### 6: Generalized Additive Models

#### John H Maindonald

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### Ideas and issues illustrated by the graphs in this vignette

Generalized Additive Models (GAMs) extend linear and generalized linear models to include smooth functions of explanatory variables, where the smoothness may be determined automatically. The graphs shown here illustrate some of the possibilities.

**Note:** Figure 6.9 shows the results of simulations. The version of this figure that is shown in Section 2 is, in order to keep to a minimum the time taken to process the vignette, for 25 simulations only. This is useful mainly as a check that the code does what is expected of it. More realistically, specify 500 or 1000 (as in the text) simulations.

# 1 Code for the Figures

```
fig6.1 <- function(plotit=TRUE){</pre>
    matchms <- data.frame(model.matrix(with(fruitchms, ~ poly(juice, 4))))</pre>
    names(matchms) <- c("Intercept", paste("poly4",1:4, sep=""))</pre>
    form <- formula(paste(paste(names(matchms), collapse="+"), "~ juice"))</pre>
    matohms$juice <- fruitohms$juice</pre>
    gph1 <- xyplot(form, data=matchms, layout=c(1,5), scales=list(tck=0.5),</pre>
                    ylab="Basis terms",
                    strip=strip.custom(strip.names=TRUE,
                    var.name="",
                    sep=expression(""),
                    factor.levels=c("Constant","Linear","Quadratic",
                                     "Cubic", "Quartic")),
                    panel=function(x,y,...){
                        llines(smooth.spline(x,y))},
                    outer=TRUE,
                    legend=list(top=list(fun=grid::textGrob,
                                 args=list(label="A: Basis functions",
                                 just="left", x=0))))
```

```
b <- coef(lm(I(ohms/1000) ~ poly(juice,4), data=fruitohms))
    matchms <- sweep(model.matrix(with(fruitchms, ~ poly(juice, 4))),</pre>
                     2, b, "*")
    matchms <- data.frame(matchms)</pre>
    names(matchms) <- c("Intercept", paste("poly4",1:4, sep=""))</pre>
    form <- formula(paste(paste(names(matchms), collapse="+"), "~ juice"))</pre>
    matohms$juice <- fruitohms$juice</pre>
    matohms$Kohms <- fruitohms$ohms/1000</pre>
    nam <- lapply(1:5, function(x)substitute(A %*% B,
                                               list(A=round(b[x],2),
                                                    B=c("Constant","Linear",
                                                    "Quadratic", "Cubic",
                                                    "Quartic")[x])))
    gph2 <- xyplot(form, data=matchms, layout=c(1,5),, scales=list(tck=0.5),</pre>
                    ylab="Add the contributions from these curves",
                    strip=strip.custom(strip.names=TRUE,
                    var.name="",
                    sep=expression(""),
                    factor.levels=as.expression(nam)),
                    panel=function(x,y,...){
                        llines(smooth.spline(x,y))},
                    outer=TRUE,
                    legend=list(top=list(fun=grid::textGrob,
                          args=list(label="B: Contribution to fitted curve",
                                    just="left", x=0))))
