

# GAM Fits for Hirsutism Data

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This document performs Generalized Additive Model (GAM) analysis on hirsutism clinical trial data. The goal is to model the Ferriman-Gallwey score at 12 months (`FGm12`) as a function of baseline measurements and treatment levels.

## 1. Data Loading and Cleaning

We begin by loading the hirsutism dataset and preparing it for analysis. This includes converting the treatment variable to a factor and handling missing or erroneous values.

```
library(mgcv)

# Load the dataset.
hirs <- read.table("hirsutism.dat", header = TRUE, sep = "\t", fill = TRUE)

# Convert Treatment to factor.
hirs$Treatment <- as.factor(hirs$Treatment)

# Correct erroneous negative FGm12 values.
neg_idx <- which(hirs$FGm12 < 0)
hirs$FGm12[neg_idx] <- 0

# Remove observations with missing values.
hirs <- na.omit(hirs)
```

Table 1: Treatment Group Counts

Value	Frequency
0	22
1	22
2	22
3	25

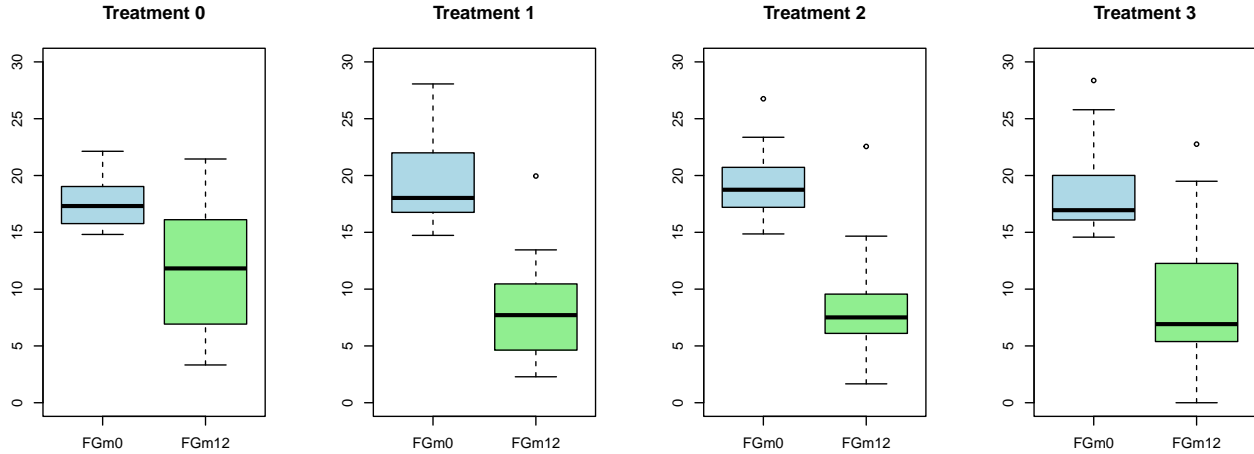
Table 2: Summary Statistics of Key Variables

FGm0	FGm12	SysPres	DiaPres	weight	height
Min. :14.57	Min. : 0.000	Min. : 88.0	Min. :46.00	Min. : 41.00	Min. :1.480
1st Qu.:16.40	1st Qu.: 5.566	1st Qu.:110.0	1st Qu.:65.00	1st Qu.: 57.00	1st Qu.:1.580
Median :17.70	Median : 8.069	Median :115.0	Median :70.00	Median : 64.00	Median :1.610
Mean :18.67	Mean : 9.066	Mean :115.9	Mean :70.04	Mean : 68.06	Mean :1.613
3rd Qu.:20.27	3rd Qu.:12.402	3rd Qu.:120.0	3rd Qu.:75.00	3rd Qu.: 74.50	3rd Qu.:1.650

FGm0	FGm12	SysPres	DiaPres	weight	height
Max. :28.36	Max. :22.759	Max. :162.0	Max. :95.00	Max. :113.00	Max. :1.800

