Day 4 - Exponential Family

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October 7, 2020

More advanced plotting, introductory simulations

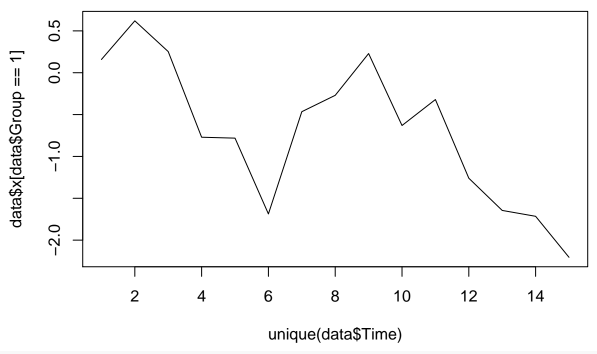
```
setwd('~/QPMR/classwork/day4Exp')
```

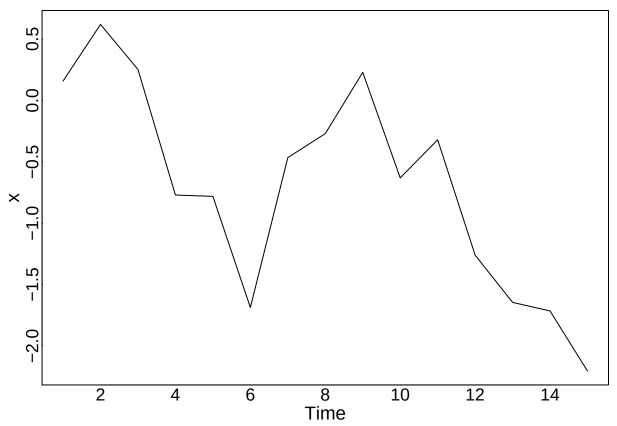
Simulate TSCS data

```
# lets create a function to generate one tscs sample with auto-correlation
tscsGen = function(rho, n.groups = 15, n.obs = 15, seed = 1523){
  set.seed(seed)
  returnMat = expand.grid(1:n.groups, 1:n.obs)
  colnames(returnMat) = c('Group', 'Time')
  returnMat$x = NA
  returnMat$x[returnMat$Time == 1] = rnorm(n.groups)
  err = numeric(n.obs*n.groups)
  err[returnMat$Time == 1] = rnorm(n.groups)
 for(g in 1:n.groups){
   for(t in 2:n.obs){
     returnMat$x[returnMat$Group == g & returnMat$Time == t] = rho * returnMat$x[returnMat$Group == g
      err[returnMat$Group == g & returnMat$Time == t] = rho * err[returnMat$Group == g & returnMat$Time
   }
  }
  err = scale(err)
  g.ef = rnorm(n.groups)
  returnMat$y = -1 + rep(g.ef, times = n.groups) + .4*returnMat$x + err
  return(returnMat)
```

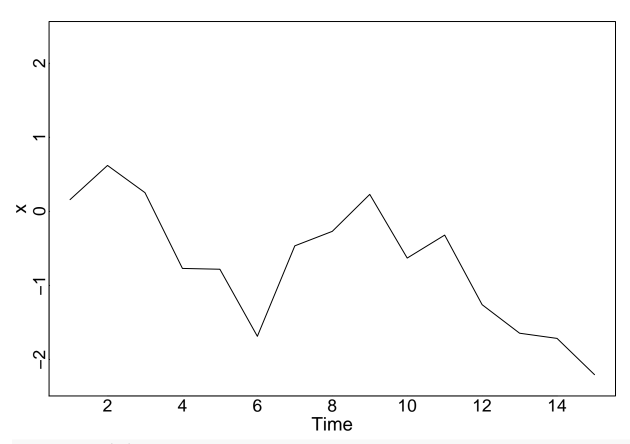
Generate one TSCS dataset at $\rho = 0.8$ and plot the variable relationships

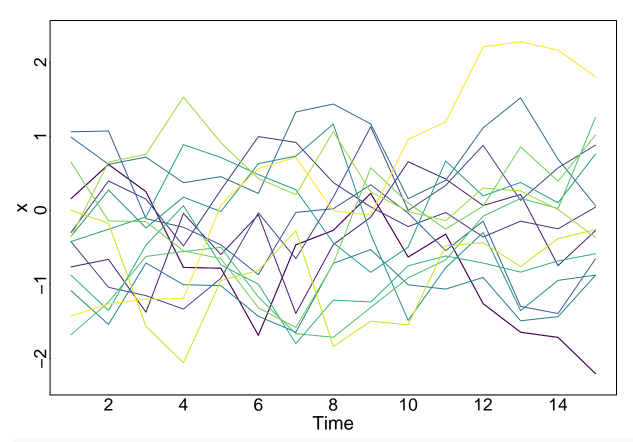
```
data = tscsGen(rho = .8)
#the goal is to plot every trend in x on one window
plot(data$Group == 1] ~ unique(data$Time), type = 'l')
```

