

# RmarkdownDemo

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## Configure

If you want to beautify your output, it always starts here. There are many options, and a few are laid out below. The **knitr** package has lots of options explained [here](#) in detail.

Part of configuring your script is loading the correct packages. Always load all packages together at the top. That way future users will know exactly what they need to install.

If you ever want someone else to be able to perfectly reproduce your results, always set the random seed at the top. Any number will do. Note that it never hurts to set the seed, *but* robust results should always stand up to random number generators.

```
set.seed(1415)
```

## Generate fake data

The **x** value is just numbers 1-100 for an x axis value. This might be time or distance, etc. For the response variable, generate a random normal distribution with the **rnorm** function, and then add a trend with the **seq** function. Then we'll add some fake treatments with **letters**.

```
x <- 1:100
y <- rnorm(100, sd=3) + seq(10.05, 20, 10/100)
z <- factor(rep(letters[1:5], each=20))
dat <- data.frame(x, y, z)
```

## Tables in knitr

This is an ugly way to preview data or display tables.

```
head(dat)
```

```

      x      y z
1 1 13.615 a
2 2 13.997 a
3 3  9.891 a
4 4 11.290 a
5 5 10.473 a
6 6 12.369 a

```

The `knitr` package has a simple built-in function for dealing with tables. This works well in either html or pdf output.

```

kable(head(dat))

```

x	y	z
1	13.615	a
2	13.997	a
3	9.891	a
4	11.290	a
5	10.473	a
6	12.369	a

## R commands embedded in prose

One of the best features in `knitr` and Rmarkdown generally, is the ability to embed real R commands in sentences, so that you can report actual values instead of constantly copying and pasting when results change a little bit.

This table has 100 rows and 3 columns. The 'x' variable starts at 1 and ends at 100.

## Explore the data

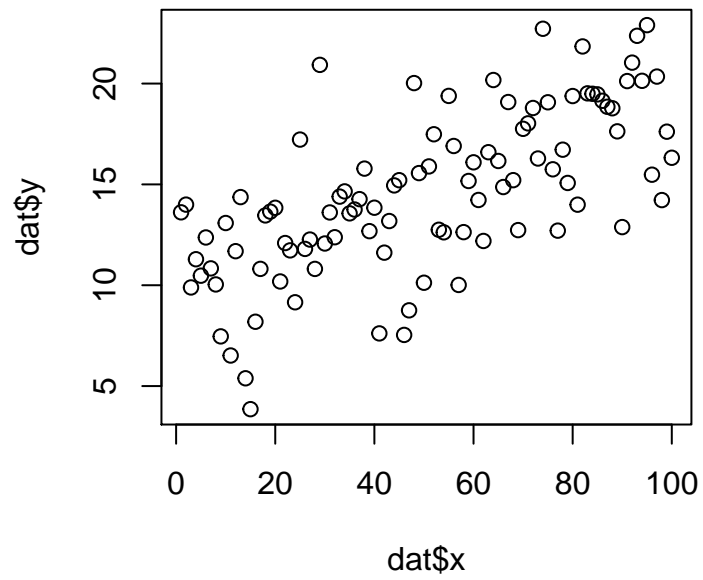
Plot the data - a trend emerges! Here are several ways to look at the data.

1. The rough R default
2. boxplots using the factors
3. points with default colors
4. same colors but with nicer plotting characters (`pch`)

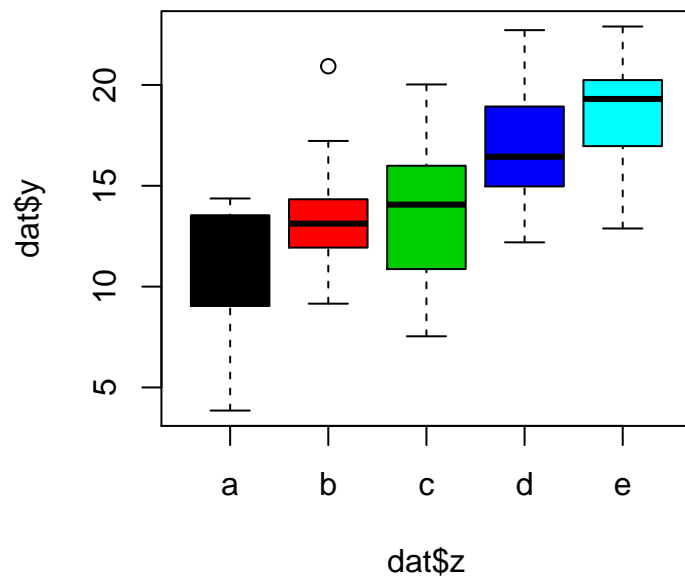
```

plot(dat$y ~ dat$x)

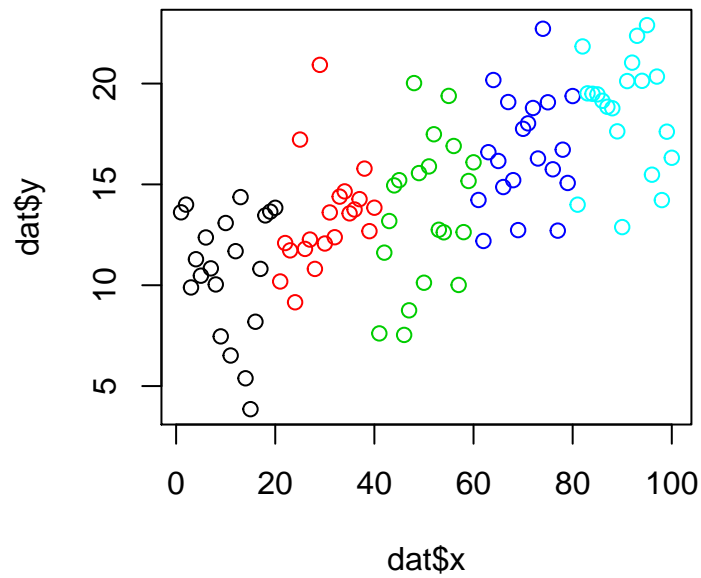
```