## 2. Exploratory Data Analysis

Here, we explore the relationships between predictors and the response variable through visualizations and correlation analysis.



```
# Compute correlations between numeric predictors and FGm12.
numeric_vars <- hirs[, c("FGm0", "SysPres", "DiaPres", "weight", "height", "FGm12")]
cor_with_target <- cor(numeric_vars, method = "pearson")[, "FGm12"]
```

Table 3: Pearson Correlations with FGm12

Variable	Correlation
FGm0	0.308
SysPres	-0.177
DiaPres	-0.079
weight	0.000
height	-0.057

The correlations are relatively low, suggesting that linear relationships alone may not capture the underlying patterns. This motivates the use of GAMs with smooth terms.

## 3. Model Fitting

We fit several GAM models of increasing complexity, starting from a simple linear model and progressively adding smooth terms and interactions.

```
set.seed(123)

# Model 0: Linear model (baseline).
gam0 <- gam(FGm12 ~ FGm0 + SysPres + DiaPres + weight + height + Treatment,
             data = hirs)
```

```

# Model 1: Full additive model with smooth terms.
gam1 <- gam(FGm12 ~ s(FGm0) + s(SysPres) + s(DiaPres) + s(weight) + s(height) + Treatment,
             data = hirs)

# Model 2: Treatment-specific smooths for FGm0.
gam2 <- gam(FGm12 ~ s(FGm0, by = Treatment) + s(SysPres) + s(DiaPres) + s(weight) + s(height) + Treatment,
             data = hirs)

# Model 3: Reduced model (remove non-significant DiaPres).
gam3 <- gam(FGm12 ~ s(FGm0, by = Treatment) + s(SysPres) + s(weight) + s(height) + Treatment,
             data = hirs)

# Model 4: Further reduction (remove height).
gam4 <- gam(FGm12 ~ s(FGm0, by = Treatment) + s(SysPres) + s(weight) + Treatment,
             data = hirs)

# Model 5: Minimal smooth model.
gam5 <- gam(FGm12 ~ s(FGm0, by = Treatment) + s(SysPres) + Treatment,
             data = hirs)

# Model 6: Only FGm0 smooth by Treatment.
gam6 <- gam(FGm12 ~ s(FGm0, by = Treatment) + Treatment,
             data = hirs)

# Model 7: Tensor product for DiaPres and weight.
gam7 <- gam(FGm12 ~ s(FGm0, by = Treatment) + s(SysPres) + te(DiaPres, weight) + Treatment,
             data = hirs)

# Model 8: Two tensor products (best candidate).
gam8 <- gam(FGm12 ~ s(FGm0, by = Treatment) + te(DiaPres, weight) + te(SysPres, height) + Treatment,
             data = hirs)

```

## 4. Model Selection

We compare models using GCV scores and ANOVA to identify the best-fitting model.

```

# Extract GCV scores for comparison.
gcv_scores <- c(
  Linear = gam0$gcv.ubre,
  GAM1 = gam1$gcv.ubre,
  GAM2 = gam2$gcv.ubre,
  GAM3 = gam3$gcv.ubre,
  GAM4 = gam4$gcv.ubre,
  GAM5 = gam5$gcv.ubre,
  GAM6 = gam6$gcv.ubre,
  GAM7 = gam7$gcv.ubre,
  GAM8 = gam8$gcv.ubre
)

```

Table 4: GCV Scores Across Models (Lower is Better)