    if(plotit){
        print(gph1, position=c(0,0,.5,1))
        print(gph2, position=c(.5,0,1,1), newpage=FALSE)
    invisible(list(gph1, gph2))
fig6.2 <- function(){</pre>
    plot(ohms ~ juice, data=fruitohms, ylim=c(0, max(ohms)*1.02))
    ## 3 (=2+1) degrees of freedom natural spline
    fitns2 <- fitted(lm(ohms ~ splines::ns(juice, df=2), data=fruitohms))</pre>
    lines(fitns2 ~ juice, data=fruitohms, col="gray40")
    ## 4 (=3+1) degrees of freedom natural spline
    fitns3 <- fitted(lm(ohms ~ splines::ns(juice, df=3), data=fruitohms))</pre>
    lines(fitns3 ~ juice, data=fruitohms, lty=2, lwd=2, col="gray40")
    legend("topright", title="D.f. for cubic regression natural spline",
           legend=c("3 [ns(juice, 2)]",
           "4 [ns(juice, 3)]"),
           lty=c(1,2), lwd=c(1,2), cex=0.8
```

```
plot(ohms ~ juice, data=fruitohms, ylim=c(0, max(ohms)*1.02))
    ## 3 (=2+1) degrees of freedom natural spline
    fitns2 <- fitted(lm(ohms ~ splines::ns(juice, df=2), data=fruitohms))</pre>
    lines(fitns2 ~ juice, data=fruitohms, col="gray40")
    ## 4 (=3+1) degrees of freedom natural spline
    fitns3 <- fitted(lm(ohms ~ splines::ns(juice, df=3), data=fruitohms))</pre>
    lines(fitns3 ~ juice, data=fruitohms, lty=2, lwd=2, col="gray40")
    legend("topright", title="D.f. for cubic regression natural spline",
           legend=c("3 [ns(juice, 2)]",
           "4 [ns(juice, 3)]"),
           lty=c(1,2), lwd=c(1,2), cex=0.8)
fig6.3 <- function(){</pre>
    ohms.lm <- lm(ohms ~ ns(juice, df=3), data=fruitohms)
    termplot(ohms.lm, partial=TRUE, se=TRUE)
fig6.4 <- function(plotit=TRUE){</pre>
    matchms2 <- model.matrix(with(fruitchms, ~ splines::ns(juice, 2)))</pre>
    matchms3 <- model.matrix(with(fruitchms, ~ splines::ns(juice, 3)))</pre>
    m <- dim(matohms3)[1]</pre>
    longdf1 <- data.frame(juice=rep(fruitohms$juice,4),</pre>
                          basis2 = c(as.vector(matohms2),rep(NA,m)),
                          basis3 = as.vector(matohms3),
                          gp = factor(rep(c("Intercept",
                          paste("spline",1:3, sep="")),
                          rep(m,4)))
    gph1 <- xyplot(basis3 ~ juice | gp, data=longdf1, layout=c(1,4),</pre>
                  scales=list(tck=0.5),
                  ylab="Basis terms", strip=FALSE,
                   strip.left=strip.custom(strip.names=TRUE,
                  var.name="",
                   sep=expression(""),
                   factor.levels=c("Constant", "Basis 1", "Basis 2",
                   "Basis 3")),
                  par.settings=simpleTheme(lty=c(2,2,1,1)),
                  panel=function(x,y,subscripts){
                       llines(smooth.spline(x,y))
                       y2 <- longdf1$basis2[subscripts]</pre>
                       if(!any(is.na(y2))) llines(smooth.spline(x,y2),lty=1)},
                   outer=TRUE.
                   legend=list(top=list(fun=grid::textGrob,
```

```
args=list(label="A:Basis functions",
                               just="left", x=0))))
    b2 <- coef(lm(I(ohms/1000) ~ splines::ns(juice,2), data=fruitohms))
    b3 <- coef(lm(I(ohms/1000) ~ splines::ns(juice,3), data=fruitohms))
    spline2 <- as.vector(sweep(matchms2, 2, b2, "*"))</pre>