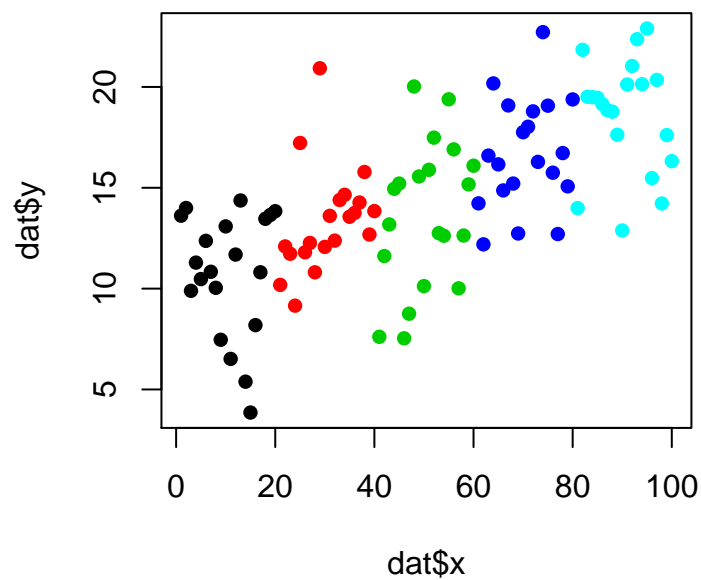
```
plot(dat$y ~ dat$z, col=unique(dat$z))
```



```
plot(dat$y ~ dat$x, col=dat$z)
```



```
plot(dat$y ~ dat$x, col=dat$z, pch=16)
```



## Linear models

Let's see if the linear and grouped trends are significant using a linear model. The model can be stored, and then we can pull out pieces as the analysis progresses.

```
lm.xy <- lm(y ~ x, data=dat)
lm.zy <- lm(y ~ z, data=dat)

kable(summary(lm.xy)$coefficients)
```

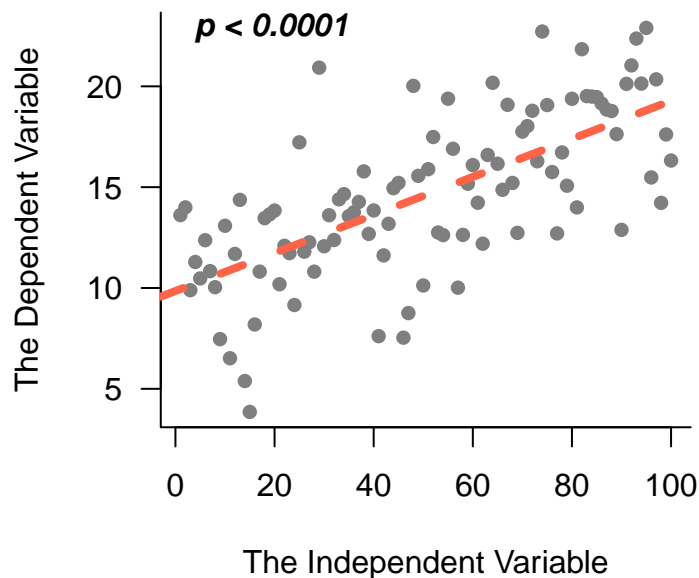
	Estimate	Std. Error	t value	Pr(> t )
(Intercept)	9.8493	0.5971	16.4959	0.0000
x	0.0943	0.0103	9.1877	0.0000

```
kable(summary(lm.zy)$coefficients)
```

	Estimate	Std. Error	t value	Pr(> t )
(Intercept)	10.7411	0.6683	16.0730	0.0000
zb	2.6180	0.9451	2.7702	0.0067
zc	2.9348	0.9451	3.1054	0.0025
zd	5.9349	0.9451	6.2798	0.0000
ze	7.8656	0.9451	8.3227	0.0000

Since we have a clear pattern, plot the line we just modeled.

```
plot(dat$y ~ dat$x, pch=16, col='gray50',
     las=1, bty='l',
     xlab='The Independent Variable',
     ylab='The Dependent Variable')
abline(lm.xy, col='tomato', lwd=4, lty=2)
text(0, max(dat$y), 'p < 0.0001', font=4, pos=4)
```



Much better.