Model	GCV
Linear	24.950
GAM1	23.146
GAM2	22.563
GAM3	22.562
GAM4	22.086
GAM5	22.815
GAM6	23.021
GAM7	21.608
GAM8	20.922

```
# ANOVA comparison of nested models.
anova(gam5, gam7, gam8, gam4, gam3, gam2, gam0, test = "F")
```

```
## Analysis of Deviance Table
##
## Model 1: FGm12 ~ s(FGm0, by = Treatment) + s(SysPres) + Treatment
## Model 2: FGm12 ~ s(FGm0, by = Treatment) + s(SysPres) + te(DiaPres, weight) +
##     Treatment
## Model 3: FGm12 ~ s(FGm0, by = Treatment) + te(DiaPres, weight) + te(SysPres,
##     height) + Treatment
## Model 4: FGm12 ~ s(FGm0, by = Treatment) + s(SysPres) + s(weight) + Treatment
## Model 5: FGm12 ~ s(FGm0, by = Treatment) + s(SysPres) + s(weight) + s(height) +
##     Treatment
## Model 6: FGm12 ~ s(FGm0, by = Treatment) + s(SysPres) + s(DiaPres) + s(weight) +
##     s(height) + Treatment
## Model 7: FGm12 ~ FGm0 + SysPres + DiaPres + weight + height + Treatment
##   Resid. Df Resid. Dev      Df Deviance      F    Pr(>F)
## 1    71.641   1378.00
## 2    57.537   904.26  14.1039   473.74 2.7106 0.0049831 **
## 3    48.856   667.87  8.6818   236.39 2.1973 0.0399574 *
## 4    66.245   1175.31 -17.3892  -507.44 2.3549 0.0097072 **
## 5    67.934   1245.75 -1.6895   -70.44 3.3647 0.0503650 .
## 6    63.890   1122.11  4.0448   123.64 2.4669 0.0564517 .
## 7    82.000   1843.55 -18.1105  -721.44 3.2147 0.0005985 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

The ANOVA table reveals significant improvements when moving from simpler to more complex models. Notably, the transition from GAM5 to GAM7 (adding `te(DiaPres, weight)`) and from GAM7 to GAM8 (adding `te(SysPres, height)`) both show significant F-statistics ( $p < 0.05$ ). The comparison also confirms that the linear model (`gam0`) is significantly outperformed by the GAM alternatives ( $p < 0.001$ ).

Based on the lowest GCV score (20.922) and ANOVA results, **Model 8** (`gam8`) is selected as the best model. It achieves 72.6% deviance explained through treatment-specific smooths for `FGm0` and tensor product interactions.

## 5. Final Model Examination

We examine the selected model in detail using summary statistics, smooth plots, and diagnostic checks.