    spline3 <- as.vector(sweep(matchms3, 2, b3, "*"))</pre>
    longdf2 <- data.frame(juice=rep(fruitohms$juice,4),</pre>
                          spline2 = c(spline2, rep(NA,m)), spline3=spline3,
                          gp = factor(rep(c("Intercept",
                                      paste("spline",1:3, sep="")),
                          rep(m,4))))
    yran <- range(c(spline2, spline3))</pre>
    yran < -c(-6, 8.5)
    gph2 <- xyplot(spline3 ~ juice | gp, data=longdf2, layout=c(1,4),</pre>
                   scales=list(tck=0.5, y=list(at=c(-4, 0, 4,8))), ylim=yran,
                  ylab="Add these contributions (ohms x 1000)", strip=FALSE,
                   strip.left=strip.custom(strip.names=TRUE,
                  var.name="",
                   sep=expression(""),
                  factor.levels=c("Const","Add 1","Add 2","Add 3")),
                  par.settings=simpleTheme(lty=c(2,2,1,1)),
                  panel=function(x,y,subscripts){
                       llines(smooth.spline(x,y))
                       y2 <- longdf2$spline2[subscripts]</pre>
                       if(!any(is.na(y2))) llines(smooth.spline(x,y2),lty=1)},
                   outer=TRUE,
                   legend=list(top=list(fun=grid::textGrob,
                               args=list(label="B: Contribution fo fitted curve",
                               just="left", x=0))))
    if(plotit){
        print(gph1, position=c(0,0,.5,1))
        print(gph2, position=c(.5,0,1,1), newpage=FALSE)
    invisible(list(gph1, gph2))
fig6.5 <- function(){</pre>
    res <- resid(lm(log(Time) ~ log(Distance), data=worldRecords))</pre>
    wr.gam <- gam(res ~ s(log(Distance)), data=worldRecords)</pre>
    plot(wr.gam, residuals=TRUE, pch=1, las=1, ylab="Fitted smooth")
```

```
fig6.6 <-
function () {
    res <- resid(lm(log(Time) ~ log(Distance), data=worldRecords))</pre>
    wr.gam <- gam(res ~ s(log(Distance)), data=worldRecords)</pre>
    gam.check(wr.gam)
fig6.7 <-
function (mf=3,nf=2)
  opar \leftarrow par(mfrow=c(mf,nf), mar=c(0.25, 4.1, 0.25, 1.1))
    set.seed(29)
                          # Ensure exact result is reproducible
    res <- resid(lm(log(Time) ~ log(Distance), data=worldRecords))</pre>
    npanels <- mf*nf
    for(i in 1:npanels){
        permres <- sample(res) # Random permutation</pre>
                                          # 0 for left-handers; 1 for right
        perm.gam <- gam(permres ~ s(log(Distance)), data=worldRecords)</pre>
        plot(perm.gam, las=1, rug=if(i<5) FALSE else TRUE, ylab="Fit")</pre>
 par(opar)
fig6.8 <- function(){</pre>
    meuse.gam <- gam(log(lead) ~ s(elev) + s(dist) + ffreq + soil,
                            data=meuse)
    plot(meuse.gam, residuals=TRUE, se=TRUE, pch=1)
    termplot(meuse.gam, terms="ffreq", se=TRUE)
    termplot(meuse.gam, terms="soil", se=TRUE)
fig6.9 <- function(nsim=1000, caption=NULL){</pre>
  opar \leftarrow par(mfrow=c(1,2), oma=c(0,0,1.6,0.6))
  if(missing(caption))captCol <- "black" else captCol <- "blue"</pre>
  if(is.null(caption))caption <- paste("Graphs are from", nsim, "simulations")
    if(!exists("meuseML.gam"))
    meuseML.gam <- gam(log(lead) ~ s(elev) + s(dist) + ffreq + soil,</pre>
                        data=meuse, method="ML")
```

