



Title of the topic

Workshop on Topic

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NBIS

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Introduction



This is a demo to showcase the RaukR presentation design guide, usage and capabilities of this presentation system. This presentation is created in RStudio using R package xaringan. This package in turn utilises the javascript library remarkjs as the underlying framework. Note that the markdown interpreter used by remarkjs (blackfriday) is different from that used by knitr (pandoc). Therefore, some features that you may be familiar with in regular RMarkdown may not work here. There are also many new features here not available in regular RMarkdown.

Note that many of the classes used in this presentation are custom built and not part of any standard package.

RMarkdown



- This is an RMarkdown presentation.
- Markdown is a simple formatting syntax for authoring documents exported to HTML, PDF etc.
- In RStudio, you create a .Rmd text file.
- Add YAML matter to the top if not already there.

```
title: "This is a title"
output:
    xaringan::moon_reader
---
```

- Click the **Knit** button for interactive render.
- Or using render() as such:

```
rmarkdown::render("report.Rmd",output_format="html_document")
```

Slides



Slide separators

Slides are separated by ---.

Incremental content on same slide is separated by -- like below.

Slide properties

Slides properties are created as below:

```
property: value
```

Slides can be named as such name: intro and then hyperlinked from elsewhere. For eg: go to [introduction] (#intro) displays as go to introduction.

Custom classes can be defined for each slide.

```
class: spaced
```

Few default options are *spaced* (increases line spacing), *middle* (center aligns content vertically) and *center* (center aligns content horizontally).

Slides can be hidden using exclude: true.

Slides can be dropped from page count using count: false.

Slides



Slide notes

Any content below ??? on a slide are notes. This is only visible in presenter mode (by pressing **P**).

Keyboard shortcuts

Press H to view keyboard shorcuts. Pressing C clones and creates two linked presentations.

Layout



The slide content can be organised into columns which can be nested if needed. Classes for 30, 40, 50, 60 and 70 have been implemented for left and right. Note that the total width must sum to 100.

Left content

Right content

Inner left content

Inner right content

Custom Divs



```
<div class="abstract">
This is a special box.
</div>
```

This is a special box.

Text Formatting



Headings can be defined as shown below.

```
## Level 2 heading
### Level 3 heading
#### Level 4 heading
##### Level 5 heading
###### Level 6 heading
```

Level 2 heading

Level 3 heading

Level 4 heading

Level 5 heading

Level 6 heading

Level 1 usage is not recommended. Use level 2 for slide titles. Use level 3 and below for other titles.

Six custom classes are defined for text scaling.

```
.largest[Largest text.]
.larger[Larger text.]
.large[Large text.]
Normal text.
.small[Small text]
.smaller[Smaller text.]
.smallest[Smallest text.]
```

Largest text. Larger text.

Large text.

Normal text.

Small text.

Smaller text.

Smallest text.

Text Formatting



Horizontal alignment of text can be adjusted as Icons from FontAwesome can be displayed shown below.

```
.left[Left aligned text.]
.center[Center aligned text.]
.right[Right aligned text.]
```

Left aligned text.

Center aligned text.

Right aligned text.

Indented quotes using >

This line is blockquoted

using the HTML <i> tag. Note that not all icons may work.

```
Here is a <i class='fa fa-calendar'></i>
calendar and a <i class='fa fa-couch'></i>
couch.
```

Here is a \rightleftharpoons calendar and a \rightleftharpoons couch.

A horizontal line can be created using ***

```
This is Bold text This is Bold text
This is Italic text This is Italic text
~~Strikethrough~~ text Strikethrough text
This is Subscript H<sub>2</sub>0 displayed as
H_2O
This is Superscript 2<sup>10</sup> displayed
as 2<sup>10</sup>
This is a [link](r-project.org) This is a link
```

Code formatting



Code can be defined inline where `this` looks like this. R code can be executed inline `r Sys.Date()` producing 2021-09-12. Code can also be defined inside code blocks.

This is code

```
This is code
```

R code is executed inside code blocks like this

```
```{r}
Sys.Date()
```
```

which shows the code and output.

```
Sys.Date()
## [1] "2021-09-12"
```

The code and results can be hidden by \``\{r,echo=FALSE,results='hide'}`.

```
data(iris)
head(iris[,1:2])
```

```
Sepal.Length Sepal.Width
## 1
             5.1
                        3.5
             4.9
## 2
                        3.0
             4.7
                        3.2
             4.6
                        3.1
             5.0
                        3.6
## 6
             5.4
                        3.9
```

Lists



Bulleted List

Bullet points are defined using the asterisk (*).

```
* Bullet 1
* Bullet 2
+ Sub-bullet 2.1
```

- Bullet 1
- Bullet 2
 - Sub-bullet 2.1

Incremental Bullets

```
* Incremental Bullet 1
--
* Incremental Bullet 2
```

Note that there is an empty line between the bullet point and the -- below.

- Incremental Bullet 1
- Incremental Bullet 2

Images • Markdown



Using Markdown

Using regular markdown.



The dimensions are based on image and/or fill up the entire available space. You have no control over the displayed dimensions.

Custom classes can be used to control size.

```
.size-20[![](assets/landing.png)]
.size-10[![](assets/landing.png)]
```

Images • HTML



Using Raw HTML

This image is 30% size.



This image is 20% size, has shadow and corners rounded.



Images • R



Using R

R chunks in RMarkdown can be used to control image display size using the arguemnt out.width.

