Let's continue with the same motivating example from last time. We will use synthetic data generated by a Brownian motion process and pretend that it is observed longitudinal data. However, before getting to the statistics, I would like to take a tiny, tidy diversion. The functions in fda and other fundamental FDA R packages require data structured in matrices. Consequently, the examples in the basic FDA reference works (listed below) construct matrices using code that seems to be convenient for the occasion. I think this makes adapting sample code to user data a little harder than it needs to be. There ought to be standard data structures for working with FDA data. I propose tibbles or data frames with function values packed into lists.

The following function generates  $n\_points$  data points for each of  $n\_curve$  Brownian motion curves that represent the longitudinal data collected from  $n\_curve$  subjects

```
library(fda)
library(tidyverse)
library(plotly)
# Function to simulate data
fake_curves <- function(n_curves = 100, n_points = 80, max_time = 100){
    ID <- lin_curves
    x <- vector(mode = "list", length = n_curves)
    t <- vector(mode = "list", length = n_curves)</pre>
    for (i in 1:n_curves)(
   t[i] <- list(sort(runif(n_points,0,max_time)))
   x[i] <- list(cumsum(rnorm(n_points)) / sqrt(n_points)))</pre>
     }
df <- tibble(ID,t,x)
names(df) <- c("ID", "Time", "Curve")
return(df)</pre>
```

Notice that each curve is associated with a unique set of random time points that lie in the interval [0, max\_time]. Not being restricted to situations where data from all subjects must be observed at the same times is a big deal. However, in practice you may encounter problems that will require curve alignment procedures. We will ignore this for now. Note that the variables Time and Curve contain lists of data points in each cell.

```
set.seed(123)
n_curves <- 40
n_points <- 80
n_points <- 80
max_time <- 100
df <- fake_curves(n_curves = n_curves,n_points = n_points, max_time = max_time)
 head(df)
```

Later on, this kind of structure will be convenient for data sets that contain both FDA curves and other scalar covariates. Note that if you are using the RStudio IDE running the function View (df) will show you an expanded view of the tibble under a tab labeled df that should look something like this

2 c(0.630078394897282, 0.821551983244717, 0.942 c(0.11769671988644	
	89, -0.100916459832916, -0
3 2 (1.61604809109122.2.55266984459013.3.73699 (1.0.1864293834494	1, 0.000395164519687163, -0
3 7 (1.01007009109121, 1.01000907799013, 3.73009 (1.0.1007293037797	82, -0.104086631330097, -0
4 c(0.119162839837372, 1.54776689596474, 1.8768 c(-0.0082238129471	7205, -0.138883014297913,
5 c(0.699845445342362, 1.9191030645743, 4.60265 c(0.10814376730727	1, 0.096037685076762, 0.017
6 c(0.115581974387169, 0.46381508000195, 2.6010 c(-0.0932650795926	503, -0.0629588948659193,

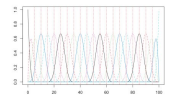
## Next, we unpack the data into a long form tibble and plot.



Now that we have the data, remember that FDA treats each curve as a basic data element living in an infinite dimensional vector space. The vectors, (XX), are random functions: (XX: \Omega \Rightarrow \text{Imathsc}(H)) where \(\text{(\Omega)}\) is an underlying probability space and \(\text{(\omega)}\) (\text{(\omega)}\) is a complete \(\text{(\omega)}\) in the reference in the \(\text{(\omega)}\) in the reference integrable \(\text{(\omega)}\) in the refere below which I have reproduced below from the previous post.

The bridge from the theory to practice is the ability to represent the random functions as a linear combination of basis vectors. This was the topic of the previous post. Here is some compact code to construct the basis

```
plot(basis)
```



This next bit of code formats the data in the long form tibble into the matrix input expected by the fda functions and creates an fda object that contains the coefficients and basis functions used to smooth data. Note the smoothing constant of lambda = .5.

 $W.obj \leftarrow Data2fd(argvals = argvals, y = y_mat, basisobj = basis, lambda = 0.5)$ 

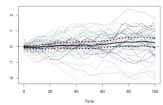
Next somewhat anticlimactically after all of the preparation and theory, we use the fda functions mean.fd () and std.fd () to calculate the pointwise mean and standard deviation from information contained in fda object. In order to use these objects to calculate the pointwise confidence interval for the mean it is necessary to construct a couple of new £da objects for the upper and lower curves. Then, we plot the smoothed curves for our data along with the pointwise mean and pointwise 95% confidence bands for the mean.

```
\label{eq:mean_def} \begin{split} & \text{M\_mean} <- \text{ mean.fd}(W.obj) \\ & \text{M\_sd } <- \text{std.fd}(W.obj) \\ & \text{F Create objects for the standard upper and lower standard deviation} \end{split}
```

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```
SE_u <- fd(basisobj = basis)
SE_l <- fd(basisobj = basis)
# Fill in the sd values
SE_l$\(\text{Scoefs} < \text{W}\) mean$\(\text{coefs} < \text{W}\) in mean$\(\text{coefs} < \text{W}\) in mean$\(\text{coefs} < \text{W}\) in mean$\(\text{V}\) mean$\(\te
```

## Smoothed Curv



Finally, we compute the covariance/correlation matrix for our sample of smoothed curves and use plotly to create an interactive plot of the three dimensional correlation surface along with a contour map

```
days <- seq(0,100, by=2)
cov_M <- var.fd(M.obj)
var_mat <- eval.bifd(days,days,cov_W)
fig <- plot.ply(x = days, y = days, z = -var_mat) %%
add_surface(contours = list(
    z = list(show=TRUE,usecolormap=TRUE, highlightcolor="#ff0000", project=list(z=TRUE))))
fig <- fig %%
    layout(scene = list(camera=list(eye = list(x=1.87, y=0.88, z=-0.64))))
fig</pre>
```

This covariance surface for the Brownian motion random walk we are using for data is interesting in its own right. There is a theoretical result that says that the estimate for the covariance function for this process is \(\text{(hat}(c)(t,s) = min(t,s)\). You can find the proof in the book by Kokoszka and Reimherr referenced below. Using the mouse to hover over some points in the graph makes this result seem plausible.

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