if(!exists("meusexML.gam"))

## Now simulate from meuseML.gam

```
simY <- simulate(meuseML.gam, nsim=nsim)</pre>
    simResults <- matrix(0, nrow=nsim, ncol=3)</pre>
    colnames(simResults) <- c( "deltaDf", "fsim", "psim")</pre>
    for(i in 1:nsim){
        mML.gam <- gam(simY[,i] ~ s(elev) + s(dist) + ffreq + soil,
                        data=meuse, method="ML")
        mxML.gam <- gam(simY[,i] ~ s(elev, dist) + ffreq + soil,</pre>
                         data=meuse, method="ML")
        aovcomp <- anova(mML.gam, mxML.gam, test="F")</pre>
        simResults[i,] <- unlist(aovcomp[2, c("Df", "F", "Pr(>F)")])
    ## Now plot the £p£-statistics and £F£-statistics
    ## against the change in degrees of freedom:
    colcode <- c("gray", "black")[1+(simResults[,"deltaDf"]>=1)]
    simResults <- as.data.frame(simResults)</pre>
    plot(psim ~ deltaDf, log="y", xlab="Change in degrees of freedom",
         ylab=expression(italic(p)*"-value"), col=colcode, data=simResults)
    abline(v=1, lty=2, col="gray")
    mtext("A", side=3, line=0.25, adj=0)
    mtext("1", side=1, at=1, line=0, cex=0.75)
    plot(fsim ~ deltaDf, log="y", xlab="Change in degrees of freedom",
         ylab=expression(italic(F)*"-statistic"), col=colcode,
         data=simResults)
    abline(v=1, lty=2, col="gray")
    mtext("1", side=1, at=1, line=0, cex=0.75)
    mtext("B", side=3, line=0.25, adj=0)
 mtext(side=3, line=0.5, adj=0.052, outer=TRUE, caption, col=captCol)
    invisible(simResults)
  par(opar)
fig6.10A <- function(){</pre>
    if(!exists("meuseML.gam"))
    meuseML.gam <- gam(log(lead) ~ s(elev) + s(dist) + ffreq + soil,</pre>
                        data=meuse, method="ML")
    plot(meuseML.gam)
    termplot(meuseML.gam, terms="ffreq", se=TRUE)
    termplot(meuseML.gam, terms="soil", se=TRUE)
    mtext(side=3, line=0.65, "A: Add effects of dist and elev", outer=TRUE,
          cex=0.8, adj=0)
```

```
fig6.10B <- function(){</pre>
    if(!exists("meusexML.gam"))
    meusexML.gam <- gam(log(lead) ~ s(elev, dist) + ffreq + soil,</pre>
                         data=meuse, method="ML")
    plot(meusexML.gam)
    termplot(meusexML.gam, terms="ffreq", se=TRUE)
    termplot(meusexML.gam, terms="soil", se=TRUE)
    mtext(side=3, line=0.65, "B: Fit surface to dist and elev", outer=TRUE,
      cex=0.8, adj=0)
fig6.10 <- function()</pre>
print("Run fig6.10A() and fig6.10B() separately")
fig6.11 <- function(){
    hand <- with(cricketer, as.vector(as.vector(unclass(left)-1)))</pre>
                                          # 0 for left-handers
                                          # 1 for right
    hand.gam <- gam(hand ~ s(year), data=cricketer, family=binomial)</pre>
    plot(hand.gam, las=1, xlab="", ylab="Pr(left-handed)",
         trans=function(x)\exp(x)/(1+\exp(x)),
         shift=mean(predict(hand.gam)))
fig6.12 <- function(mf=3,nf=2){
    opar \leftarrow par(mfrow=c(mf,nf), mar=c(0.25, 4.1, 0.25, 1.1))
    npanel <- mf*nf
    for(i in 1:npanel){
        hand <- sample(c(0,1), size=nrow(cricketer), replace=TRUE,
                        prob=c(0.185, 0.815))
                                          # 0 for left-handers
                                          # 1 for right
        hand.gam <- gam(hand ~ s(year), data=cricketer, family=binomial)
        plot(hand.gam, las=1, xlab="",
             rug=if(i<4)FALSE else TRUE,</pre>
             trans=function(x)\exp(x)/(1+\exp(x)),
             shift=mean(predict(hand.gam)))
        par(opar)
```

```
fig6.13 <- function(){
    rtlef <- data.frame(with(cricketer, as(table(year, left), "matrix")))</pre>
    rtlef <- within(rtlef, year <- as.numeric(rownames(rtlef)))</pre>
    right.gam <- gam(right ~ s(year), data=rtlef, family=poisson)</pre>
    left.gam <- gam(left ~ s(year), data=rtlef, family=poisson)</pre>
    rtlef <- within(rtlef,</pre>
                 {fitright <- predict(right.gam, type="response")</pre>
                  fitleft <- predict(left.gam, type="response")})</pre>
    key.list <- list(text=expression("Right-handers", "Left-handers",</pre>
        "Left-handers "%*%" 4"),
                      corner=c(0,1), x=0, y=0.985,
                      points=FALSE, lines=TRUE)
    parset <- simpleTheme(col=c("blue", "purple", "purple"),</pre>
                            lty=c(1,1,2), lwd=c(2,2,1)
    gph <- xyplot(fitright+fitleft+I(fitleft*4) ~ year, data=rtlef,</pre>
                   auto.key=key.list, par.settings=parset,tck=-0.05,
                   xlab="",
                   ylab="Number of cricketers\nborn in given year",
                   type="1", ylim=c(0,70))
    print(gph)
```

