Exploratory Data Analysis Course Notes

Xing Su

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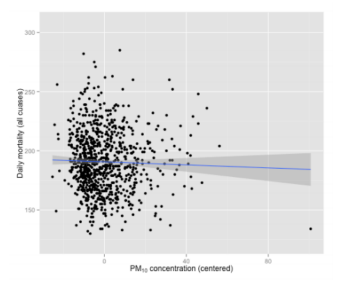
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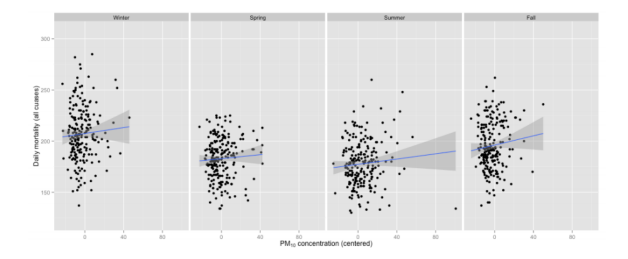
Principle of Analytic Graphics

- Principle 1: Show Comparisons
 - always comparative (compared to what)
 - randomized trial compare control group to test group
 - evidence for a hypothesis is always relative to another competing hypothesis
- Principle 2: Show causality/mechanism/explanation/systematic structure
 - form hypothesis to evidence showing a relationship (causal framework, why something happened)
- Principle 3: Show multivariate data
 - more than 2 variables because the real world is multivariate
 - show as much data on a plot as you can
 - example

Loading required package: pacman



- slightly negative relationship between pollution and mortality
 - when split up by season, the relationships are all positive \rightarrow season = confounding variable



• Principle 4: Integration of evidence

- use as many modes of evidence/displaying evidence as possible (modes of data presentation)
- integrate words/numbers/images/diagrams (information rich)
- analysis should drive the tool

• Principle 5: Describe/document evidence with appropriate labels/scales/sources

- add credibility to that data graphic
- Principle 6: Content is the most important
 - analytical presentations ultimately stand/fall depending on quality/relevance/integrity of content

Exploratory Graphs (examples)

- Purpose: understand data properties, find pattern in data, suggest modeling strategies, debug
- Characteristics: made quickly, large number produced, gain personal understanding, appearances and presentation are aren't as important

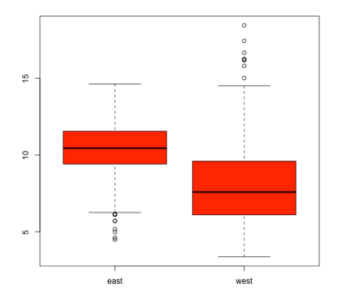
One Dimension Summary of Data

- summary(data) = returns min, 1st quartile, median, mean, 3rd quartile, max
- boxplot(data, col = "blue") = produces a box with middles 50% highlighted in the specified color
 - whiskers = $\pm 1.58IQR/\sqrt{n}$
 - * $IQR = interquartile range, Q_3 Q_1$
 - box = 25%, median, 75%

- histograms (data, col = "green") = produces a histogram with specified breaks and color
 - breaks = 100 = the higher the number is the smaller/narrower the histogram columns are
- rug(data) = density plot, add a strip under the histogram indicating location of each data point
- barplot(data, col = wheat) = produces a bar graph, usually for categorical data
- Overlaying Features
- abline(h/v = 12) = overlays horizontal/vertical line at specified location
 - col = "red" = specifies color
 - lwd = 4 = line width
 - lty = 2 = line type

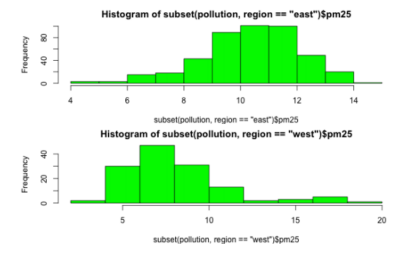
Two Dimensional Summaries

- multiple/overlay 1D plots (using lattice/ggplot2)
- box plots: boxplot(pm25 ~ region, data = pollution, col = "red")



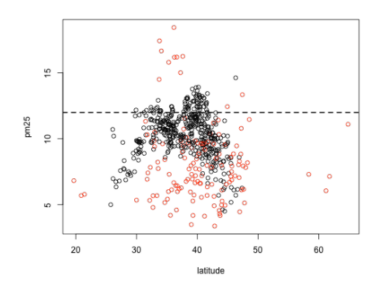
• histogram:

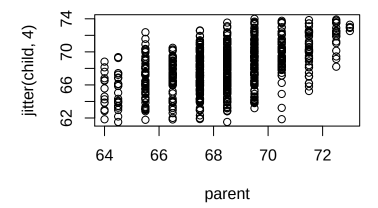
- par(mfrow = c(2, 1), mar = c(4, 4, 2, 1)) = set margin
- hist(subset(pollution, region == "east")\$pm25, col = "green") = first histogram
- hist(subset(pollution, region == "west") pm25, col = "green") = second histogram



\bullet scatterplot

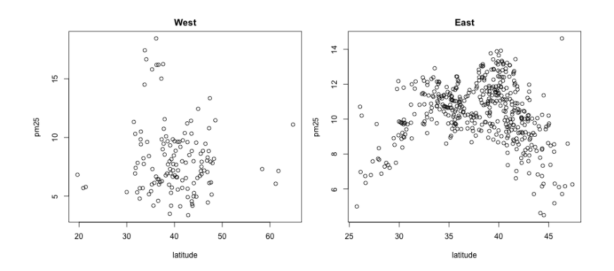
- with(pollution, plot(latitude, pm25, col = region))
- abline(h = 12, lwd = 2, lty = 2) = plots horizontal dotted line
- plot(jitter(child, 4)~parent, galton) = spreads out data points at the same position to simulate measurement error/make high frequency more visibble





ullet multiple scatter plots

- par(mfrow = c(1, 2), mar = c(5, 4, 2, 1)) = sets margins
- with(subset(pollution, region == "west"), plot(latitude, pm25, main = "West")) =
 left scatterplot
- with(subset(pollution, region == "east"), plot(latitude, pm25, main = "East")) =
 right scatterplot



Process of Making a Plot/Considerations

- where will plot be made? screen or file?
- how will plot be used? viewing on screen/web browser/print/presentation?
- large amount of data vs few points?
- need to be able to dynamically resize?
- plotting system: base, lattice, ggplot2?

Base Plotting

- blank canvas, "artist's palette", start with plot function
- annotations text, lines, points, axis
- convenient, but cannot go back when started (need to plan ahead)
- everything need to be manually set carefully to be able to achieve the desired effect (margins)
- core plotting/graphics engine in R encapsulated in the following
 - graphics: plotting functions for vase graphing system (plot, hist, boxplot, text)
 - grDevices: contains all the code implementing the various graphics devices (x11, PDF, PostScript, PNG, etc)
- Two phase: initialize, annotate
- calling plot(x, y) or hist(x) will launch a graphics device and draw a plot on device
 - if no argument specified, default called
 - parameters documented in "?par"
 - Note: it is some times necessary to convert column/variable to factor to make plotting easier
 - * airquality <- transform(airquality, Month = factor(month))

Base Graphics Functions and Parameters

• arguments

- pch: plotting symbol (default = open circle)
- lty: line type (default is solid)
 - * 0=blank, 1=solid (default), 2=dashed, 3=dotted, 4=dotdash, 5=longdash, 6=twodash
- lwd: line width (integer)
- col: plotting color (number string or hexcode, colors() returns vector of colors)
- xlab, ylab: x-y label character strings
- cex: numerical value giving the amount by which plotting text/symbols should be magnified relative to the default
 - * cex = 0.15 * variable: plot size as an additional variable
- par() function = specifies global graphics parameters, affects all plots in an R session (can be overridden)
 - las: orientation of axis labels
 - bg: background color
 - mar: margin size (order = bottom left top right)
 - oma: outer margin size (default = 0 for all sides)
 - mfrow: number of plots per row, column (plots are filled row-wise)
 - mfcol: number of plots per row, column (plots are filled column-wise)
 - can verify all above parameters by calling par("parameter")

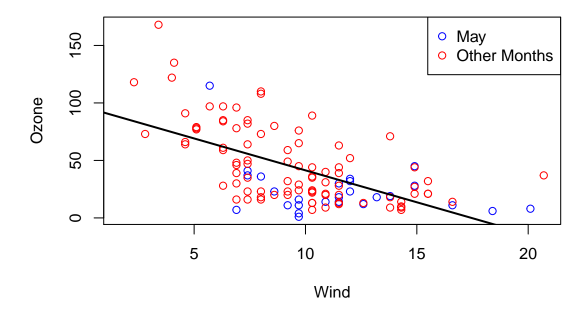
• plotting functions

- lines: adds liens to a plot, given a vector of x values and corresponding vector of y values
- points: adds a point to the plot
- text: add text labels to a plot using specified x,y coordinates
- title: add annotations to x,y axis labels, title, subtitles, outer margin
- mtext: add arbitrary text to margins (inner or outer) of plot
- axis: specify axis ticks

Base Plot Example

```
p_load("datasets")
# type ="n" sets up the plot and does not fill it with data
with(airquality, plot(Wind, Ozone, main = "Ozone and Wind in New York City", type = "n"))
# subsets of data are plotted here using different colors
with(subset(airquality, Month == 5), points(Wind, Ozone, col = "blue"))
with(subset(airquality, Month != 5), points(Wind, Ozone, col = "red"))
legend("topright", pch = 1, col = c("blue", "red"), legend = c("May", "Other Months"))
model <- lm(Ozone ~ Wind, airquality)
# regression line is produced here
abline(model, lwd = 2)</pre>
```

