# Generalized Additive Models (GAMs) CMDA 4654 Project 2

Group 19: Brady Bolton, Eryk Jesse, Charles Lee, Dan Schlicht



Replying to @millerdl

140 char vrsn

1 GAMs are just GLMs
2 GAMs fit wiggly terms
3 use + s(foo) not foo in frmla
4 use method = "REML"
5 gam.check()

2:37 PM · Mar 16, 2017 · TweetDeck

- Type of generalized linear model
- Response variable depends on smooth functions  $f_i(x_i)$
- General structure of a GAM:  $g(E(Y)) = \beta_0 + f_1(x_1) + f_2(x_2) + \cdots + f_n(x_n)$
- Smooth functions can be many different things (polynomials, splines, weighted means, etc)

The main package used for GAMs is mgcv.

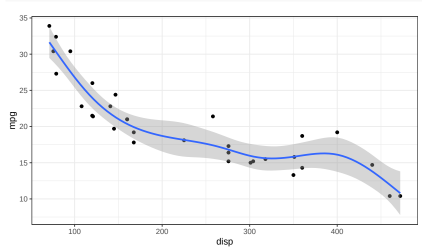
Mixed GAM Computation Vehicle with Automatic

**Smoothness Estimation** 

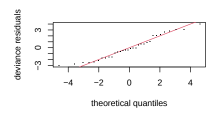


```
library(mgcv)
data("mtcars")
mtcars_gam <-
  gam(mpg ~ s(disp), data = mtcars, method = "REML")
summary(mtcars_gam)
Family: gaussian
Link function: identity
Formula:
mpg ~ s(disp)
Parametric coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 20.0906 0.3788 53.04 <2e-16 ***
---
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
Approximate significance of smooth terms:
         edf Ref.df F p-value
s(disp) 4.884 5.904 36.3 <2e-16 ***
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
R-sq.(adi) = 0.874 Deviance explained = 89.4%
-REML = 74.101 Scale est. = 4.5918 n = 32
```

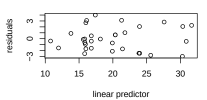
```
library(ggplot2)
ggplot(data = mtcars, aes(x = disp, y = mpg)) +
    theme_bw() + geom_point() +
    geom_smooth(method = "gam", formula = y ~ s(x))
```



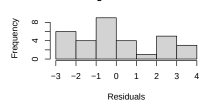
gam.check(mtcars\_gam)



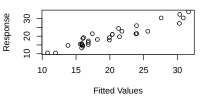
#### Resids vs. linear pred.



Histogram of residuals



#### Response vs. Fitted Values



```
data("iris")
```

```
# df <- read.csv('health_data.csv')
# show the data</pre>
```

### **Conclusion**

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### References