
0.8,fill="skyblue")+
    labs(title="plot for residuals of train data from linear model")+
    theme(plot.title = element_text(hjust=0.5))+
    geom_vline(data=lm.residuals, aes(xintercept = 0), colour="red")+
    geom_density()</pre>
```

plot for residuals of train data from linear model

-50

-75

• Goodness of fit of a model can be checked by plotting histogram plot of residuals. If the residuals are following normal distribution then we can say that our model is fitting the data properly.

0

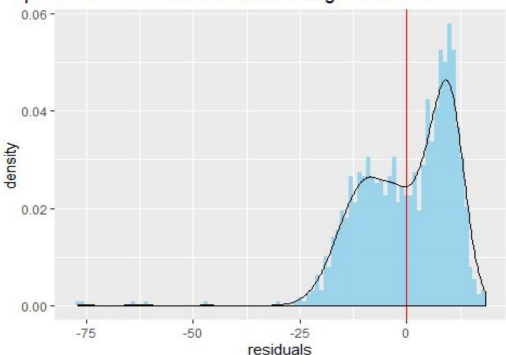
-25

residuals

• We can observe from the histogram of residuals for linear model that the residuals are not following the normal distribution(we can clearly see its a bimodal distribution). This says that the model is not fitting the data properly.

```
glm.residuals <- data.frame(residuals = residuals(glm.fit))
ggplot(data=glm.residuals,mapping=aes(x=residuals))+
    geom_histogram(binwidth=1,aes(y=..density..),fill="skyblue",alpha=0.8)+
    labs(title="plot for residuals of train data from generalised linear
model")+
    theme(plot.title = element_text(hjust=0.5))+
    geom_vline(data=glm.residuals, aes(xintercept = 0), colour="red")+
    geom_density()</pre>
```

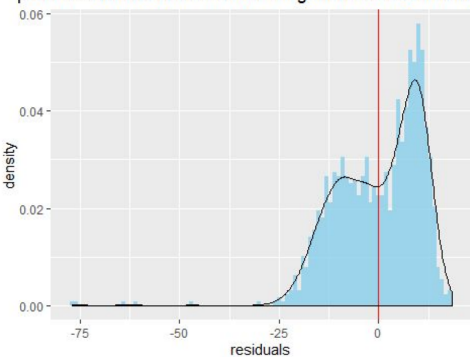
plot for residuals of train data from generalised linear mod



- We can observe from the histogram of residuals for generalised linear model that the residuals are not following the normal distribution. This says that the model is not fitting the data properly.
- We can also see that there are some outliers towards the left in the histogram which has very high residual value.

```
gam.residuals <- data.frame(residuals = residuals(gam.fit))
ggplot(data=gam.residuals, mapping=aes(x=residuals)) +
    geom_histogram(binwidth=1,aes(y = ..density..), alpha =
0.8,fill="skyblue")+
    labs(title="plot for residuals of train data from generative additive
model")+
    theme(plot.title = element_text(hjust=0.5))+
    geom_vline(data=gam.residuals, aes(xintercept = 0), colour="red")+
    geom_density()</pre>
```

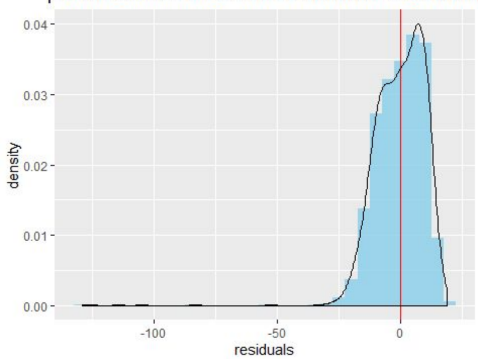
plot for residuals of train data from generative additive mo



- As generative additive model is obtained by combining more than one glm model the residuals of this model is also coming out to be following the similar distribution.
- We can observe from the histogram of residuals for generalised additive model that the residuals are not following the normal distribution. This says that the model is not fitting the data properly.