Ozone and Wind in New York City

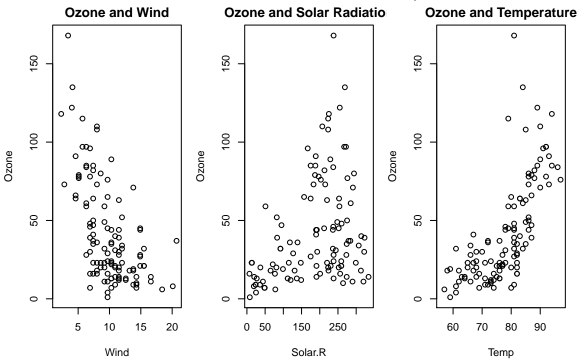


Multiple Plot Example

• Note: typing example(points) in R will launch a demo of base plotting system and may provide some helpful tips on graphing

```
# this expression sets up a plot with 1 row 3 columns, sets the margin and outer margins
par(mfrow = c(1, 3), mar = c(4, 4, 2, 1), oma = c(0, 0, 2, 0))
with(airquality, {
    # here three plots are filled in with their respective titles
    plot(Wind, Ozone, main = "Ozone and Wind")
    plot(Solar.R, Ozone, main = "Ozone and Solar Radiation")
    plot(Temp, Ozone, main = "Ozone and Temperature")
    # this adds a line of text in the outer margin*
    mtext("Ozone and Weather in New York City", outer = TRUE)}
)
```

Ozone and Weather in New York City



Graphics Device

- A graphic device is something where you can make a plot appear
 - window on screen (screen device) = quick visualizations and exploratory analysis
 - $-\mathbf{pdf}$ (file device) = plots that may be printed out or incorporated in to document
 - PNG/JPEG (file device) = plots that may be printed out or incorporated in to document
 - scalable vector graphics (SVG)
- When a plot is created in R, it has to be sent to a graphic device
- Most common is screen device
 - quartz() on Mac, windows() on Windows, x11() on Unix/Linux
 - ?Devices = lists devices found

• Plot creation

- screen device
 - * call plot/xplot/qplot \rightarrow plot appears on screen device \rightarrow annotate as necessary \rightarrow use
- file devices
 - * explicitly call graphics device \rightarrow plotting function to make plot (write to file) \rightarrow annotate as necessary \rightarrow explicitly close graphics device with dev.off()

• Graphics File Devices

- Vector Formats (good for line drawings/plots w/ solid colors, a modest number of points)
 - * pdf: useful for line type graphics, resizes well, usually portable, not efficient if too many points
 - * svg: XML based scalable vector graphics, support animation and interactivity, web based
 - * win.metafile: Windows metafile format
 - * postscript: older format, resizes well, usually portable, can create encapsulated postscript file, Windows often don't have postscript viewer (postscript = predecessor of PDF)
- Bitmap Formats (good for plots w/ large number of points, natural scenes/webbased plots)
 - * png: Portable Network Graphics, good for line drawings/image with solid colors, uses lossless compression, most web browsers read this natively, good for plotting a lot of data points, does not resize well
 - * JPEG: good for photographs/natural scenes/gradient colors, size efficient, uses lossy compression, good for plotting many points, does not resize well, can be read by almost any computer/browser, not great for line drawings (aliasing on edges)
 - * tiff: common bitmap format supports lossless compression
 - * bmp: native Windows bitmapped format

• Multiple Open Graphics Devices

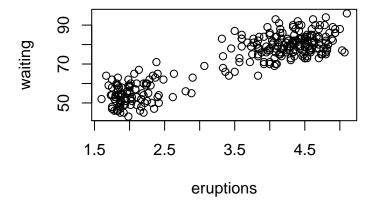
- possible to open multiple graphics devices (screen, file, or both)
- plotting occurs only one device at a time
- dev.cur() = returns the currently active device
- every open graphics device is assigned an integer ≥ 2
- dev.set(<integer>) = change the active graphics device <integer> = number associated with
 the graphics device you want to switch to

Copying plots

- dev.copy() = copy a plot from one device to another
- dev.copy2pdf() = specifically for copying to PDF files
- Note: copying a plot is not an exact operation, so the result may not be identical to the original
- example

```
## Create plot on screen device
with(faithful, plot(eruptions, waiting))
## Add a main title
title(main = "Old Faithful Geyser data")
```

Old Faithful Geyser data



```
## Copy my plot to a PNG file
dev.copy(png, file = "geyserplot.png")
## Don't forget to close the PNG device!
dev.off()
```

lattice Plotting System

- p_load("lattice") = load lattice system
- implemented using the lattice and grid packages
 - lattice package = contains code for producing *Trellis* graphics (independent from base graphics system)
 - grid package = implements the graphing system; lattice build on top of grid
- all plotting and annotation is done with single function call
 - margins/spacing/labels set automatically for entire plot, good for putting multiple on the screen
 - good for conditioning plots \rightarrow examining same plots over different conditions how y changes vs x across different levels of z
 - panel functions can be specified/customized to modify the subplots
- lattice graphics functions return an object of class "trellis", where as base graphics functions plot data directly to graphics device
 - print methods for lattice functions actually plots the data on graphics device
 - trellis objects are auto-printed
 - trellis.par.set() \rightarrow can be used to set global graphic parameters for all trellis objects
- hard to annotate, awkward to specify entire plot in one function call
- cannot add to plot once created, panel/subscript functions hard to prepare

lattice Functions and Parameters

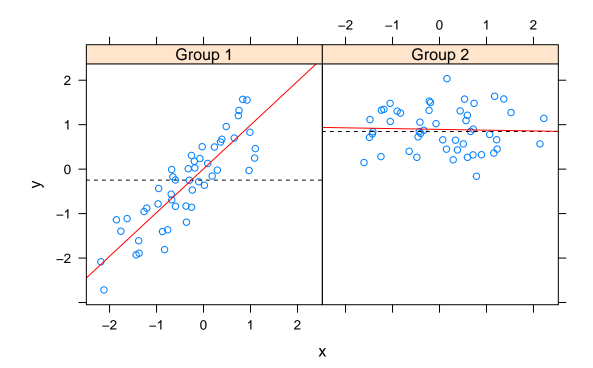
• Funtions

- xyplot() = main function for creating scatterplots
- bwplot() = box and whiskers plots (box plots)
- histogram() = histograms
- stripplot() = box plot with actual points
- dotplot() = plot dots on "violin strings"
- splom() = scatterplot matrix (like pairs() in base plotting system)
- levelplot()/contourplot() = plotting image data
- Arguments for xyplot(y ~ x | f * g, data, layout, panel)
 - default blue open circles for data points
 - formula notation is used here (\sim) = left hand side is the y-axis variable, and the right hand side is the x-axis variable
 - f/g = conditioning/categorical variables (optional)
 - * basically creates multi-panelled plots (for different factor levels)
 - * * indicates interaction between two variables
 - * intuitively, the xyplot displays a graph between x and y for every level of f and g
 - data = the data frame/list from which the variables should be looked up
 - * if nothing is passed, the parent frame is used (searching for variables in the workspace)
 - * if no other arguments are passed, defaults will be used
 - layout = specifies how the different plots will appear
 - * layout = c(5, 1) = produces 5 subplots in a horizontal fashion
 - * padding/spacing/margin automatically set
 - [optional] panel function can be added to control what is plotted inside each panel of the plot

- * panel functions receive x/y coordinates of the data points in their panel (along with any additional arguments)
- * ?panel.xyplot = brings up documentation for the panel functions
- * Note: no base plot functions can be used for lattice plots

lattice Example

```
p_load("lattice")
set.seed(10)
x <- rnorm(100)
f <- rep(0:1, each = 50)
y <- x + f - f * x+ rnorm(100, sd = 0.5)
f <- factor(f, labels = c("Group 1", "Group 2"))
## Plot with 2 panels with custom panel function
xyplot(y ~ x | f, panel = function(x, y, ...) {
    # call the default panel function for xyplot
    panel.xyplot(x, y, ...)
    # adds a horizontal line at the median
    panel.abline(h = median(y), lty = 2)
    # overlays a simple linear regression line
    panel.lmline(x, y, col = 2)
})</pre>
```



ggplot2 Plotting System

- p_load("ggplot2") = loads ggplot2 package
- implementation of Grammar of Graphics by Leland Wilkinson, written by Hadley Wickham (created RStudio)

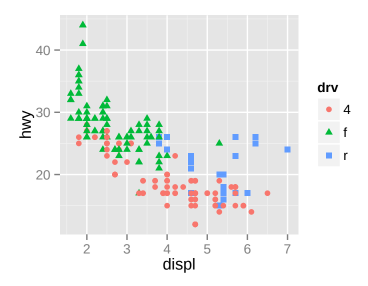
"In brief, the grammar tells us that a statistical graphic is a mapping from data to aesthetic attributes (color, shape, size) of geometric objects (points, lines, bars). The plot may also contain statistical transformations of the data and is drawn on a specific coordinate system"

- grammar graphics plot, splits the different between base and lattice systems
- automatically sets spacings/text/tiles but also allows annotations to be added
- default makes a lot of choices, but still customizable

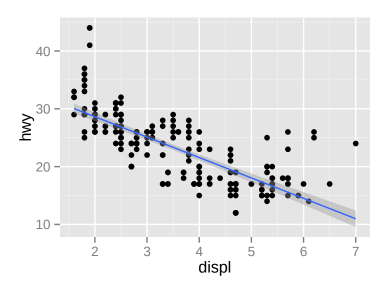
ggplot2 Functions and Parameters

- basic components of a ggplot2 graphic
 - data frame = source of data
 - aesthetic mappings = how data are mapping to color/size (x vs y)
 - **geoms** = geometric objects like points/lines/shapes to put on page
 - **facets** = conditional plots using factor variables/multiple panels
 - stats = statistical transformations like binning/quantiles/smoothing
 - scales = scale aesthetic map uses (i.e. male = red, female = blue)
 - coordinate system = system in which data are plotted
- qplot(x, y, data, color, geom) = quick plot, analogous to base system's plot() function
 - default style: gray background, white gridlines, x and y labels automatic, and solid black circles for data points
 - data always comes from data frame (in unspecified, function will look for data in workspace)
 - plots are made up of aesthetics (size, shape, color) and geoms (points, lines)
 - Note: capable of producing quick graphics, but difficult to customize in detail
- factor variables: important for graphing subsets of data = they should be labelled with specific information, and not just 1, 2, 3
 - color = factor1 = use the factor variable to display subsets of data in different colors on the same plot (legend automatically generated)
 - shape = factor2 = use the factor variable to display subsets of data in different shapes on the same plot (legend automatically generated)
 - example

```
p_load("ggplot2")
qplot(displ, hwy, data = mpg, color = drv, shape = drv)
```

