Plotting model residuals with plotres

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August 29, 2024

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1 Introduction

Residual plots are important for checking linear models, but they also are useful for other types of model. They can be used for an overview of the model's performance, to check for outliers, and to check if the response should be transformed.

The plotres function in the plotmo R package [10] makes it easy to plot residuals for a wide variety of R models. Figure 1 shows an example. It was produced with the following code:

library(rpart); library(plotmo)	<pre># plotres is in the plotmo package</pre>
library(earth); data(ozone1)	# get the ozone data
<pre>rpart.mod <- rpart(03 ~ ., data = ozone1)</pre>	<pre># generate an rpart model</pre>
<pre>plotres(rpart.mod)</pre>	<pre># plot its residuals</pre>



Figure 1: An example plotres plot. This example is for an rpart model.

Since this is an **rpart** model [14], plotres draws the model tree at the top left [8]. Different figures will be drawn in the top left for other types of model (Section 5).

The bottom left plot is a standard Residuals vs Fitted plot of the training data. In this example we see the quantized fits characteristic of **rpart** models. Each vertical line of points corresponds to a leaf of the tree. (We could use plotres's **jitter** argument to jitter the points to reduce overplotting caused by quantization here.) The red line is a lowess smooth.

In case 220, for example, we see that although the model estimates a high ozone level (the fitted value is 23), the observed level was actually a lot higher. From the QQ plot (bottom right) we see that this residual is indeed a bit unusual. Since it's the biggest absolute residual, it determines the right bound of the Cumulative Distribution plot (top right).

A variety of other residual plots can be generated by plotres as described on its help page.



Figure 2: Model of the Boston housing data. Note from the right figure that many house prices are artificially clipped at 50.

Residual plots of the Boston housing data

Residual plots reveal an often-overlooked issue in the popular Boston housing dataset [3].

We build a model which estimates house prices given various characteristics of the neighborhood. The left side of Figure 2 shows the model residuals i.e. the difference between the predicted and actual house prices in the training data. This example uses a gbm model, although any popular non-parametric modeling technique could be used:

```
data(Boston, package="MASS")  # boston housing data
library(gbm)
mod <- gbm(medv ~ ., data=Boston, n.trees=5000, interact=4)
library(plotmo)
plotres(mod, which=3)  # left figure (which=3 for residuals plot)
```

In the residuals plot the largest residual is for case 373, and we see an artificial line which starts at this residual and extends diagonally downwards to the right.¹ Investigation of these cases reveals that there are 16 houses with a price of 50 (right of Figure 2). It's likely that this price is artificial, i.e., the maximum price of a house was clipped to 50 (in units of \$1000, the data is from a 1970 US census).

If we remove these cases and rebuild the model we see a more healthy residual plot (middle of Figure 2):

This isn't to say that these artificial cases *should* be removed, only that we should be aware of them when deciding how to model the data. The well-known StatLib California housing data (e.g. [4] Section 10.14) exhibits the same issue in a more pronounced form (965 of the house values are clipped at \$500001).

¹Such lines are common for quantized responses, but not for continuous values like house prices.

2 Some examples

Here are some examples which illustrate plotres on various models.

```
library(earth) # for ozone1 data
data(ozone1)
lm.mod <- lm(03 ~ ., data=ozone1)</pre>
                                                            ## linear model
plotres(lm.mod)
earth.mod <- earth(03 ~ ., data=ozone1, degree=2)</pre>
                                                            ## earth
plotres(earth.mod) # equivalent to plot.earth
library(rpart)
                                                            ## rpart
rpart.mod <- rpart(03 ~ ., data=ozone1)</pre>
plotres(rpart.mod)
library(tree)
                                                            ## tree
tree.mod <- tree(03<sup>~</sup>., data=ozone1)
plotres(tree.mod)
library(randomForest)
                                                            ## randomForest
set.seed(2015)
rf.mod <- randomForest(03~., data=ozone1)</pre>
plotres(rf.mod)
library(gbm)
                                                            ## gbm
set.seed(2015)
gbm.mod <- gbm(03~., data=ozone1, dist="gaussian",
                interact=2, shrink=.01, n.trees=1000)
plotres(gbm.mod)
library(nnet)
                                                            ## nnet
set.seed(2015)
nnet.mod <- nnet(03~., data=scale(ozone1), size=2, decay=0.01, trace=FALSE)</pre>
plotres(nnet.mod, type="raw")
library(neuralnet)
                                                            ## neuralnet
set.seed(2015)
nn.mod <- neuralnet(03<sup>^</sup>humidity+temp, data=scale(ozone1), hidden=2)
plotres(nn.mod)
                                                            ## caret
library(caret)
set.seed(2015)
caret.earth.mod <- train(03~., data=ozone1, method="earth",</pre>
                           tuneGrid=data.frame(degree=2, nprune=10))
plotres(caret.earth.mod, type="raw")
```

This definitely isn't an exhaustive list of models supported by plotres. The packages used in the above code are [2, 5, 6, 11-15].