```
rlm.residuals <- data.frame(residuals = residuals(rlm.fit))
ggplot(data=rlm.residuals,mapping=aes(x=residuals)) +
    geom_histogram(binwidth=5,aes(y = ..density..), alpha =
0.8,fill="skyblue")+
    labs(title="plot for residuals of train data from robust linear model")+
    theme(plot.title = element_text(hjust=0.5))+
    geom_vline(data=rlm.residuals, aes(xintercept = 0), colour="red")+
    geom_density()</pre>
```

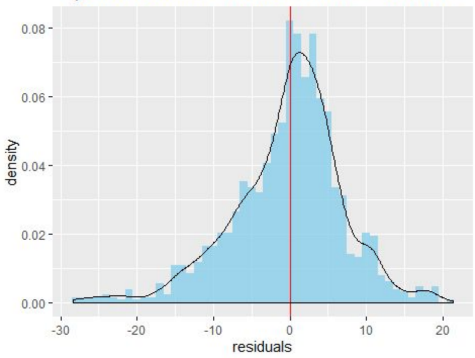
plot for residuals of train data from robust linear model



- We can observe that the residuals of robust linear are following normal distribution with some outliers lying towards the left in the histogram plot.
- Hence we can say that robust linear model is properly fitting our data.

```
loess.residuals <- data.frame(residuals = residuals(loess.fit))
ggplot(data=loess.residuals,mapping=aes(x=residuals)) +
    geom_histogram(binwidth=1,aes(y = ..density..), alpha =
0.8,fill="skyblue")+
    labs(title="plot for residuals of train data from loess model")+
    theme(plot.title = element_text(hjust=0.5))+
    geom_vline(data=loess.residuals, aes(xintercept = 0), colour="red")+
    geom_density()</pre>
```

plot for residuals of train data from loess model

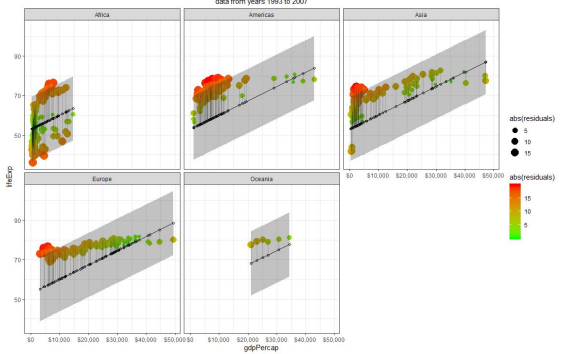


- We can observe that the residuals are lying symmetric the mean 0 which is expected. and also We can observe that the residuals of loess model are following normal distribution.
- Hence we can say that loess is perfectly fitting our data.
- Out of all the model loess is fitting the data perfectly but loess is not preferred for the datasets greater than 1000 samples as it is computationally not storable. It requires $O(n^2)$ in memory.

Question3: Plot a graph showing the prediction on test data points, linear regression line fitting the data. Also draw the area covered by prediction intervals. Facet the data based on continents.