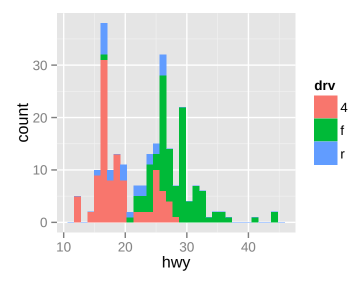


- \bullet adding statistics: geom = c("points", "smooth") = add a smoother/"low S"
 - "points" plots the data themselves, "smooth" plots a smooth mean line in blue with an area of 95% confidence interval shaded in dark gray
 - \mathtt{method} = "lm" = additional argument method can be specified to create different lines/confidence intervals
 - * lm = linear regression
 - example



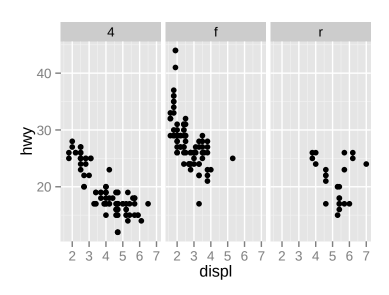
- histograms: if only one value is specified, a histogram is produced
 - fill = factor1 = can be used to fill the histogram with different colors for the subsets (legend automatically generated)

- example

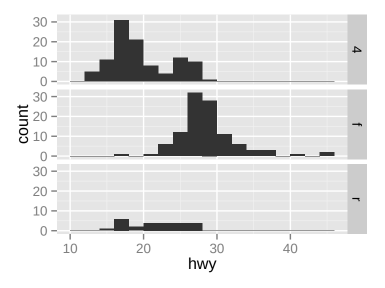


- facets: similar to panels in lattice, split data according to factor variables
 - facets = rows ~ columns = produce different subplots by factor variables specified
 (rows/columns)
 - "." indicates there are no addition row or column
 - facets = . ~ columns = creates 1 by col subplots
 - facets = row ~ . = creates row row by 1 subplots
 - labels get generated automatically based on factor variable values
 - example

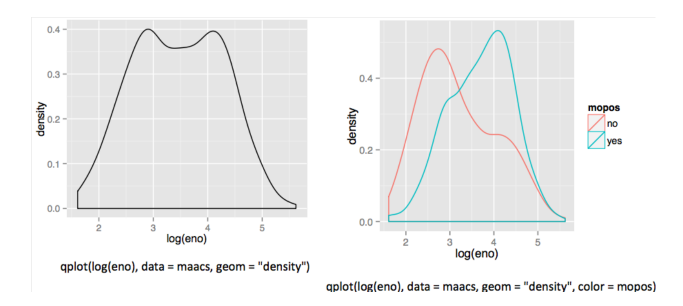
qplot(displ, hwy, data = mpg, facets = . ~ drv)



qplot(hwy, data = mpg, facets = drv ~ ., binwidth = 2)



- density smooth: smooths the histograms into a line tracing its shape
 - geom = "density" = replaces the default scatterplot with density smooth curve
 - example



- ggplot()
 - built up in layers/modularly (similar to base plotting system)
 - * data \rightarrow overlay summary \rightarrow metadata/annotation
 - g <- ggplot(data, aes(var1, var2))</pre>
 - * initiates call to ggplot and specifies the data frame that will be used

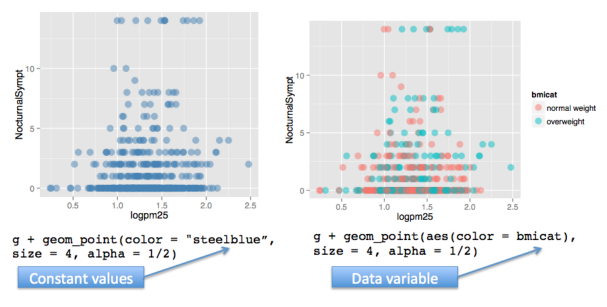
- * aes(var1, var2) = specifies aesthetic mapping, or var1 = x variable, and var2 = y variable
- * summary(g) = displays summary of ggplot object
- * print(g) = returns error ("no layer on plot") which means the plot does know how to draw the data yet
- g + geom_point() = takes information from g object and produces scatter plot
- + geom_smooth() = adds low S mean curve with confidence interval
 - * method = "lm" = changes the smooth curve to be linear regression
 - * size = 4, linetype = 3 = can be specified to change the size/style of the line
 - * se = FALSE = turns off confidence interval
- + facet_grid(row ~ col) = splits data into subplots by factor variables (see facets from qplot())
 - * conditioning on continous variables is possible through cutting/making a new categorical variable
 - * cutPts <- quantiles(df\$cVar, seq(0, 1, length=4), na.rm = TRUE) = creates quantiles where the continuous variable will be cut
 - · seq(0, 1, length=4) = creates 4 quantile points
 - · na.rm = TRUE = removes all NA values
 - * df\$newFactor <- cut(df\$cVar, cutPts) = creates new categorical/factor variable by using the cutpoints
 - \cdot creates n-1 ranges from n points = in this case 3

- annotations:

- * xlab(), ylab(), labs(), ggtitle() = for labels and titles
 - · labs(x = expression("log " * PM[2.5]), y = "Nocturnal") = specifies x and y labels
 - expression() = used to produce mathematical expressions
- * geom functions = many options to modify
- * theme() = for global changes in presentation
 - example: theme(legend.position = "none")
- * two standard themes defined: theme_gray() and theme_bw()
- * base_family = "Times" = changes font to Times

- aesthetics

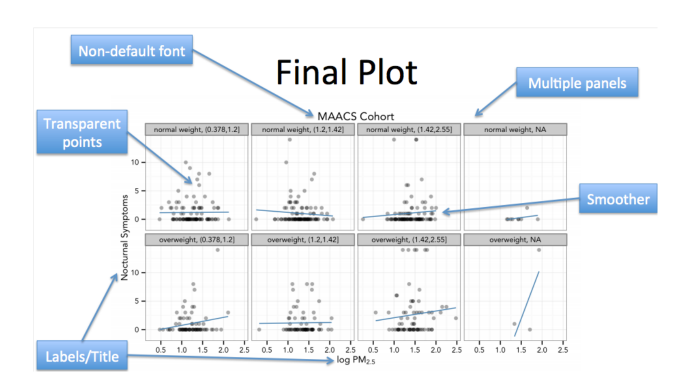
- * + geom_point(color, size, alpha) = specifies how the points are supposed to be plotted on the graph (style)
 - · Note: this translates to geom_line()/other forms of plots
 - · color = "steelblue" = specifies color of the data points
 - · aes(color = var1) = wrapping color argument this way allows a factor variable to be assigned to the data points, thus subsetting it with different colors based on factor variable values
 - \cdot size = 4 = specifies size of the data points
 - \cdot alpha = 0.5 = specifies transparency of the data points
- $*\ example$



- axis limits

- * + ylim(-3, 3) = limits the range of y variable to a specific range
 - · Note: ggplot will exclude (not plot) points that fall outside of this range (outliers), potentially leaving gaps in plot
- * + coord_cartesian(ylim(-3, 3)) = this will limit the visible range but plot all points of the data

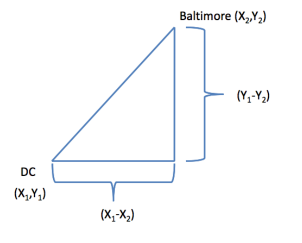
ggplot2 Comprehensive Example

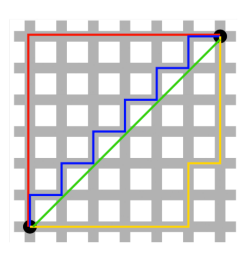


Hierarchical Clustering

- useful for visualizing high dimensional data, organizes things that are close into groups
- agglomerative approach (most common) bottom up
 - 1. start with data
 - 2. find closest pairs, put them together (create "super point" and remove original data)
 - 3. find the next closest
 - 4. repeat = yields a tree showing order of merging (dendrogram)
 - requires
 - * merging approach: how to merge two points
 - * distance metric: calculating distance between two points
 - * continuous Euclidean distance $\rightarrow \sqrt{(A_1 A_2)^2 + (B_1 B_2)^2 + \cdots + (Z_1 Z_2)^2}$
 - * continuous $correlation\ similarity o$ how correlated two data points are
 - * binary Manhattan distance ("city block distance") $\rightarrow |A_1 A_2| + |B_1 B_2| + \cdots + |Z_1 Z_2|$

Warning in readPNG("figures/25.png"): libpng warning: iCCP: known incorrect
sRGB profile





Euclidean Distance

Binary Distance

Procedure for Constructing Hierarchical Clusters (hclust function)

- 1. calculate all pair wise distances between all points to see which points are closest together
 - dist(data.frame(x=x, y=y) = returns pair wise distances for all of the (x,y) coordinates
 - Note: dist() function uses Euclidean distance by default
- 2. group two closest points from the calculated distances and merge them to a single point
- 3. find the next two closest points and merge them, and repeat
- 4. order of clustering is shown in the dendrogram

Approaches for Merging Points/Clusters

• the approach is specified in the argument method = "complete" or "average" in hclust() function

- average linkage = taking average of the x and y coordinates for both points/clusters (center of mass effectively)
- *complete linkage* = to measure distance of two clusters, take the two points in the clusters that are the furthest apart
- Note: two approaches may produce different results so it's a good idea to use both approaches to validate results