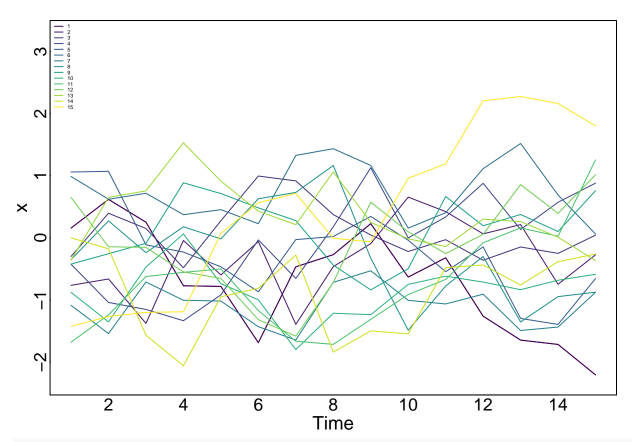


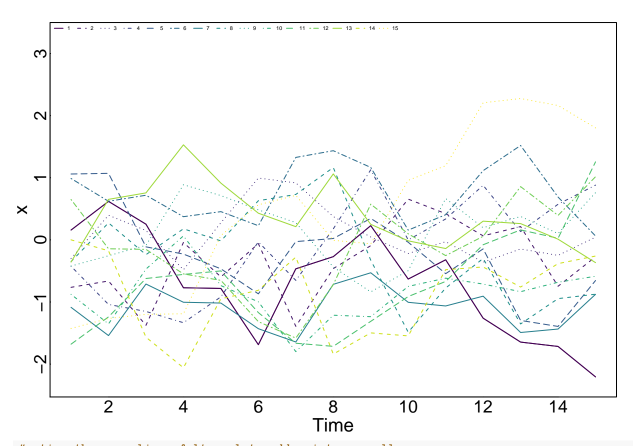


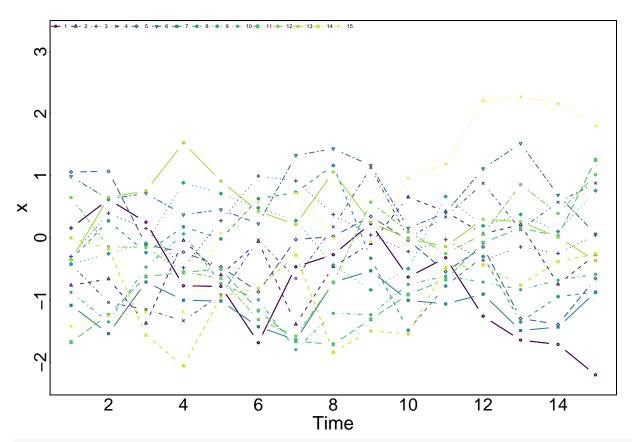
Loading required package: viridisLite

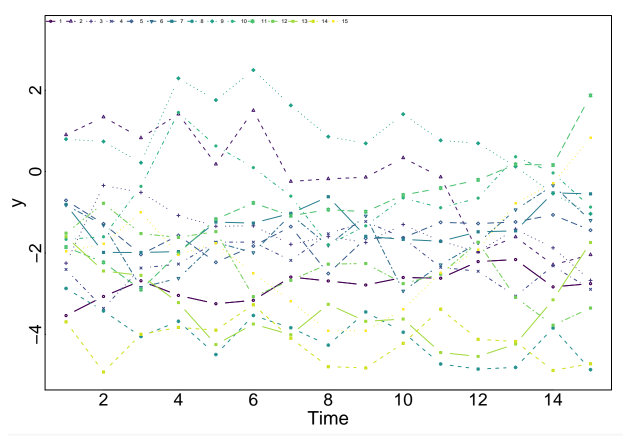


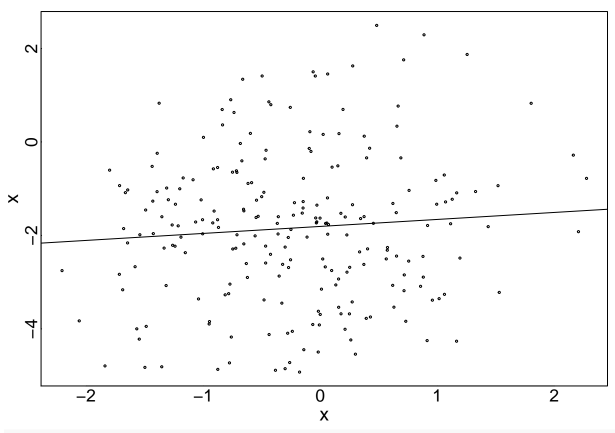


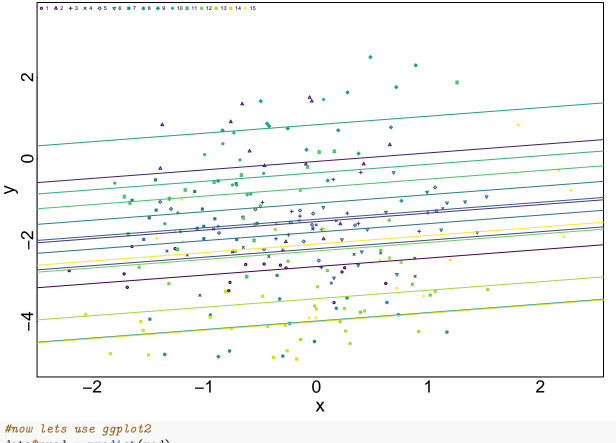




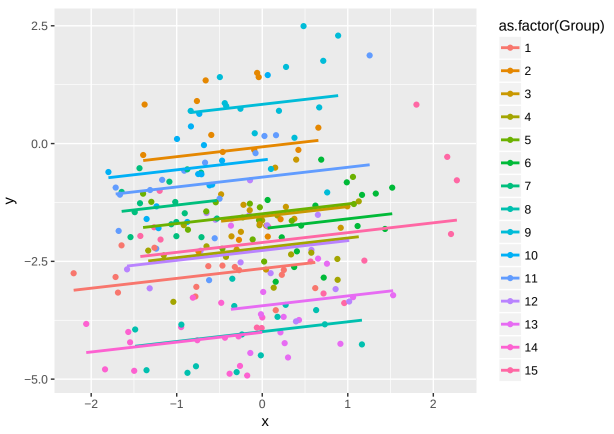


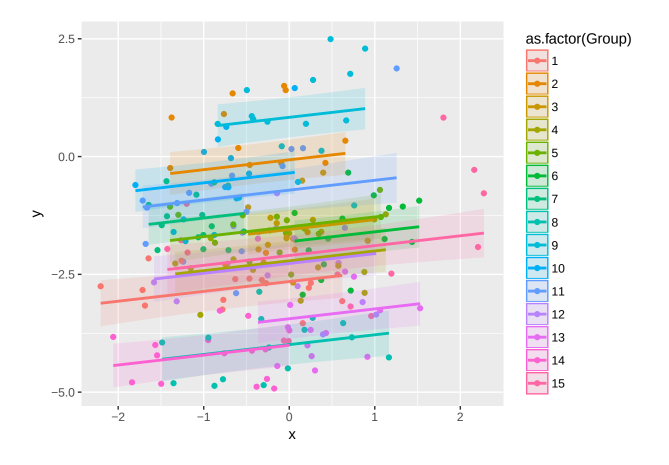






```
#now lets use ggplot2
data$pred = predict(mod)
library(ggplot2)
ggplot(data, aes(x = x, y = y, color = as.factor(Group)) ) +
    geom_point() +
    geom_line(aes(y = pred), size = 1)
```





Analyze the generated TSCS data and plot the results

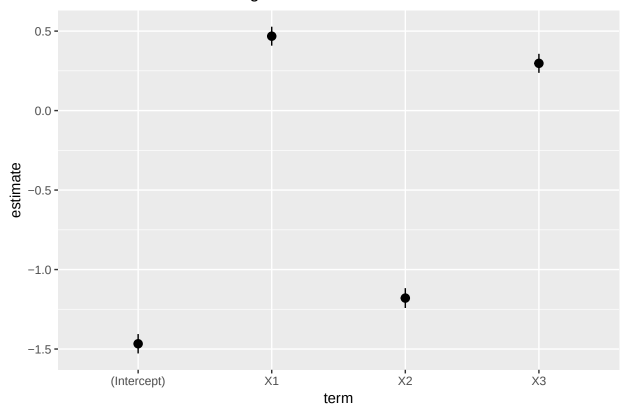
```
#we want to simulate a bunch of data sets at a range of rho, run a linear model, and check for coverage
simAnalysis = function(rho = seq(0, .9, by = .1),
                       n.datasets = 1000, ...){ #the ... notation allows additional arguments to be sen
  datas = lapply(1:n.datasets, function(y){
   lapply(rho, function(x){
      dd = tscsGen(x, seed = 100*y + 10*x)
     mod = lm(y \sim x + as.factor(Group) - 1, data)
      cov = confint(mod)[1,1] < .4 & confint(mod)[1,2] > .4
     pos = confint(mod)[1,1] > 0
     return(list(cov, pos))
  })})
  datas = unlist(datas)
  #so, it looped through each rho nested within each dataset
  datas.cov = datas[seq(1, length(datas), by = 2)]
  datas.pos = datas[seq(2, length(datas), by = 2)]
  covsByRho = numeric(length(rho))
  possByRho = numeric(length(rho))
  for(i in 1:length(rho)){
    covsByRho[i] = mean(datas.cov[seq(i, length(datas.cov), by = length(rho))])
   possByRho[i] = mean(datas.pos[seq(i, length(datas.pos), by = length(rho))])
 return(list(covsByRho, possByRho))
} #hw - parallelize this code
```