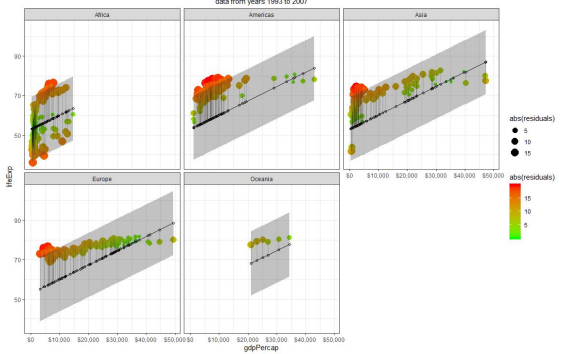
```
temp <- predict(lm.fit,test.data,interval = "predict")</pre>
test.data$predicted <- temp[,1]</pre>
test.data$residuals <- test.data$lifeExp-test.data$predicted
sd1 <- sd(test.data$predicted)</pre>
ggplot(test.data,mapping=aes(x=gdpPercap,y=lifeExp))+
geom_point(test.data,mapping=aes(x=gdpPercap,y=lifeExp,color=abs(residuals),s
ize=abs(residuals)))+
  geom_point(mapping=aes(y=predicted), shape=1)+
  geom line(mapping=aes(y=predicted))+
  scale_color_continuous(low="green", high="red")+
  geom_segment(aes(xend=gdpPercap,yend=predicted),alpha=.2)+
  theme bw()+
  facet_wrap(continent~.)+
geom_ribbon(mapping=aes(ymin=test.data$predicted-1.96*sd1,ymax=test.data$pred
icted+1.96*sd1), alpha = 0.3)+
  labs(title="plot for actual vs predicted values of lifeExpectancy of
different continents using lm", subtitle = "data from years 1993 to 2007")+
  theme(plot.title = element_text(hjust=0.5),plot.subtitle =
element text(hjust=0.5))+
  scale_x_continuous(labels = scales::dollar)
```

plot for actual vs predicted values of lifeExpectancy of different continents using Im data from years 1993 to 2007



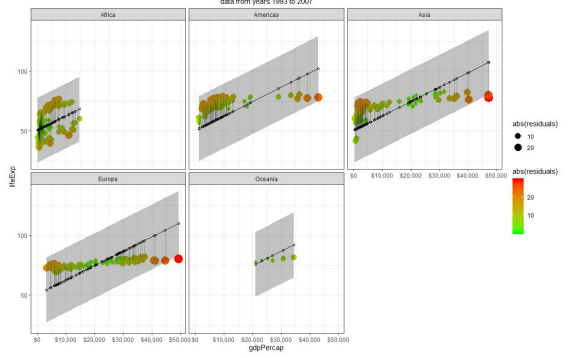
```
test.data$predicted <- predict(glm.fit,test.data)</pre>
test.data$residuals <- test.data$lifeExp-test.data$predicted
sd2 <- sd(test.data$predicted)</pre>
ggplot(test.data,mapping=aes(x=gdpPercap,y=lifeExp))+
geom_point(test.data,mapping=aes(x=gdpPercap,y=lifeExp,color=abs(residuals),s
ize=abs(residuals)))+
  geom_point(mapping=aes(y=predicted), shape=1)+
  geom_line(mapping=aes(y=predicted))+
  scale_color_continuous(low="green",high="red")+
  geom_segment(aes(xend=gdpPercap,yend=predicted),alpha=.2)+
  theme bw()+
  facet_wrap(continent~.)+
geom_ribbon(mapping=aes(ymin=test.data$predicted-1.96*sd2,ymax=test.data$pred
icted+1.96*sd2), alpha = 0.3)+
  labs(title="plot for actual vs predicted values of lifeExpectancy of
different continents using glm", subtitle = "data from years 1993 to 2007")+
  theme(plot.title = element_text(hjust=0.5),plot.subtitle =
element_text(hjust=0.5))+
  scale_x_continuous(labels = scales::dollar)
```

plot for actual vs predicted values of lifeExpectancy of different continents using glm data from years 1993 to 2007



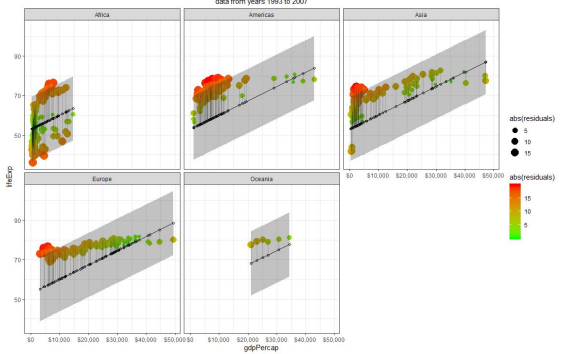
```
test.data$predicted <- predict(rlm.fit,test.data)</pre>
test.data$residuals <- test.data$lifeExp-test.data$predicted
sd3 <- sd(test.data$predicted)</pre>
ggplot(test.data,mapping=aes(x=gdpPercap,y=lifeExp))+
geom_point(test.data,mapping=aes(x=gdpPercap,y=lifeExp,color=abs(residuals),s
ize=abs(residuals)))+
  geom_point(mapping=aes(y=predicted), shape=1)+
  geom_line(mapping=aes(y=predicted))+
  scale_color_continuous(low="green",high="red")+