```

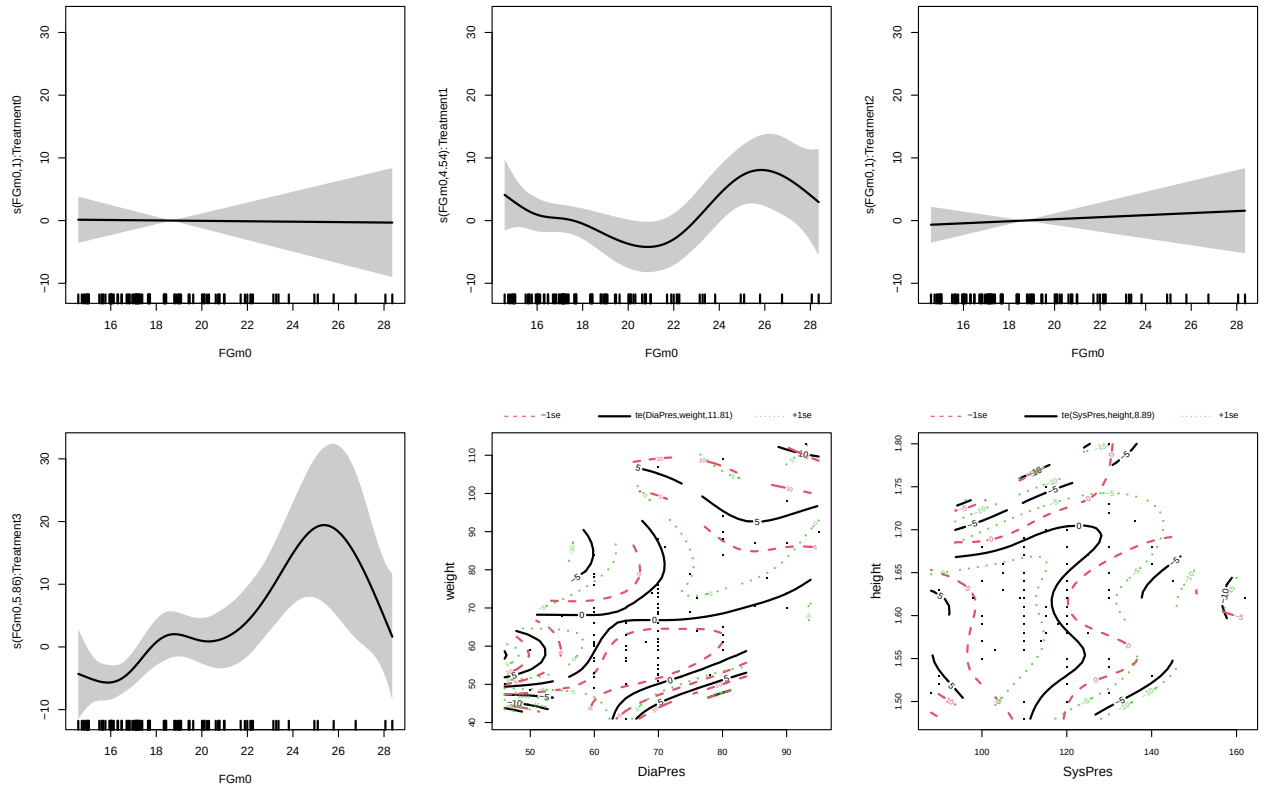
summary(gam8)