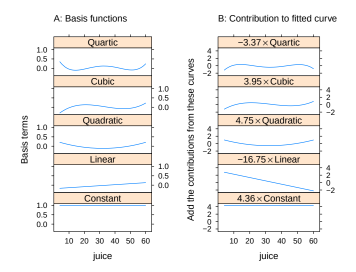
```
countAccs <- function(data=airAccs, dateCol="Date",</pre>
                       fromDate = as.Date("2006-01-01"),
                       by="1 day", prefix="num"){
  dfCount <- eventCounts(data, dateCol=dateCol,</pre>
                        from= fromDate, by=by,
                        prefix=prefix)
  dfCount[, "day"] <- julian(dfCount[,"Date"], origin=fromDate)</pre>
  dfCount
##
fig6.14 <- function()</pre>
    print("Run the separate functions fig6.14A() and fig6.14B()")
fig6.14A <- function(fromDate=as.Date("2006-01-01"), basis.df=50){</pre>
  if(!exists('dfDay06'))dfDay06 <- countAccs(fromDate=fromDate)</pre>
year <- seq(from=fromDate, to=max(dfDay06$Date), by="1 year")</pre>
atyear=julian(year, origin=fromDate)
dfDay06.gam <-
  gam(formula = num ~ s(day, k=basis.df), family = quasipoisson,
      data = dfDay06)
av <- mean(predict(dfDay06.gam))</pre>
plot(dfDay06.gam, xaxt="n", shift=av, trans=exp, rug=FALSE, xlab="",
     ylab="Estimated rate per day")
```

## 2 Show the Figures

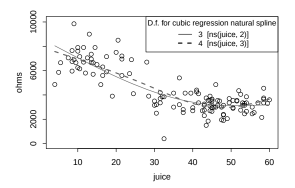
fig6.1()

```
pkgs <- c("DAAG","mgcv","splines","gamclass")
z <- sapply(pkgs, require, character.only=TRUE, warn.conflicts=FALSE)
if(any(!z)){
  notAvail <- paste(names(z)[!z], collapse=", ")
  print(paste("The following packages require to be installed:", notAvail))
}

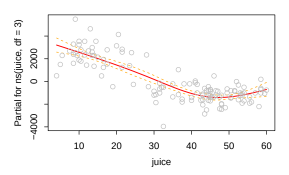
if(!exists("meuse")){
  msg <- "Cannot find package 'sp',"
      if(!require("sp"))
      return(paste(msg, "cannot do graph."))
  data("meuse", package="sp", envir = environment())
  }
}</pre>
```



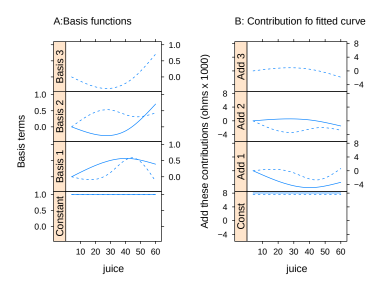
### fig6.2()



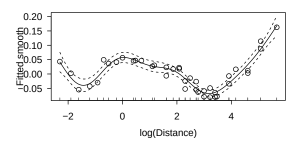
# fig6.3()



### fig6.4()



### fig6.5()



### fig6.6()

Method: GCV Optimizer: magic

Smoothing parameter selection converged after 6 iterations.

The RMS GCV score gradiant at convergence was 7.277e-07.

The Hessian was positive definite.