  geom_segment(aes(xend=gdpPercap,yend=predicted),alpha=.2)+
  theme bw()+
  facet_wrap(continent~.)+
geom_ribbon(mapping=aes(ymin=test.data$predicted-1.96*sd3,ymax=test.data$pred
icted+1.96*sd3), alpha = 0.3)+
  labs(title="plot for actual vs predicted values of lifeExpectancy of
different continents using rlm", subtitle = "data from years 1993 to 2007")+
  theme(plot.title = element_text(hjust=0.5),plot.subtitle =
element_text(hjust=0.5))+
  scale_x_continuous(labels = scales::dollar)
```

plot for actual vs predicted values of lifeExpectancy of different continents using rlm data from years 1993 to 2007



```
test.data$predicted <- predict(gam.fit,test.data)</pre>
test.data$residuals <- test.data$lifeExp-test.data$predicted
sd4<-sd(test.data$predicted)</pre>
ggplot(test.data,mapping=aes(x=gdpPercap,y=lifeExp))+
geom_point(test.data,mapping=aes(x=gdpPercap,y=lifeExp,color=abs(residuals),s
ize=abs(residuals)))+
  geom_point(mapping=aes(y=predicted), shape=1)+
  geom_line(mapping=aes(y=predicted))+
  scale_color_continuous(low="green",high="red")+
  geom_segment(aes(xend=gdpPercap,yend=predicted),alpha=.2)+
geom_ribbon(mapping=aes(ymin=test.data$predicted-1.96*sd4,ymax=test.data$pred
icted+1.96*sd4), alpha = 0.3)+
  theme bw()+
  facet_wrap(continent~.)+
  labs(title="plot for actual vs predicted values of lifeExpectancy of
different continents using gam", subtitle = "data from years 1993 to 2007")+
  theme(plot.title = element_text(hjust=0.5),plot.subtitle =
element_text(hjust=0.5))+
  scale_x_continuous(labels = scales::dollar)
```

plot for actual vs predicted values of lifeExpectancy of different continents using gam data from years 1993 to 2007



```
test.data$predicted <- predict(loess.fit,test.data)</pre>
test.data$residuals <- test.data$lifeExp-test.data$predicted
test.data <- test.data[complete.cases(test.data),]</pre>
sd5<-sd(test.data$predicted)</pre>
ggplot(test.data,mapping=aes(x=gdpPercap,y=lifeExp))+
geom_point(test.data,mapping=aes(x=gdpPercap,y=lifeExp,color=abs(residuals),s
ize=abs(residuals)))+
  geom_point(mapping=aes(y=predicted), shape=1)+
  geom_line(mapping=aes(y=predicted))+
  scale_color_continuous(low="green", high="red")+
  geom_segment(aes(xend=gdpPercap,yend=predicted),alpha=.2)+
geom_ribbon(mapping=aes(ymin=test.data$predicted-1.96*sd5,ymax=test.data$pred
icted+1.96*sd5),alpha=0.3)+
  theme_bw()+
  facet_wrap(continent~.)+
  labs(title="plot for actual vs predicted values of lifeExpectancy of
different continents using loess", subtitle = "data from years 1993 to 2007")+
  theme(plot.title = element_text(hjust=0.5),plot.subtitle =
element text(hjust=0.5))+
  scale_x_continuous(labels = scales::dollar)
```

plot for actual vs predicted values of lifeExpectancy of different continents using loess data from years 1993 to 2007