Characteristics of Hierarchical Clustering Algorithms

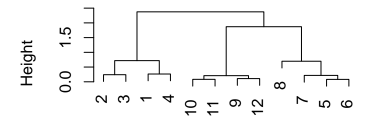
- clustering result/plot maybe *unstable*
 - changing few points/outliers could lead to large changes
 - change different distance metrics to see how sensitive the clustering is
 - change merging strategy
 - scaling of variables could affect the clustering (if one unit/measurement is much larger than another)
- deterministic = running the hclust function with same parameters and the same data will produce the same plot
- determining how many clusters there are (where to cut) may not always be clear
- primarily used for exploratory data analysis, to see over all pattern in data if there is any at all

hclust Function and Example

- hh <- hclust(dist(dataFrame)) function = produces a hierarchical cluster object based on pair wise distances from a data frame of x and y values
 - dist() = defaults to Euclidean, calculates the distance/similarity between two observations; when applied to a data frame, the function applies the $\sqrt{(A_1 A_2)^2 + (B_1 B_2)^2 + ... + (Z_1 Z_2)^2}$ formula to every pair of rows of data to construct a matrix of distances between the roes
 - * order of the hierarchical cluster is derived from the distance
 - plot(hh) = plots the dendrogram
 - automatically sorts column and row according to cluster
 - names(hh) = returns all parameters of the hclust object
 - * hh\$order = returns the order of the rows/clusters from the dendrogram
 - * hh\$dist.method = returns method for calculating distance/similarity
- Note: dendrogram that gets generated DOES NOT show how many clusters there are, so cutting (at 2.0 level for example) must be done to determine number of clusters must be a convenient and sensible point
- hclust Example

```
set.seed(1234)
x <- rnorm(12,mean=rep(1:3,each=4),sd=0.2)
y <- rnorm(12,mean=rep(c(1,2,1),each=4),sd=0.2)
dataFrame <- data.frame(x=x,y=y)
distxy <- dist(dataFrame)
hClustering <- hclust(distxy)
plot(hClustering)</pre>
```

Cluster Dendrogram



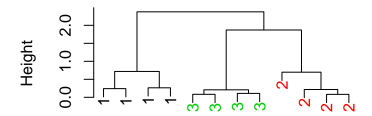
distxy hclust (*, "complete")

myplcclust Function and Example

• Note: myplcclust = a function to plot hclust objects in color (clusters labeled 1 2 3 etc.), but must know how many clusters there are initially

```
myplclust <- function(hclust, lab = hclust$labels,</pre>
    lab.col = rep(1, length(hclust$labels)), hang = 0.1, ...) {
    ## modifiction of plclust for plotting hclust objects *in colour*! Copyright
    ## Eva KF Chan 2009 Arguments: hclust: hclust object lab: a character vector
    ## of labels of the leaves of the tree lab.col: colour for the labels;
    ## NA=default device foreground colour hang: as in hclust & plclust Side
    ## effect: A display of hierarchical cluster with coloured leaf labels.
    y <- rep(hclust$height, 2)
    x <- as.numeric(hclust$merge)</pre>
    y \leftarrow y[which(x < 0)]
    x \leftarrow x[which(x < 0)]
    x \leftarrow abs(x)
    y <- y[order(x)]
    x <- x[order(x)]
    plot(hclust, labels = FALSE, hang = hang, ...)
    text(x = x, y = y[hclust$order] - (max(hclust$height) * hang), labels = lab[hclust$order],
col = lab.col[hclust\$order], srt = 90, adj = c(1, 0.5), xpd = NA, ...)
}
# example
dataFrame \leftarrow data.frame(x = x, y = y)
distxy <- dist(dataFrame)</pre>
hClustering <- hclust(distxy)
myplclust(hClustering, lab = rep(1:3, each = 4), lab.col = rep(1:3, each = 4))
```

Cluster Dendrogram



distxy hclust (*, "complete")

heatmap Function and Example

- heatmap(data.matrix) function = similar to image(t(x))
 - good for visualizing high-dimension matrix data, runs hierarchical analysis on rows and columns of table
 - yellow = high value, red = low value
 - Note: the input must be a numeric matrix, so as.matrix(data.frame) can be used to convert if necessary
- \bullet example

```
set.seed(12345)
data <- matrix(rnorm(400), nrow = 40)
heatmap(data)</pre>
```

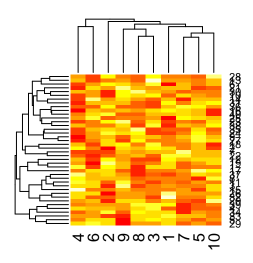
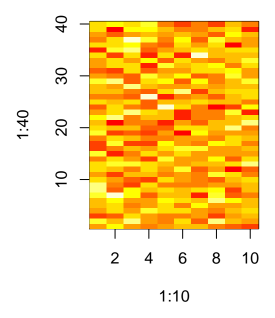


image Function and Example

- image(x, y, t(dataMatrix)[, nrow(dataMatrix):1]) = produces similar color grid plot as the heatmap() without the dendrograms
 - t(dataMatrix)[, nrow(dataMatrix)]
 - * t(dataMatrix) = transpose of dataMatrix, this is such that the plot will be displayed in the same fashion as the matrix (rows as values on the y axis and columns as values on the x axis)
 - example 40 x 10 matrix will have graph the 10 columns as x values and 40 rows as y values
 - * [, nrow(dataMatrix)] = subsets the data frame in reverse column order; when combined with the t() function, it reorders the rows of data from 40 to 1, such that the data from the matrix is displayed in order from top to bottom
 - · Note: without this statement the rows will be displayed in order from bottom to top, as that is in line with the positive y axis
 - -x, y = used to specify the values displayed on the x and y axis
 - * Note: must be in increasing order
- example

image(1:10, 1:40, t(data)[, nrow(data):1])



K-means Clustering

- similar to hierarchical clustering, focuses on finding things that are close together
 - define close, groups, visualizing/interpreting grouping

• partitioning approach

- 1. set number of clusters initially
- 2. find centroids for each cluster
- 3. assign points to the closest centroid
- 4. recalculate centroid
- 5. repeat = yields estimate of cluster centroids and which cluster each point belongs to

- requires

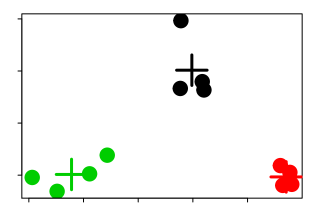
- * distance metric
- * initial number of clusters
- * initial guess as to where the cluster centroids are

Procedure for Constructing K-means Clusters (kmeans function)

- 1. choose three random points as the starting centroids
- 2. take each of the data points and assign it to the closest centroid (creating a cluster around each starting point)
- 3. take each cluster and recalculate the centroid (taking the mean) with its enclosed data points
- 4. repeat step 2 and 3 (reassign points to centroids and update centroid locations) until a stable result is achieved

• example

```
set.seed(1234)
x <- rnorm(12, mean=rep(1:3, each=4), sd=0.2)
y \leftarrow rnorm(12, mean=rep(c(1,2,1), each=4), sd=0.2)
dataFrame <- data.frame(x=x,y=y)</pre>
# specifies initial number of clusters to be 3
kmeansObj <- kmeans(dataFrame,centers=3)</pre>
names(kmeansObj)
## [1] "cluster"
                       "centers"
                                       "totss"
                                                       "withinss"
## [5] "tot.withinss" "betweenss"
                                       "size"
                                                       "iter"
## [9] "ifault"
# returns cluster assignments
kmeansObj$cluster
   [1] 3 3 3 3 1 1 1 1 2 2 2 2
par(mar=rep(0.2,4))
plot(x,y,col=kmeansObj$cluster,pch=19,cex=2)
points(kmeansObj$centers,col=1:3,pch=3,cex=3,lwd=3)
```



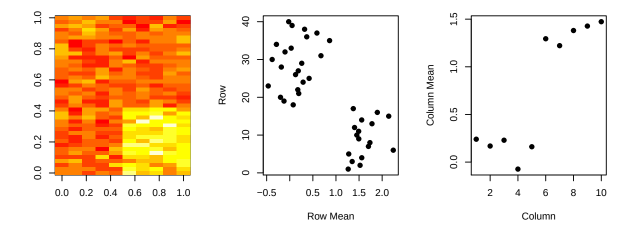
Characteristics of K-means Clustering Algorithms

- requires number of clusters initially
 - pick by eye/intuition
 - pick by cross validation/information theory, etc. [link]
- not deterministic (starting points chosen at random)
 - useful to run the algorithms a few times with different starting points to validate results

Dimension Reduction

- two kinds of problems that relate to high-dimension dataset/matrix with many variables
 - 1. find a new set (smaller) of variables that are uncorrelated and explain as much variance of data as possible
 - normally many variables are not independent (i.e. height vs weight)
 - statistical problem, commonly solved with PCA
 - 2. find a lower rank matrix (best matrix created with fewer variables) that still explains the data
 - data compression problem, commonly solved SVD
- \bullet example
 - **Note**: we are arbitrarily introduced pattern in data: we flip a coin and if the it is heads, we replace the row with [0, 0, 0, 0, 3, 3, 3, 3, 3]
 - here we plot the patterns in rows and columns (already sorted)

```
for(i in 1:40){
  # flip a coin
  coinFlip <- rbinom(1,size=1,prob=0.5)</pre>
  # if coin is heads add a common pattern to that row
  if(coinFlip){
    data[i,] \leftarrow data[i,] + rep(c(0,3), each=5)
  }
# hierarchical clustering
hh <- hclust(dist(data))</pre>
dataOrdered <- data[hh$order,]</pre>
# create 1 x 3 panel plot
par(mfrow=c(1,3))
# heat map (sorted)
image(t(dataOrdered)[,nrow(dataOrdered):1])
# row means (40 rows)
plot(rowMeans(dataOrdered),40:1,,xlab="Row Mean",ylab="Row",pch=19)
# column means (10 columns)
plot(colMeans(dataOrdered),xlab="Column",ylab="Column Mean",pch=19)
```



Singular Value Decomposition (SVD)