The estimated model rank was 10 (maximum possible: 10)

Model rank = 10 / 10

Basis dimension (k) checking results. Low p-value (k-index<1)

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'.

```
$\rm k^{\,\prime}$ edf k-index p-value s(log(Distance)) 9.00 8.32 1.16 0.77
```

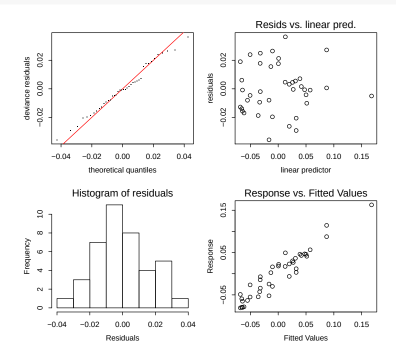
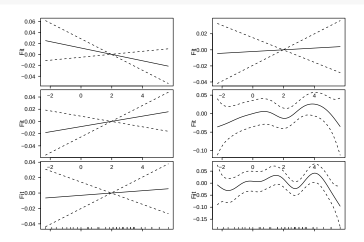
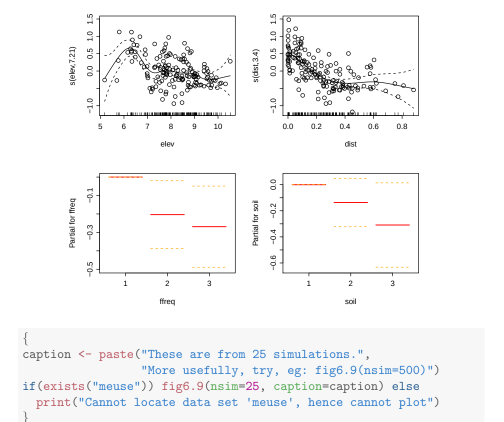


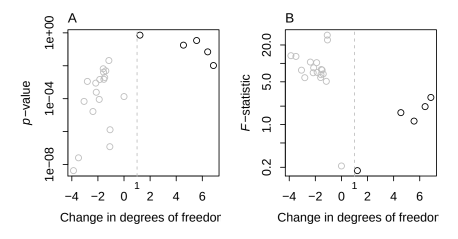
fig6.7()



if(exists("meuse")) fig6.8() else
 print("Cannot locate data set 'meuse', hence cannot plot")

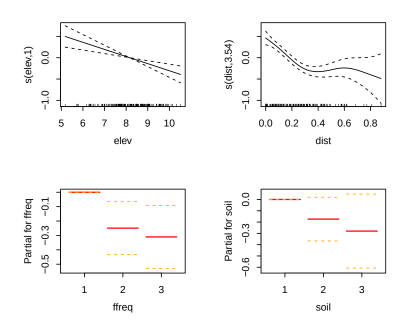


### These are from 25 simulations. More usefully, try, eg: fig6.9(nsim=500)



```
if(exists("meuse")) fig6.10A() else
    print("Cannot locate data set 'meuse', hence cannot plot")
```

#### A: Add effects of dist and elev



```
if(exists("meuse")) fig6.10B() else
    print("Cannot locate data set 'meuse', hence cannot plot")
```

#### B: Fit surface to dist and elev

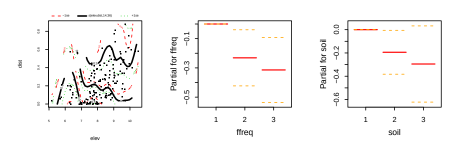


fig6.11()

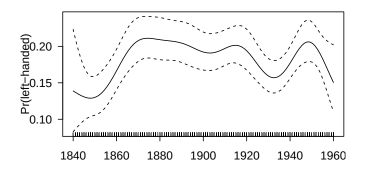


fig6.12()

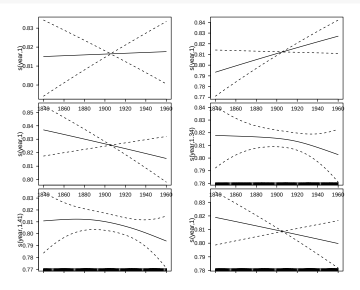
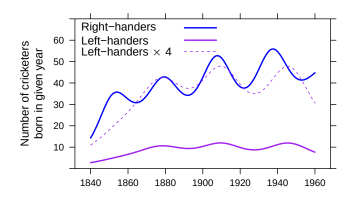


fig6.13()



```
{
  fig6.14A(fromDate=as.Date("2010-01-01"), basis.df=50)
  fig6.14B(fromDate=as.Date("2010-01-01"), basis.df=50)
}
```