## Improve the plots

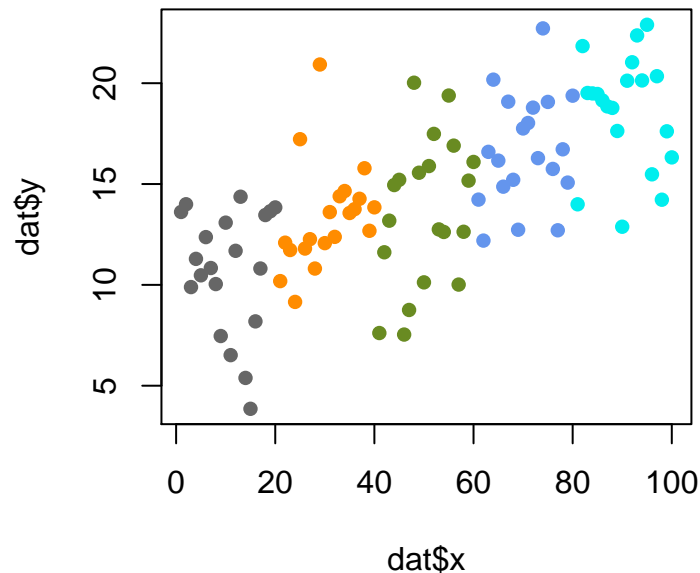
Default R colors are useful but not that aesthetic. So we can assign them however we want to. Assigning them to the same dataframe keeps all data points lined up perfectly. ***R does not line up your data***

*automatically!! You have to make sure everything is lined up before you can trust results!!*

```
dat$col <- 'gray40'
dat$col[dat$z == 'b'] <- 'darkorange'
dat$col[dat$z == 'c'] <- 'olivedrab4'
dat$col[dat$z == 'd'] <- 'cornflowerblue'
dat$col[dat$z == 'e'] <- 'cyan2'
colors5 <- unique(dat$col)
```

And check the new colors. Note they are now a bit more colorblind-proof.

```
plot(dat$y ~ dat$x, col=dat$col, pch=16)
```



## Aggregate data across treatments

The linear model created above was ok for  $x$  vs  $y$ . However, what if we want to take advantage of groups instead of just the simple linear relationship?

Let's create a dataset that combines data from each group (a, b, c, d, e). The **aggregate** function is perfect.

Also, R does not have a default function for standard error, so we'll create one. Creating functions in R is pretty simple, and becomes mandatory anytime you are going to repeat lines of code over and over.

```
se <- function(a) {
  sd(a)/sqrt(length(a))
}
```

First, create an empty data frame, and then fill in the row and column names. Next fill in the columns with the **aggregate** function.

```
grouped <- data.frame(matrix(0, nrow=nlevels(dat$z), ncol=3))
names(grouped) <- c('mean', 'sd', 'se')
row.names(grouped) <- levels(dat$z)
grouped$mean <- aggregate(dat$y, by=list(dat$z), FUN='mean')$x
```

```
grouped$sd <- aggregate(dat$y, by=list(dat$z), FUN='sd')$x
grouped$se <- aggregate(dat$y, by=list(dat$z), FUN='se')$x
grouped
```

```
      mean    sd    se
a 10.74 3.072 0.6870
b 13.36 2.589 0.5789
c 13.68 3.632 0.8121
d 16.68 2.731 0.6106
e 18.61 2.805 0.6272
```

Create a simple model that now takes advantage of the replicated regression study design.

```
lm.RepReg <- lm(y ~ as.numeric(z), data=dat)
kable(summary(lm.RepReg)$coefficients)
```

	Estimate	Std. Error	t value	Pr(> t )
(Intercept)	8.8974	0.7011	12.6908	0.0000
as.numeric(z)	1.9048	0.2114	9.0111	0.0000

Plot the data showing error bars (or standard deviation bars in this example).

- The `arrows` function is one of the easiest way to create error bars.
- After the error bars are in place, plot the colored points on top.

```
y.lim <- c(min(grouped$mean - grouped$sd),
           max(grouped$mean + grouped$sd))
plot(grouped$mean ~ c(1:5), type='n',
     ylim=y.lim, las=1, bty='n',
     xlab = 'This other variable',
     ylab = 'The response variable')
arrows(x0 = c(1:5), y0 = grouped$mean + grouped$sd,
       x1 = c(1:5), y1 = grouped$mean - grouped$sd,
       col='gray50', angle=90, code=3, length=0.08, lwd=2)
points(grouped$mean ~ c(1:5),
       pch=21, bg=colors5, col='gray20', cex=2)
points(dat$y ~ jitter(as.numeric(dat$z)),
       col=alpha(dat$col, alpha=.3), pch=16)
abline(lm.RepReg, col='tomato', lty=2, lwd=2)
legend('topleft',
      legend=row.names(grouped),
      pt.bg=colors5, pch=21, pt.cex=1.5,
      bty='n', text.col='gray30', y.intersp=.8)
```