- Let X = matrix which each variable in column (measurement) and each observation in row (subject)
- SVD in this case is a **matrix decomposition** process, in which X is divided into *three* separate matrices as follows:

$$X = UDV^T$$

- -U = left singular vector, orthogonal matrix (columns independent of each other)
- -D = singular values, diagonal matrix
- -V = right singular vector, orthogonal matrix (columns independent of each other)
- **Note**: orthogonal implies that a matrix is always invertible $[A^{-1} = A^T]$ and that the product of the matrix and its transpose equals the identity matrix $[AA^T = I]$
 - * when a orthogonal matrices, A, is multiplied by another matrix, B, it is effectively a linear transformation in that the length and angles of B are preserved
- **Note**: diagonal implies that any value outside of the main diagonal $(\searrow) = 0$
 - * example

$$A = \begin{bmatrix} 1 & 0 & 0 \\ 0 & 2 & 0 \\ 0 & 0 & 3 \end{bmatrix}$$

• Note: scale of data matters for SVD/PCA (scaling the data may help), patterns detected maybe mixed together, and computation is intensive for these operations

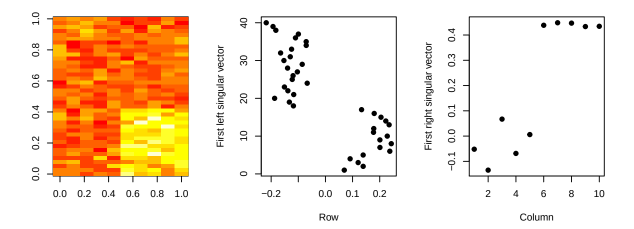
Principal Components Analysis (PCA)

- first scale the variables and run SVD on normalized matrix
 - scaling = subtract each column by its mean and divide by its standard deviation
- principal components = the right singular values or the V matrix

SVD and PCA Example

- \bullet U and V Matrices
 - $-s \leftarrow svd(data) = performs SVD on data (n \times m matrix) and splits it into u, v, and d matrices$
 - * su = n \times m \text{ matrix} \rightarrow \text{horizontal variation}$
 - * s\$d = 1 × m vector \rightarrow vector of the singular/diagonal values
 - · diag(s\$d) = $m \times m$ diagonal matrix
 - * sv = m \times m \text{ matrix} \rightarrow \text{vertical variation}$
 - * s\$u %*% diag(s\$d) %*% t(s\$v) = returns the original data $\rightarrow X = UDV^T$
 - scale(data) = scales the original data by subtracting each data point by its column mean and dividing by its column standard deviation

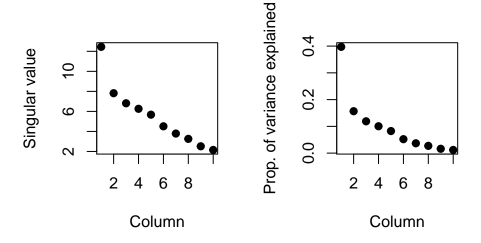
```
# running svd
svd1 <- svd(scale(dataOrdered))
# create 1 by 3 panel plot
par(mfrow=c(1,3))
# data heatmap (sorted)
image(t(dataOrdered)[,nrow(dataOrdered):1])
# U Matrix - first column
plot(svd1$u[,1],40:1,,xlab="Row",ylab="First left singular vector",pch=19)
# V vector - first column
plot(svd1$v[,1],xlab="Column",ylab="First right singular vector",pch=19)</pre>
```



ullet D Matrix and Variance Explained

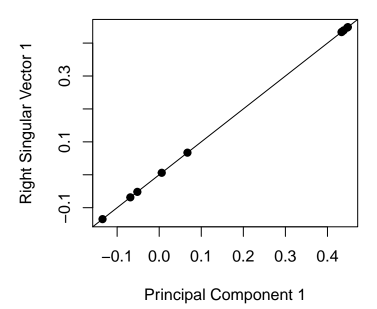
- d matrix (s\$d vector) captures the singular values, or variation in data that is explained by that particular component (variable/column/dimension)
- **proportion of variance Explained** = converting the singular values to variance (square the values) and divide by the total variance (sum of the squared singular values)
 - * effectively the same pattern as the singular values, just converted to percentage
 - * in this case, the first component/dimension, which captures the shift in means (see previous plot) of SVD captures about 40% of the variation

```
# create 1 x 2 panel plot
par(mfrow=c(1,2))
# plot singular values
plot(svd1$d,xlab="Column",ylab="Singular value",pch=19)
# plot proportion of variance explained
plot(svd1$d^2/sum(svd1$d^2),xlab="Column",ylab="Prop. of variance explained",pch=19)
```



• Relationship to PCA

- p <- prcomp(data, scale = TRUE) = performs PCA on data specified
 - * scale = TRUE = scales the data before performing PCA
 - * returns prcomp object
 - * summary(p) = prints out the principal component's standard deviation, proportion of variance, and cumulative proportion
- PCA's rotation vectors are equivalent to their counterparts in the V matrix from the SVD



```
# summarize PCA
summary(pca1)
```

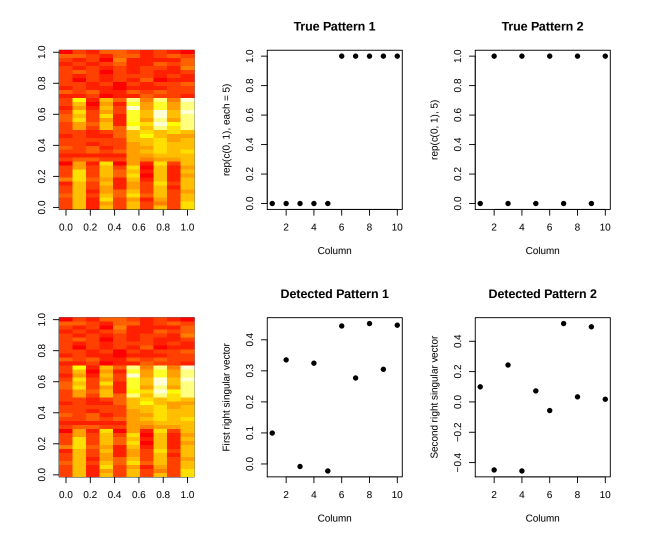
```
## Importance of components:
##
                             PC1
                                    PC2
                                            PC3
                                                   PC4
                                                           PC5
                                                                   PC6
                                                                            PC7
                          1.9930 1.2518 1.0905 1.0024 0.90836 0.72211 0.60630
## Standard deviation
## Proportion of Variance 0.3972 0.1567 0.1189 0.1005 0.08251 0.05214 0.03676
## Cumulative Proportion 0.3972 0.5539 0.6728 0.7733 0.85582 0.90797 0.94473
##
                              PC8
                                      PC9
                                              PC10
```

```
## Standard deviation 0.52145 0.40286 0.34423
## Proportion of Variance 0.02719 0.01623 0.01185
## Cumulative Proportion 0.97192 0.98815 1.00000
```

• More Complex Patterns

- SVD can be used to *detect unknown patterns* within the data (we rarely know the true distribution/pattern about the population we're analyzing)
- however, it may be hard to pinpoint exact patterns as the principal components may confound each other
 - * in the example below, you can see that the two principal components that capture the most variation have both horizontal shifts and alternating patterns captured in them

```
set.seed(678910)
# setting pattern
data <- matrix(rnorm(400), nrow = 40)</pre>
for(i in 1:40){
  # flip a coin
  coinFlip1 <- rbinom(1,size=1,prob=0.5)</pre>
  coinFlip2 <- rbinom(1,size=1,prob=0.5)</pre>
  # if coin is heads add a common pattern to that row
  if(coinFlip1){
    data[i,] \leftarrow data[i,] + rep(c(0,5), each=5)
  }
  if(coinFlip2){
    data[i,] \leftarrow data[i,] + rep(c(0,5),5)
  }
}
hh <- hclust(dist(data)); dataOrdered <- data[hh$order,]</pre>
# perform SVD
svd2 <- svd(scale(dataOrdered))</pre>
par(mfrow=c(2,3))
image(t(dataOrdered)[,nrow(dataOrdered):1])
plot(rep(c(0,1),each=5),pch=19,xlab="Column", main="True Pattern 1")
plot(rep(c(0,1),5),pch=19,xlab="Column",main="True Pattern 2")
image(t(dataOrdered)[,nrow(dataOrdered):1])
plot(svd2$v[,1],pch=19,xlab="Column",ylab="First right singular vector",
    main="Detected Pattern 1")
plot(svd2$v[,2],pch=19,xlab="Column",ylab="Second right singular vector",
    main="Detected Pattern 2")
```

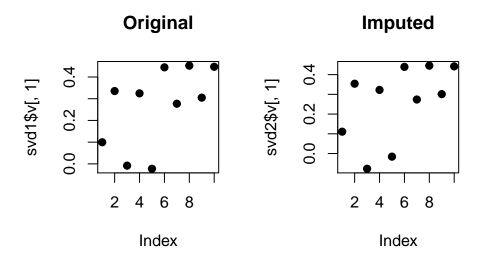


• Missing Data

- SVD cannot be performed on dataset with NA values
- impute package from Bioconductor can help approximate missing values from surrounding values
 - * impute.knn function takes the missing row and imputes the data using the k nearest neighbors to that row
 - \cdot k=10 = default value (take the nearest 10 rows)

```
## The "impute" library is available from http://bioconductor.org
## biocLite() is the recommended way to install Bioconductor packages.
## If the "impute" library is not already installed, you can uncomment
## the following three lines to install it from Bioconductor.
#source("http://bioconductor.org/biocLite.R")
#biocLite() # install core packages
#biocLite(c("impute"), suppressUpdates=TRUE) # install "impute" library
library(impute)
data2 <- dataOrdered
# set random samples = NA
data2[sample(1:100,size=40,replace=FALSE)] <- NA</pre>
```

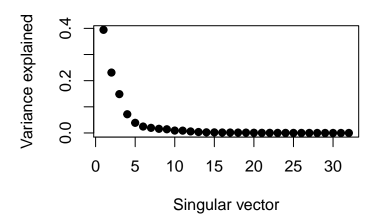
```
data2 <- impute.knn(data2)$data
svd1 <- svd(scale(data0rdered)); svd2 <- svd(scale(data2))
par(mfrow=c(1,2))
plot(svd1$v[,1],pch=19, main="Original")
plot(svd2$v[,1],pch=19, main="Imputed")</pre>
```