3 Limitations

Plotres is designed primarily for displaying standard **response** – **fitted** residuals for regression models with a single continuous response. For some model types it supports multiple responses and other kinds of residuals.

In general, it won't work on models that don't save the call or data with the model in a standard way. For further discussion please see *Accessing the model data* in the *plotmo* vignette.

4 Generating the residuals and calling predict

Plotres first tries to get the residuals by calling the **residuals** method for the model. If the call fails (which it will for models that don't have a **residuals** method), plotres must figure out the residuals manually. It does that using **predict**.

Plotres tries to use sensible default arguments for predict, but they won't always be correct (plotres can't know about the predict method for every kind of model). Change the defaults if necessary using plotres arguments with a predict. prefix. Plotres passes any argument prefixed with predict. directly to predict, after removing the prefix.

For example, predict.gbm has a n.trees argument, which plotres defaults to the total number of trees. But that can be changed, for example:

Use trace = 1 with plotres to see the arguments passed to predict and friends.

5 The which=1 plot and the "w1." prefix

The top left plot is a model-specific plot.² We call this plot the "which=1" plot. What gets plotted here depends on the model class. For example, for earth models this is a model selection plot and for glmnet models it's a coefficient profile plot.

Nothing will be displayed for some models. This isn't really an issue. You will see three instead of four plots when you call plotres with the default arguments.

For some models, the which=1 plot is called with default arguments programmed into plotres. Use trace = 1 to see those arguments.

²This plot is included by default, since by default which = 1:4.

Change the arguments passed to this plot by using plotres arguments with a w1. prefix. Plotres passes any argument prefixed with w1. directly to the which=1 plot, after removing the prefix. For example, w1.ylim=c(-10,20). Note the w1. prefix; plain ylim=c(-10,20) affects the residual plots, not this plot. See also the example below. It's very easy to forget the w1. prefix.

It may happen that the which=1 plot is only partially plotted, or messes up the page for further plots. For example, plots that use grid graphics will cause problems. In that case, call plotres with a which argument that excludes 1 (don't use the default which). We would be interested in hearing about models that cause this kind of bad behavior.

An example

Here's a which=1 plot showing glmnet taming the longley data (Figure 3):

```
library(glmnet); library(plotmo); data(longley)
mod <- glmnet(data.matrix(longley[,1:6]), longley[,7])
plotres(mod, which=1)  # left side of the figure</pre>
```

Change the parameters passed to the plot using the w1. prefix:

<pre>plotres(mod, which=1,</pre>	# right side of the figure
w1.xvar="norm",	<pre># pass xvar="norm" to the plot</pre>
w1.col=1:3)	# change the color scheme



Figure 3: A glmnet [1] model of the longley data [7]. Just the which=1 plot is shown. Left plotres(glmnet.mod) Right plotres(glmnet.mod, w1.xvar="norm", w1.col=1:3))

6 Notes on glmnet and glmnetUtils models

The following code plots an example glmnet model (Figure 4). The previous page also has some examples.

```
library(glmnet)
library(plotmo)
x <- matrix(rnorm(100 * 10), 100, 10)  # n=100 p=10
y <- x[,1] + x[,2] + 3 * rnorm(100)  # y depends only on x[,1] and x[,2]
mod <- glmnet(x, y)
plotres(mod)</pre>
```

For the which=1 plot (Figure 4 top left), plotres makes an internal call to plot_glmnet (in the plotmo package). This function is much like plot.glmnet in the glmnet package. It has an extra xvar option "rlambda", which plotres uses by default. (This allows s to be plotted and variable names on the right of the plot.) Override that default by passing say w1.xvar="norm" to plotres.

The plot is annotated with a vertical gray line showing the s parameter. This is the penalty lambda that plotres passes internally to predict.glmnet when generating the residuals for the residual plots (Section 4). By default, plotres calls predict.glmnet with type="response" and s=0 (no penalty). Change that by passing say predict.s=.02 to plotres. (In fact, you can pass any argument to predict.glmnet by prefixing the argument with predict. as described in Section 4.)

When using plotres, arguments such as ylim can be passed on to plot_glmnet using a w1. prefix as described in Section 5, for example w1.ylim=c(-3,3). When calling plot_glmnet directly, don't use the w1. prefix.

Use w1.s.col=0 to remove the s vertical gray line, or w1.s.col="green" to make it green (for example).

Plotres can also be used with cv.glmnet models (it will invoke plot.cv.glmnet for the which=1 plot).

For multiple response models, use plotres's **nresponse** argument to select which response is plotted. The type.coef argument of plot.multnet isn't supported.

s=0 glmnet(x=x, y=y)



Figure 4: A glmnet model plotted with plotres.

In this example, all variables except V1 and V2 are noise variables.

To calculate the residuals, s=0 was passed to predict.glmnet, as indicated by the vertical gray line in the top left plot.