##
## Family: gaussian
## Link function: identity
##
## Formula:
## FGm12 ~ s(FGm0, by = Treatment) + te(DiaPres, weight) + te(SysPres,
##      height) + Treatment
##
## Parametric coefficients:
##             Estimate Std. Error t value Pr(>|t|)
## (Intercept) 11.1839    0.9629 11.615 2.71e-16 ***
## Treatment1 -3.4750    1.3632 -2.549  0.0137 *
## Treatment2 -1.8591    1.3216 -1.407  0.1653
## Treatment3 -2.8704    1.3489 -2.128  0.0379 *
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Approximate significance of smooth terms:
##                  edf Ref.df     F p-value
## s(FGm0):Treatment0 1.000 1.000 0.006 0.941025
## s(FGm0):Treatment1 4.540 5.395 2.495 0.036942 *
## s(FGm0):Treatment2 1.000 1.000 0.215 0.644806
## s(FGm0):Treatment3 5.864 6.824 4.834 0.000398 ***
## te(DiaPres,weight) 11.814 13.226 2.293 0.016420 *
## te(SysPres,height) 8.885 10.700 1.977 0.056693 .
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## R-sq.(adj) =  0.542 Deviance explained = 72.6%
## GCV = 20.922 Scale est. = 12.392 n = 91

```

The model summary reveals:

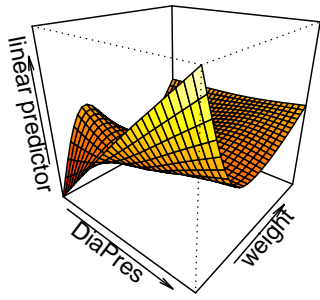
- **Parametric coefficients:** Treatment1 ( $p = 0.014$ ) and Treatment3 ( $p = 0.038$ ) show significant reductions in FGm12 scores relative to the control group. Treatment2 does not reach statistical significance ( $p = 0.165$ ).
- **Smooth terms:** `s(FGm0):Treatment1` and `s(FGm0):Treatment3` are significant, indicating nonlinear baseline effects for these treatment groups. The tensor product `te(DiaPres, weight)` is significant ( $p = 0.016$ ), while `te(SysPres, height)` is marginally significant ( $p = 0.057$ ).



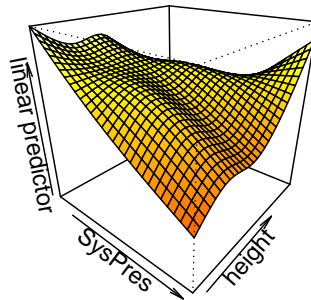
The smooth plots reveal:

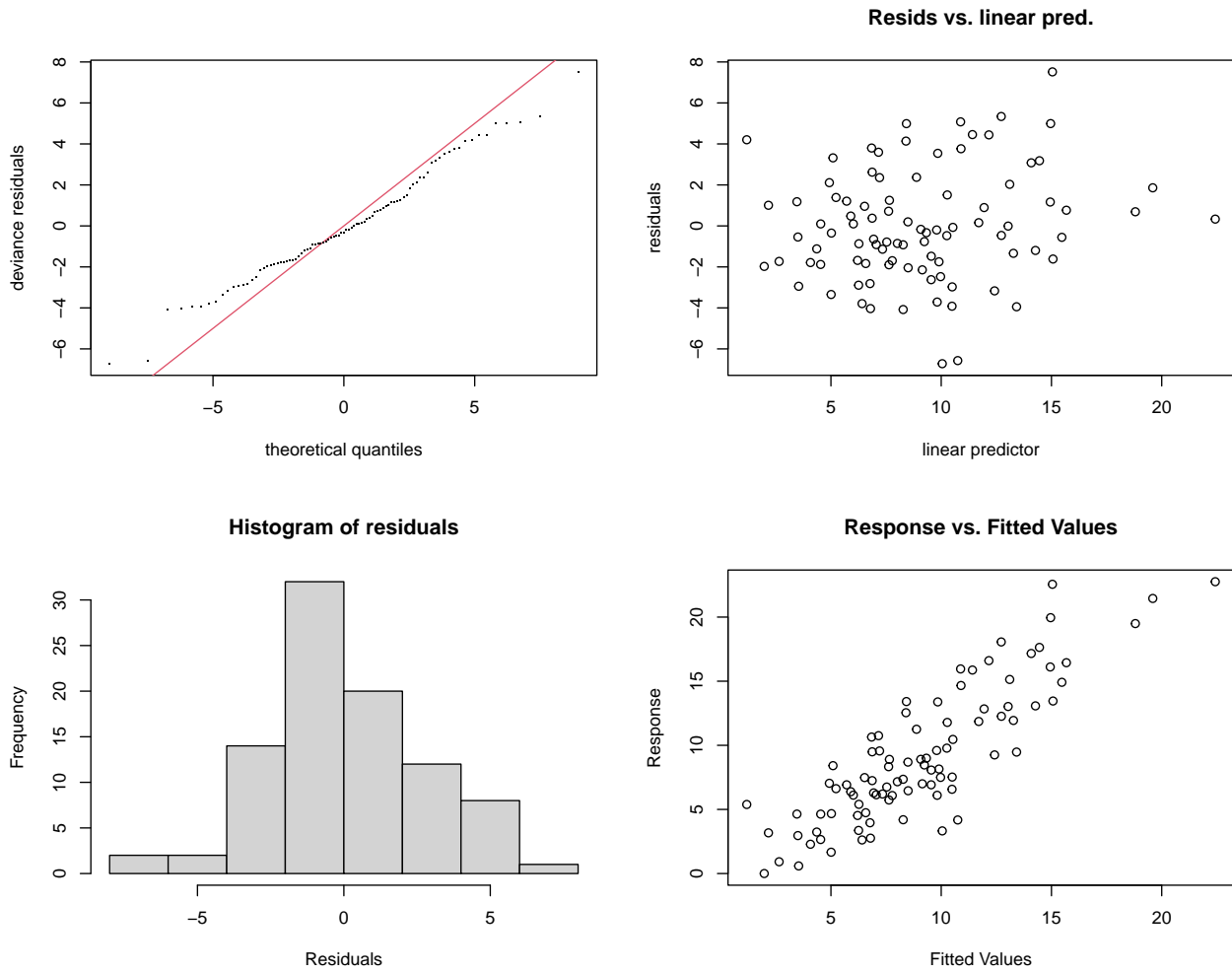
- **Treatment 0 and 2:** Flat effects of  $FGm0$  ( $edf \approx 1$ ), indicating essentially linear or no relationship between baseline hirsutism and final outcome for these groups.
- **Treatment 1:** A nonlinear pattern ( $edf = 4.54$ ) with higher  $FGm12$  values at elevated baseline levels.
- **Treatment 3:** The most complex relationship ( $edf = 5.86$ ) showing strong nonlinear dependence on baseline severity.
- **Tensor products:** Capture complex interactions, with  $te(DiaPres, weight)$  using 11.8 effective degrees of freedom.

### Interaction: DiaPres x Weight



### Interaction: SysPres x Height