Create Approximations/Data Compression

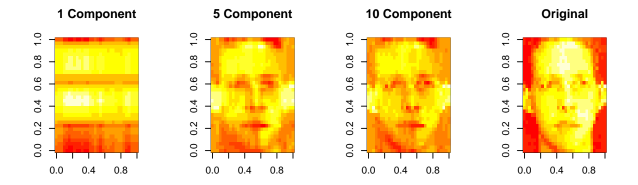
- SVD can be used to create lower rank representation, or compressed representation of data
- ullet if we look at the variance explained plot below, most of the variation is explained by the first few principal components

```
# load faceData
load("figures/face.rda")
# perform SVD
svd3 <- svd(scale(faceData))
plot(svd3$d^2/sum(svd3$d^2),pch=19,xlab="Singular vector",ylab="Variance explained")</pre>
```



• approximations can thus be created by taking the first few components and using matrix multiplication with the corresponding U, V, and D components

```
approx1 <- svd3$u[,1] %*% t(svd3$v[,1]) * svd3$d[1]
approx5 <- svd3$u[,1:5] %*% diag(svd3$d[1:5])%*% t(svd3$v[,1:5])
approx10 <- svd3$u[,1:10] %*% diag(svd3$d[1:10])%*% t(svd3$v[,1:10])
# create 1 x 4 panel plot
par(mfrow=c(1,4))
# plot original facedata
image(t(approx1)[,nrow(approx1):1], main = "1 Component")
image(t(approx5)[,nrow(approx5):1], main = "5 Component")
image(t(approx10)[,nrow(approx10):1], main = "10 Component")
image(t(faceData)[,nrow(faceData):1], main = "0riginal")</pre>
```



Color Packages in R Plots

- proper use of color can help convey the message by improving clarity/contrast of data presented
- default color schemes for most plots in R are fairly terrible, so some external packages are helpful

grDevices Package

- colors() function = lists names of colors available in any plotting function
- colorRamp function
 - takes any set of colors and return a function that takes values between 0 and 1, indicating the extremes of the color palette (e.g. see the gray function)
 - pal <- colorRamp(c("red", "blue")) = defines a colorRamp function
 - pal(0) returns a 1 x 3 matrix containing values for RED, GREEN, and BLUE values that range from 0 to $255\,$
 - pal(seq(0, 1, len = 10)) returns a 10 x 3 matrix of 10 colors that range from RED to BLUE (two ends of spectrum defined in the object)
 - example

```
# define colorRamp function
pal <- colorRamp(c("red", "blue"))
# create a color
pal(0.67)</pre>
```

```
## [,1] [,2] [,3]
## [1,] 84.15 0 170.85
```

• colorRampPalette function

- takes any set of colors and return a function that takes integer arguments and returns a vector of colors interpolating the palette (like heat.colors or topo.colors)
- pal <- colorRampPalette(c("red", "yellow")) defines a colorRampPalette function
- pal(10) returns 10 interpolated colors in hexadecimal format that range between the defined ends of spectrum
- example

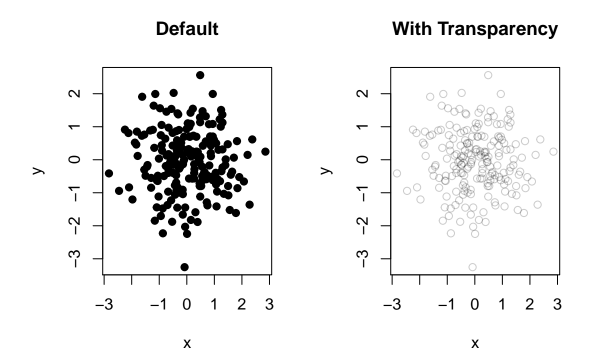
```
# define colorRampPalette function
pal <- colorRampPalette(c("red", "yellow"))
# create 10 colors
pal(10)</pre>
```

```
## [1] "#FF0000" "#FF1C00" "#FF3800" "#FF5500" "#FF7100" "#FF8D00" "#FFAA00"
## [8] "#FFC600" "#FFE200" "#FFFF00"
```

• rgb function

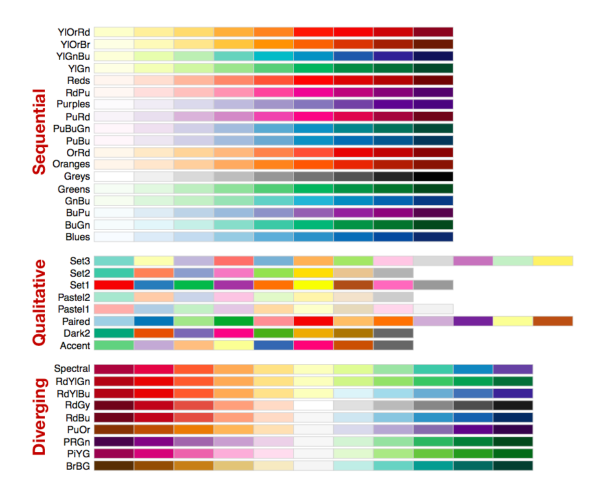
- red, green, and blue arguments = values between 0 and 1
- alpha = 0.5 = transparency control, values between 0 and 1
- returns hexadecimal string for color that can be used in plot/image commands
- colorspace package cna be used for different control over colors
- example

```
x <- rnorm(200); y <- rnorm(200)
par(mfrow=c(1,2))
# normal scatter plot
plot(x, y, pch = 19, main = "Default")
# using transparency shows data much better
plot(x, y, col = rgb(0, 0, 0, 0.2), main = "With Transparency")</pre>
```



RColorBrewer Package

- can be found on CRAN that has predefined color palettes
 - p_load("RColorBrewer")
- types of palettes
 - Sequential = numerical/continuous data that is ordered from low to high
 - Diverging = data that deviate from a value, increasing in two directions (i.e. standard deviations from the mean)
 - Qualitative = categorical data/factor variables
- ullet palette information from the RColorBrewer package can be used by colorRamp and colorRampPalette functions
- available colors palettes

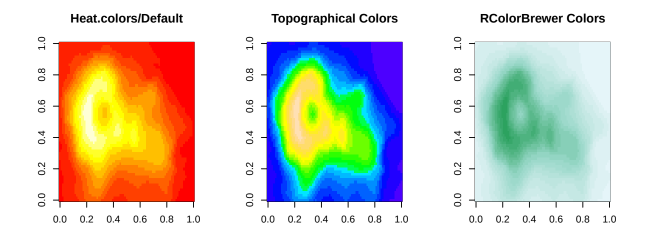


• brewer.pal(n, "BuGn") function

- n = number of colors to generated
- "BuGn" = name of palette
 - * ?brewer.pal list all available palettes to use
- returns list of n hexadecimal colors

\bullet example

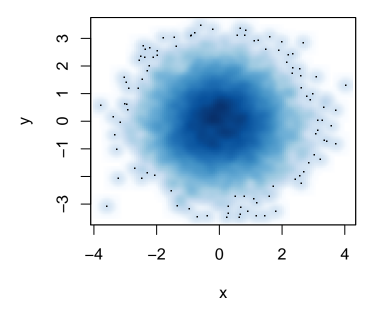
```
p_load("RColorBrewer")
# generate 3 colors using brewer.pal function
cols <- brewer.pal(3, "BuGn")
pal <- colorRampPalette(cols)
par(mfrow=c(1,3))
# heat.colors/default
image(volcano, main = "Heat.colors/Default")
# topographical colors
image(volcano, col = topo.colors(20), main = "Topographical Colors")
# RColorBrewer colors
image(volcano, col = pal(20), main = "RColorBrewer Colors")</pre>
```



• smoothScatter function

- used to plot large quantities of data points
- creates 2D histogram of points and plots the histogram
- default color scheme = "Blues" palette from RColorBrewer package
- example

```
x <- rnorm(10000); y <- rnorm(10000)
smoothScatter(x, y)</pre>
```



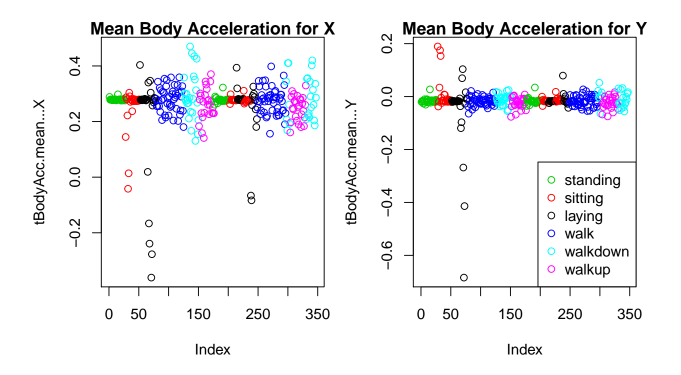
Case Study: Human Activity Tracking with Smart Phones

Loading Training Set of Samsung S2 Data from UCI Repository

```
# load data frame provided
load("samsungData.rda")
# table of 6 types of activities
table(samsungData$activity)
##
##
             sitting standing
                                   walk walkdown
                                                    walkup
     laying
                1286
                                                      1073
##
       1407
                          1374
                                   1226
                                              986
```

Plotting Average Acceleration for First Subject

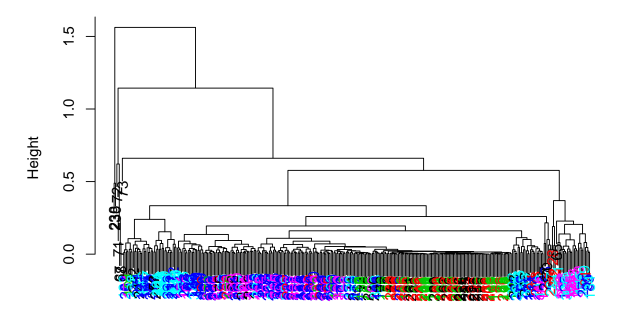
```
# set up 1 x 2 panel plot
par(mfrow=c(1, 2), mar = c(5, 4, 1, 1))
# converts activity to a factor variable
samsungData <- transform(samsungData, activity = factor(activity))
# find only the subject 1 data
sub1 <- subset(samsungData, subject == 1)
# plot mean body acceleration in X direction
plot(sub1[, 1], col = sub1$activity, ylab = names(sub1)[1],
    main = "Mean Body Acceleration for X")
# plot mean body acceleration in Y direction
plot(sub1[, 2], col = sub1$activity, ylab = names(sub1)[2],
    main = "Mean Body Acceleration for Y")
# add legend
legend("bottomright",legend=unique(sub1$activity),col=unique(sub1$activity), pch = 1)</pre>
```