Coefficient plots

```
set.seed(99)
X = matrix(rnorm(1000*3), ncol = 3, nrow = 1000)
y = -1.5 + X%*%c(.5, -1.2, .3) + rnorm(1000)
mod = lm(y \sim X)
coef(mod)
## (Intercept)
                        Х1
                                    X2
## -1.4667980
                 0.4680407 -1.1794593
                                         0.2972580
confint(mod)
##
                    2.5 %
                              97.5 %
## (Intercept) -1.5277685 -1.4058275
                0.4083956 0.5276857
## X1
## X2
               -1.2418564 -1.1170621
## X3
                0.2373861 0.3571299
plot(coef(mod), 1:4, yaxt = 'n', pch = 18,
     xlim = c(min(confint(mod)), max(confint(mod))),
     ylab = '', xlab = 'Estimates')
segments(x0 = confint(mod)[,1], x1 = confint(mod)[,2],
         y0 = 1:4, y1 = 1:4)
```

```
abline(v=0, lty = 3)
axis(2, at = 1:4,
     tick = F, labels = c(expression(beta['0']),
                           expression(beta['1']),
                           expression(beta['2']),
                           expression(beta['3'])),
     las = 1)
\beta_3
\beta_2
\beta_1
\beta_0
      -1.5
                                         -0.5
                       -1.0
                                                           0.0
                                                                            0.5
                                      Estimates
library(broom)
coef = tidy(mod, conf.int = T)
coef
##
             term
                    estimate std.error statistic
                                                                      conf.low
                                                           p.value
## 1 (Intercept) -1.4667980 0.03107017 -47.209209 2.440994e-256 -1.5277685
## 2
               X1 0.4680407 0.03039474 15.398738 3.725627e-48 0.4083956
```

Coefficients of a linear regression model



Exponential family

- Fundamental to generalized linear models
- Puts PDF's in a generalized form
- Subfunctions are contained within the exponent component of the natural exponential function
- The isolated subfunctions quite naturally produce a small number of statistics that compactly summarize even large datasets without any loss of information
- Sufficient statistic: For some parameter contains all the information available in a given dataset about that parameter
- Suppose we consider a one-parameter conditional probability density function or probability mass function for the random variable Z of the form $F(z|\zeta)$, it is exponential if it can be written as:

$$f(z|\zeta) = \exp[t(z)u(\zeta)]r(z)s(\zeta)$$

$$= \exp\left[u(\zeta)\sum_{i=1}^{n} t(z_i) + \sum_{i=1}^{n} \log(r(z_i)) + n\log(s(\zeta))\right]$$

• Canonical form:

$$f(y|\theta) = \exp[y\theta - b(\theta) + c(y)]$$

- Simply sum over dimension of θ for higher dimensionality
- Derive the exponential family form of the following:
 - Poisson

$$f(y|\mu) = \frac{e^{-u}\mu^y}{y!}$$

- Binomial (with known n)

$$f(y|n,p) = \binom{n}{p} p^y (1-p)^{n-y}$$

– Normal (for both μ and σ^2)

$$f(y|\mu, \sigma^2) = \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left[-\frac{1}{2\sigma^2}(y-\mu)^2\right]$$

- Gamma

$$f(y|\mu,\delta) = \left(\frac{\delta}{\mu}\right)^{\delta} \frac{1}{\Gamma(\delta)} y^{\delta-1} \exp\left[\frac{-\delta y}{\mu}\right]$$

- Negative binomial

$$f(y|r,p) = \binom{r+y-1}{y} p^r (1-p)^y$$

- Multinomial (assume three outcomes)

$$p(\mathbf{Y}_i = r | \mathbf{X}) = \frac{\exp(\mathbf{X}_i \beta_r)}{1 + \sum_{s=1}^{k-1} \exp(\mathbf{X} \beta_s))}$$