```

## 
## Method: GCV   Optimizer: magic
## Smoothing parameter selection converged after 18 iterations.
## The RMS GCV score gradient at convergence was 1.072644e-06 .
## The Hessian was positive definite.
## Model rank =  88 / 88
## 
## Basis dimension (k) checking results. Low p-value (k-index<1) may
## indicate that k is too low, especially if edf is close to k'.
## 
##          k'    edf k-index p-value
## s(FGm0):Treatment0 9.00  1.00    1.05    0.66
## s(FGm0):Treatment1 9.00  4.54    1.05    0.63
## s(FGm0):Treatment2 9.00  1.00    1.05    0.69
## s(FGm0):Treatment3 9.00  5.86    1.05    0.66
## te(DiaPres,weight) 24.00 11.81    1.09    0.88
## te(SysPres,height) 24.00  8.89    0.86    0.03 *
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

```

The diagnostic plots indicate:

- **Q-Q plot:** Residuals follow theoretical quantiles reasonably well, with minor deviations at the tails.

- **Residuals vs. fitted:** No systematic patterns, suggesting homoscedasticity.
- **Histogram:** Approximately normal distribution of residuals.
- **Response vs. fitted:** Positive correlation between observed and predicted values.

The basis dimension check shows adequate k values for most smooth terms ( $k\text{-index} > 1$ ). However, `te(SysPres, height)` displays a k-index of 0.86 with  $p = 0.03$ , suggesting potential under-smoothing. Since the effective degrees of freedom ( $\text{edf} = 8.89$ ) is well below the basis dimension ( $k' = 24$ ), this is likely a minor concern and no corrective action is required.

## 6. Concluding Remarks

The Generalized Additive Model analysis demonstrates that smooth terms and tensor product interactions substantially improve model fit compared to a simple linear model. The final model (`gam8`) achieves the lowest GCV score (20.922) and explains 72.6% of the variability in `FGm12` by:

- Modeling treatment-specific nonlinear effects of baseline hirsutism levels (`FGm0`).
- Capturing interactions between physiological variables (`DiaPres × weight` and `SysPres × height`) through tensor product smooths.

Key findings from the clinical perspective:

- **Treatment efficacy varies:** Treatments 1 and 3 show statistically significant reductions in Ferriman-Gallwey scores relative to the control group, while Treatment 2 does not reach significance.
- **Baseline severity matters differently by treatment:** For Treatments 0 and 2, baseline hirsutism has minimal impact on final outcomes. For Treatments 1 and 3, participants with higher baseline FG values show stronger nonlinear responses, suggesting these treatments may be more effective for patients with moderate baseline severity.
- **Physiological interactions:** The significant interaction between diastolic blood pressure and weight suggests that body composition and cardiovascular factors may influence treatment response, warranting further clinical investigation.