Clustering Based on Only Average Acceleration

```
# load myplclust function
source("myplclust.R")
# calculate distance matrix
distanceMatrix <- dist(sub1[,1:3])
# form hclust object
hclustering <- hclust(distanceMatrix)
# run myplclust on data
myplclust(hclustering, lab.col = unclass(sub1$activity))</pre>
```

Cluster Dendrogram



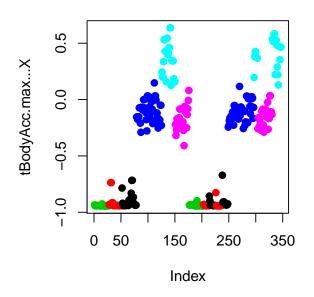
distanceMatrix hclust (*, "complete")

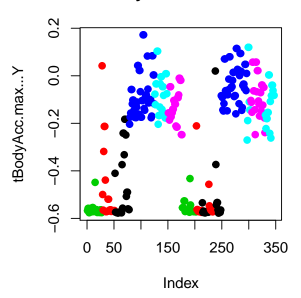
Plotting Max Acceleration for the First Subject

```
# create 1 x 2 panel
par(mfrow=c(1,2))
# plot max accelecrations in x and y direction
plot(sub1[,10],pch=19,col=sub1$activity,ylab=names(sub1)[10],
    main = "Max Body Acceleration for X")
plot(sub1[,11],pch=19,col = sub1$activity,ylab=names(sub1)[11],
    main = "Max Body Acceleration for Y")
```



Max Body Acceleration for Y

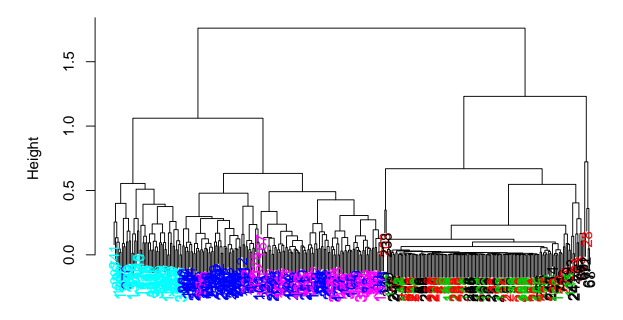




Clustering Based on Maximum Acceleration

```
# calculate distance matrix for max distances
distanceMatrix <- dist(sub1[,10:12])
hclustering <- hclust(distanceMatrix)
myplclust(hclustering,lab.col=unclass(sub1$activity))</pre>
```

Cluster Dendrogram



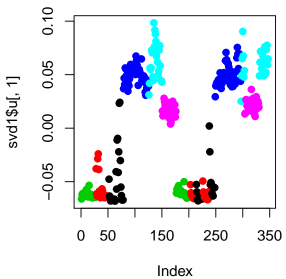
distanceMatrix hclust (*, "complete")

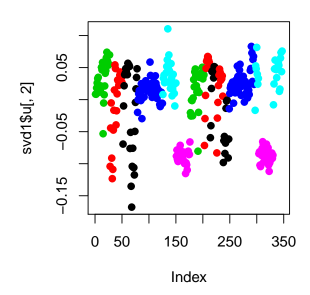
Singular Value Decomposition

```
# perform SVD minus last two columns (subject and activity)
svd1 = svd(scale(sub1[,-c(562,563)]))
# create 1 x 2 panel plot
par(mfrow=c(1,2))
# plot first two left singular vector
# separate moving from non moving
plot(svd1$u[,1],col=sub1$activity,pch=19, main = "First Left Singular Vector")
plot(svd1$u[,2],col=sub1$activity,pch=19, main = "Second Left Singular Vector")
```



Second Left Singular Vector

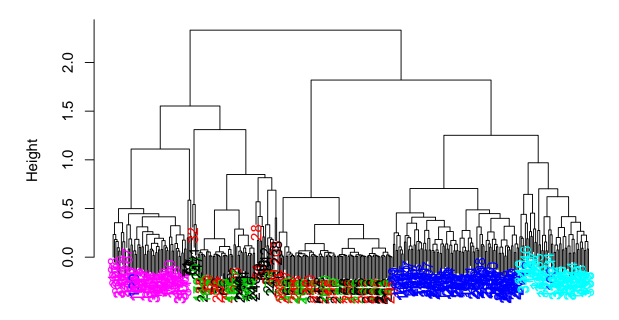




New Clustering with Maximum Contributers

```
# find the max contributing feature
maxContrib <- which.max(svd1$v[,2])
# recalculate distance matrix
distanceMatrix <- dist(sub1[, c(10:12,maxContrib)])
hclustering <- hclust(distanceMatrix)
myplclust(hclustering,lab.col=unclass(sub1$activity))</pre>
```

Cluster Dendrogram



distanceMatrix hclust (*, "complete")

```
# name of max contributing factor
names(samsungData)[maxContrib]
```

```
## [1] "fBodyAcc.meanFreq...Z"
```

K-means Clustering (nstart=1, first try)

```
# specify 6 centers for data
kClust <- kmeans(sub1[,-c(562,563)],centers=6)
# tabulate 6 clusteres against 6 activity but many clusters contain multiple activities
table(kClust$cluster,sub1$activity)</pre>
```

```
##
##
       laying sitting standing walk walkdown walkup
                     0
                                    95
##
             0
            16
                    12
                               7
                                     0
                                                      0
##
     2
                                               0
                    33
##
     3
            24
                              46
                                               0
                                                      0
##
     4
            10
                     2
                               0
                                     0
                                               0
                                                      0
                     0
                               0
                                                      0
##
     5
                                     0
                                              49
                                               0
                                                     53
##
```

K-means clustering (nstart=100, first try)

```
# run k-means algorithm 100 times
kClust <- kmeans(sub1[,-c(562,563)],centers=6,nstart=100)
# tabulate results
table(kClust$cluster,sub1$activity)</pre>
```

```
##
##
       laying sitting standing walk walkdown walkup
##
     1
             0
                     37
                              51
                                     0
                                               0
             3
                     0
                               0
                                     0
                                               0
                                                     53
##
     2
##
     3
            18
                     10
                               2
                                     0
                                               0
                                                      0
                                                      0
##
             0
                     0
                               0
                                     0
                                              49
     4
                     0
                               0
                                     0
                                               0
                                                      0
##
     5
            29
                     0
                               0
                                    95
                                               0
                                                      0
##
     6
             0
```

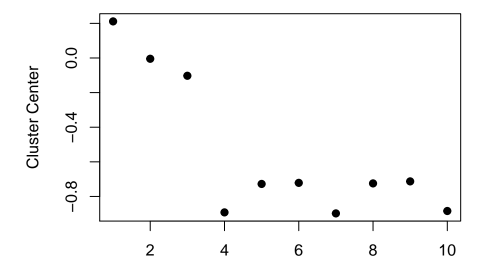
K-means clustering (nstart=100, second try)

```
# run k-means algorithm 100 times
kClust <- kmeans(sub1[,-c(562,563)],centers=6,nstart=100)
# tabulate results
table(kClust$cluster,sub1$activity)</pre>
```

```
##
##
       laying sitting standing walk walkdown walkup
##
     1
            18
                     10
                               2
                                     0
                                               0
                                                       0
     2
            0
                     0
                               0
                                     0
                                              49
                                                       0
##
##
     3
            29
                     0
                               0
                                     0
                                               0
                                                       0
                                               0
                                                       0
##
     4
             0
                     37
                              51
                                     0
                                                       0
##
     5
             0
                     0
                               0
                                    95
                                               0
##
     6
             3
                     0
                               0
                                     0
                                               0
                                                     53
```

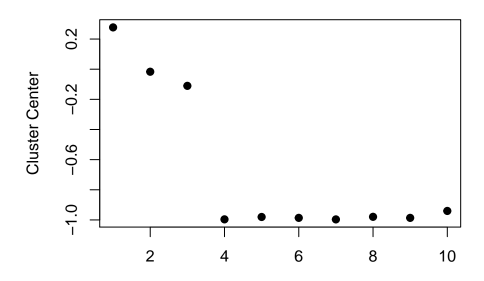
Cluster 1 Variable Centers (Laying)

```
# plot first 10 centers of k-means for laying to understand which features drive the activity plot(kClust$center[1,1:10],pch=19,ylab="Cluster Center",xlab="")
```



Cluster 2 Variable Centers (Walking)

plot first 10 centers of k-means for laying to understand which features drive the activity $\protect\operatorname{plot}(kClust\protects(4,1:10],pch=19,ylab="Cluster Center",xlab="")$



Case Study: Fine Particle Pollution in the U.S. from 1999 to 2012

Read Raw Data from 1999 and 2012

```
# read in raw data from 1999
pm0 <- read.table("pm25_data/RD_501_88101_1999-0.txt", comment.char = "#", header = FALSE, sep = "|", n
# read in headers/column lables
cnames <- readLines("pm25_data/RD_501_88101_1999-0.txt", 1)</pre>
# convert string into vector
cnames <- strsplit(substring(cnames, 3), "|", fixed = TRUE)</pre>
# make vector the column names
names(pm0) <- make.names(cnames[[1]])</pre>
# we are interested in the pm2.5 readings in the "Sample. Value" column
x0 <- pm0$Sample.Value
# read in the data from 2012
pm1 <- read.table("pm25_data/RD_501_88101_2012-0.txt", comment.char = "#", header = FALSE, sep = "|",
    na.strings = "", nrow = 1304290)
# make vector the column names
names(pm1) <- make.names(cnames[[1]])</pre>
# take the 2012 data for pm2.5 readings
x1 <- pm1$Sample.Value
```