The \mathbf{x} matrix has no column names, so the variables have been automatically named with a \mathbf{V} prefix. The variable names are spread out to prevent overplotting, but we can still make out which curve is associated with which variable.

7 Notes on gbm models

Plotres uses the plot_gbm function (in the plotmo package) for plotting gbm models. Figure 5 shows an example, generated with the following code. (Only the which=1 plot_gbm plot is shown; the residual plots are the same as for other models.)

plotres(gbm.mod)

The vertical dotted lines and the corresponding colored numbers along the top of the plot show the number of trees selected by various criteria.

The OOB curve is the out-of-bag improvement, similar to the gbm.perf plot. The OOB curve is on a different scale to the other curves, so shouldn't really be shown on the same plot. We force it in by rescaling and shifting it. Although helpful for comparing model-selection criteria, this can also lead to confusion: the scale on the left doesn't apply to the OOB curve. A dashed curve is used as a reminder.

The vertical gray line shows the n.trees parameter. This is the number of trees that plotres passes internally to predict.gbm when generating the residuals (Section 4), although the residual plots aren't shown here. In the residual plots, only train.fraction of the residuals will be shown if train.fraction was specified in the original call to gbm.

When generating the residuals, by default plotres calls predict.gbm with type="response" and n.trees set to all the trees. Change that by passing say predict.n.trees=200 to plotres. (In fact, you can pass any argument to predict.gbm by prefixing the argument with predict. as described in Section 4.)

When using plotres, arguments such as ylim can be passed on to plot_gbm using a w1. prefix as described in Section 5, for example w1.ylim=c(-3,3). When calling plot_gbm directly, don't use the w1. prefix.



Figure 5: A gbm model.

Just the plotres which=1 plot is shown (same as calling plot_gbm directly).

The plot is a little crowded because both the train.fraction and cv.folds arguments were used in the call to gbm when building the model.

8 Comparison to plot.lm

The function plot.lm automatically standardizes residuals for some of the plots. In contrast, with plotres we must explicitly specify when the residuals should be standardized. This is because standardization isn't possible or appropriate for most of the models that plotres is designed for.

The different which numbering scheme used by plotres is mostly an historical legacy (from the days when this function was part of the earth package).

As a somewhat academic exercise, here's how plot.lm can be emulated with plotres (Figure 9):

```
plotlm <- function(object) # similar to plot.lm</pre>
Ł
    plotres(object, which=3, center=FALSE,
                                                          # residuals vs fitted
            caption=paste(deparse(object$call), collapse=" "))
    plotres(object, which=4, standardize=TRUE)
                                                          # QQ plot
                                                          # scale-location plot
    plotres(object, which=6, standardize=TRUE, main="Scale-Location")
    plotres(object, which=3, versus=4, standardize=TRUE) # leverage plot
}
fit <- lm(Volume ~ ., data = trees)</pre>
                                       # simple linear model
par(mfrow = c(2,2), oma = c(0,0,3,0)) # four plots on page, space for caption
plot(fit)
                                         # call plot.lm
                                         # call our version of plot.lm
plotlm(fit)
```

9 FAQ and common error messages

Please see the FAQ and *Common error messages* in the plotmo vignette.

See also the *Notes on some packages* in that vignette.

Most of the difficulties associated with plotting residuals by functions like plotres arise because the model-building function neglected to include some essential fields in the model, for example the call. See the vignette *Guidelines for S3 Regression Models* [9].

Why is nothing displayed for which=1?

See Section 5. For some model classes, nothing will displayed for which=1. By default, plotres will plot three instead of four plots. (Package authors may contact the main-tainer of plotres to add which=1 support for their model plot; this is usually a trivial extension to plotres.)

Residuals vs Fitted Normal Q-Q 9 310 Standardized residuals 310 2 LC. 00 Residuals 8 0 0 °0 8 0 0⁰ 0 Ϋ́ ī 0 0₁₈ 018 70 -2 10 20 40 50 60 2 30 0 -1 1 Fitted values **Theoretical Quantiles** Scale-Location Residuals vs Leverage с 310 1.5 VIStandardized residuals 310 Standardized residuals 0.5 018 2 62 °0 0 03 0 oC 1.0 0 C 000 0 0 0 Ф 0.5 0 0 С g ĩ 0 റ് 0 0 \cap Cook's distance₁₈ 0.0 0.5 0.00 10 20 30 40 50 60 70 0.05 0.10 0.15 0.20 Fitted values Leverage

plot.lm

plot.Im emulated with plotres



Figure 6: plot.lm and plotres.

How do I pass arguments to predict() called internally by plotres?

Plotres passes any argument prefixed with **predict**. directly to **predict**, after removing the prefix.

For example plotres(glmnet.mod, predict.s = .8) will pass s = .8 to plotres's internal calls to predict.glmnet.

For more information, please see the description of the "..." argument in the help page of plotres.

References

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