Summaries for Both Periods

```
# generate 6 number summaries
summary(x1)
##
     Min. 1st Qu. Median
                            Mean 3rd Qu.
                                            Max.
                                                   NA's
## -10.00
           4.00
                    7.63
                            9.14 12.00 909.00
                                                  73133
summary(x0)
##
     Min. 1st Qu. Median Mean 3rd Qu.
                                            Max.
                                                   NA's
##
           7.20 11.50 13.74 17.90 157.10 13217
# calculate % of missing values, Are missing values important here?
data.frame(NA.1990 = mean(is.na(x0)), NA.2012 = mean(is.na(x1)))
##
      NA.1990
                 NA.2012
## 1 0.1125608 0.05607125
```

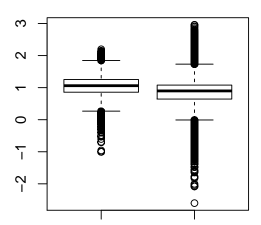
Make a boxplot of both 1999 and 2012

```
par(mfrow = c(1,2))
# regular boxplot, data too right skewed
boxplot(x0, x1, main = "Regular Boxplot")
# log boxplot, significant difference in means, but more spread
boxplot(log10(x0), log10(x1), main = "log Boxplot")
```

Regular Boxplot

200 400 600 800

log Boxplot

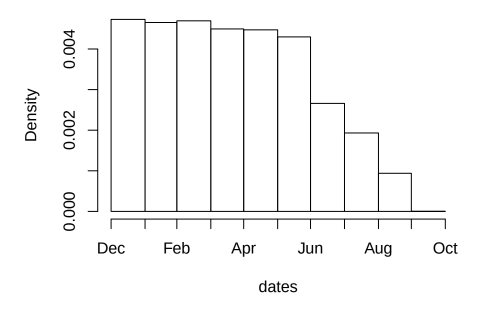


Check for Negative Values in 'x1'

0

```
# summary again
summary(x1)
      Min. 1st Qu. Median
                              Mean 3rd Qu.
                                               Max.
                                                       NA's
   -10.00
              4.00
                                     12.00 909.00
                                                      73133
                      7.63
                              9.14
# create logical vector for
negative <- x1 < 0
# count number of negatives
sum(negative, na.rm = T)
## [1] 26474
# calculate percentage of negatives
mean(negative, na.rm = T)
## [1] 0.0215034
# capture the date data
dates <- pm1$Date
dates <- as.Date(as.character(dates), "%Y%m%d")</pre>
# plot the histogram
hist(dates, "month") ## Check what's going on in months 1--6
```

Histogram of dates



Check Same New York Monitors at 1999 and 2012

```
# find unique monitors in New York in 1999
site0 <- unique(subset(pm0, State.Code == 36, c(County.Code, Site.ID)))
# find unique monitors in New York in 2012
site1 <- unique(subset(pm1, State.Code == 36, c(County.Code, Site.ID)))
# combine country codes and siteIDs of the monitors
site0 <- paste(site0[,1], site0[,2], sep = ".")
site1 <- paste(site1[,1], site1[,2], sep = ".")
# find common monitors in both
both <- intersect(site0, site1)
# print common monitors in 1999 and 2012
print(both)</pre>
```

```
## [1] "1.5" "1.12" "5.80" "13.11" "29.5" "31.3" "63.2008"
## [8] "67.1015" "85.55" "101.3"
```

Find how many observations available at each monitor

```
# add columns for combined county/site for the original data
pm0$county.site <- with(pm0, paste(County.Code, Site.ID, sep = "."))
pm1$county.site <- with(pm1, paste(County.Code, Site.ID, sep = "."))
# find subsets where state = NY and county/site = what we found previously
cnt0 <- subset(pm0, State.Code == 36 & county.site %in% both)
cnt1 <- subset(pm1, State.Code == 36 & county.site %in% both)
# split data by the county/size values and count oberservations
sapply(split(cnt0, cnt0$county.site), nrow)</pre>
```

```
29.5
                                              31.3
                                                      5.80 63.2008 67.1015
##
      1.12
              1.5
                    101.3
                             13.11
##
       61
               122
                     152
                                61
                                        61
                                               183
                                                        61
                                                               122
                                                                       122
##
     85.55
##
        7
sapply(split(cnt1, cnt1$county.site), nrow)
      1.12
                                      29.5
                                                      5.80 63.2008 67.1015
##
               1.5
                     101.3
                             13.11
                                              31.3
##
        31
                64
                        31
                                31
                                        33
                                                15
                                                        31
                                                                30
     85.55
##
##
       31
```

Choose Monitor where County = 63 and Side ID = 2008

```
# filter data by state/county/siteID
pm1sub <- subset(pm1, State.Code == 36 & County.Code == 63 & Site.ID == 2008)
pm0sub <- subset(pm0, State.Code == 36 & County.Code == 63 & Site.ID == 2008)
# there are 30 observations from 2012, and 122 from 1999
dim(pm1sub)</pre>
```

[1] 30 29

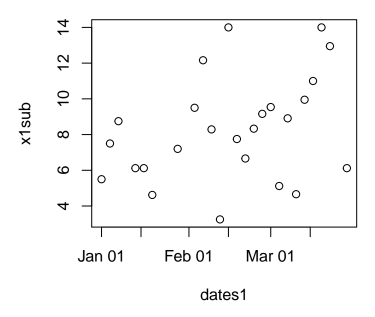
```
dim(pm0sub)
```

[1] 122 29

Plot Data for 2012

```
# capture the dates of the subset of data
dates1 <- pm1sub$Date
# capture measurements for the subset of data
x1sub <- pm1sub$Sample.Value
# convert dates to appropriate format
dates1 <- as.Date(as.character(dates1), "%Y%m%d")
# plot pm2.5 value vs time
plot(dates1, x1sub, main = "PM2.5 Polution Level in 2012")</pre>
```

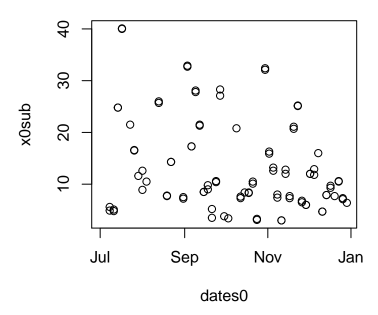
PM2.5 Polution Level in 2012



Plot data for 1999

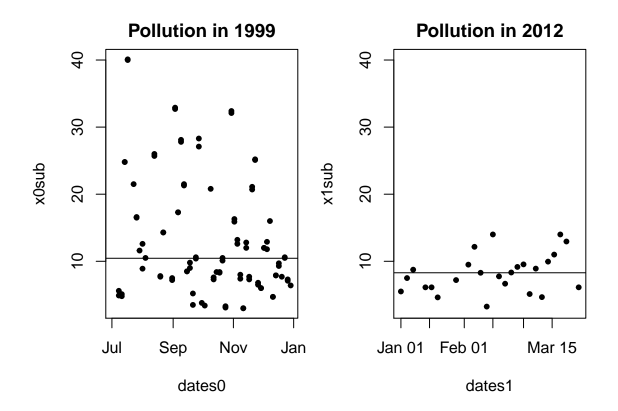
```
# capture the dates of the subset of data
dates0 <- pm0sub$Date
# convert dates to appropriate format
dates0 <- as.Date(as.character(dates0), "%Y%m%d")
# capture measurements for the subset of data
x0sub <- pm0sub$Sample.Value
# plot pm2.5 value vs time
plot(dates0, x0sub, main = "PM2.5 Polution Level in 1999")</pre>
```

PM2.5 Polution Level in 1999



Panel Plot for Both Years

```
# find max range for data
rng <- range(x0sub, x1sub, na.rm = T)
# create 1 x 2 panel plot
par(mfrow = c(1, 2), mar = c(4, 4, 2, 1))
# plot time series plot for 1999
plot(dates0, x0sub, pch = 20, ylim = rng, main="Pollution in 1999")
# plot the median
abline(h = median(x0sub, na.rm = T))
# plot time series plot for 2012
plot(dates1, x1sub, pch = 20, ylim = rng, main="Pollution in 2012")
# plot the median
abline(h = median(x1sub, na.rm = T))</pre>
```



Find State-wide Means and Trend

```
# divide data by state and find the mean of pollution level for 1999
mn0 <- with(pm0, tapply(Sample.Value, State.Code, mean, na.rm = T))
# divide data by state and find the mean of pollution level for 1999
mn1 <- with(pm1, tapply(Sample.Value, State.Code, mean, na.rm = T))
# convert to data frames while preserving state names
d0 <- data.frame(state = names(mn0), mean = mn0)
d1 <- data.frame(state = names(mn1), mean = mn1)
# merge the 1999 and 2012 means by state
mrg <- merge(d0, d1, by = "state")
# dimension of combined data frame
dim(mrg)</pre>
```

[1] 52 3

```
# first few lines of data
head(mrg)
```

```
## state mean.x mean.y
## 1 1 19.956391 10.126190
## 2 10 14.492895 11.236059
## 3 11 15.786507 11.991697
## 4 12 11.137139 8.239690
## 5 13 19.943240 11.321364
## 6 15 4.861821 8.749336
```

```
# plot the pollution levels data points for 1999
with(mrg, plot(rep(1, 52), mrg[, 2], xlim = c(.8, 2.2), ylim = c(3, 20),
    main = "PM2.5 Pollution Level by State for 1999 & 2012",
    xlab = "", ylab = "State-wide Mean PM"))
# plot the pollution levels data points for 2012
with(mrg, points(rep(2, 52), mrg[, 3]))
# connected the dots
segments(rep(1, 52), mrg[, 2], rep(2, 52), mrg[, 3])
# add 1999 and 2012 labels
axis(1, c(1, 2), c("1999", "2012"))
```

PM2.5 Pollution Level by State for 1999 & 2012