This image is displayed at a size of 250 pixels.

```
```{r,out.width=250}
knitr::include_graphics('assets/landing.png')
```



# Math expressions



Some examples of rendering equations.

$$e^{i\pi}+1=0$$
 
$$rac{E imes X^2\prod I}{2+7}=432$$
 
$$\sum_{i=1}^n X_i$$
 
$$\int_0^{2\pi} \sin x \ dx$$

$$egin{aligned} \left(\sum_{i=1}^{n}i
ight)^2 &= \left(rac{n(n-1)}{2}
ight)^2 = rac{n^2(n-1)^2}{4} \ X &\sim \mathrm{N}(0,1) \ Y &\sim \chi^2_{n-p} \ R &\equiv X/Y \sim t_{n-p} \end{aligned} \ P(|X-\mu| > k) &= P(|X-\mu|^2 > k^2) \ &\leq rac{\mathbb{E}\left[|X-\mu|^2
ight]}{k^2} \ &\leq rac{\mathrm{Var}[X]}{k^2} \end{aligned}$$

# Tables • kable



The most simple table using kable from R package knitr.

knitr::kable(head(iris), 'html')

Sepal.Length	Sepal.Width	Petal.Length	Petal.Width	Species
5.1	3.5	1.4	0.2	setosa
4.9	3.0	1.4	0.2	setosa
4.7	3.2	1.3	0.2	setosa
4.6	3.1	1.5	0.2	setosa
5.0	3.6	1.4	0.2	setosa
5.4	3.9	1.7	0.4	setosa

## Tables • kableExtra



More advanced table using kableExtra and formattable.

	Sepal.Length	Sepal.Width	Petal.Length	Petal.Width	Species
1	5.1	3.5	1.4	0.2	setosa
2	4.9	3.0	1.4	0.2	setosa
51	7.0	3.2	4.7	1.4	versicolor
52	6.4	3.2	4.5	1.5	versicolor
105	6.5	3.0	5.8	2.2	virginica
106	7.6	3.0	6.6	2.1	virginica

# Tables • DT



Interactive table using R package DT.

library(DT)

DT::datatable(iris[1:20,],options=list(pageLength=7))

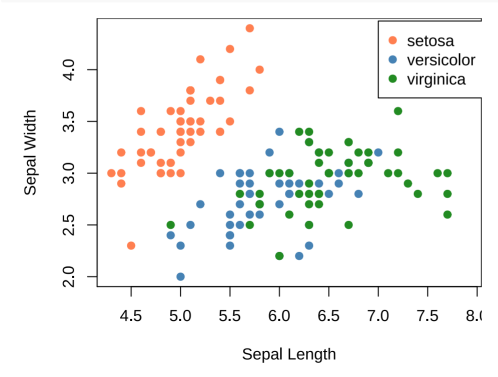
Show 7				Search:		
	Sepal.Length 🖣	Sepal.Width 🖣	Petal.Length 🔷	Petal.Width 🔷	Species	\$
1	5.1	3.5	1.4	0.2	setosa	
2	4.9	3	1.4	0.2	setosa	
3	4.7	3.2	1.3	0.2	setosa	
4	4.6	3.1	1.5	0.2	setosa	
5	5	3.6	1.4	0.2	setosa	
6	5.4	3.9	1.7	0.4	setosa	
7	4.6	3.4	1.4	0.3	setosa	
Showing 1 to 7	7 of 20 entries			Previous 1 2	2 3	Next

## Static plots • Base Plot



Plots using base R are widely used and may be good enough for most situations. But they lack a consistent coding framework.

```
{par(mar=c(5,5,0,0))
 plot(x=iris$Sepal.Length,y=iris$Sepal.Width,
 col=c("coral","steelblue","forestgreen")[iris$Species],
 xlab="Sepal Length",ylab="Sepal Width",pch=19)
 legend(x=7,y=4.47,legend=c("setosa","versicolor","virginica"),
 col=c("coral","steelblue","forestgreen"),pch=19)}
```

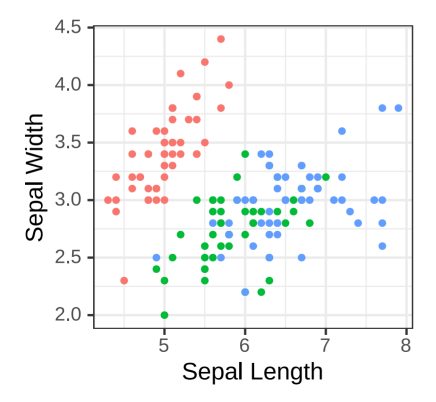


# Static plots • ggplot2



R package ggplot2 is a versatile and almost complete plotting solution.

```
iris %>%
 ggplot(aes(x=Sepal.Length,y=Sepal.Width,col=Species))+
 geom_point(size=2)+
 labs(x="Sepal Length",y="Sepal Width")+
 theme_bw(base_size=18)
```



## **Species**

- setosa
- versicolor
- virginica

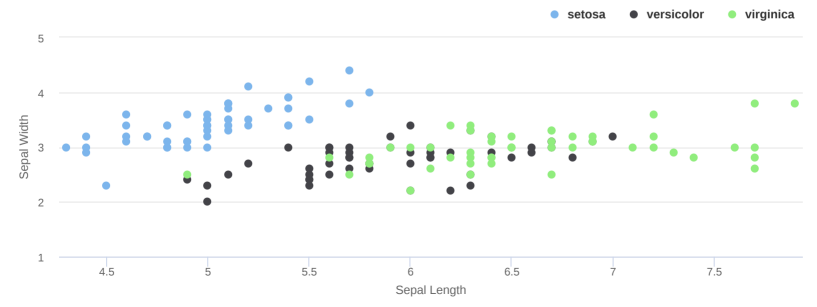
# Interactive plots • highcharter



R package <a href="highcharter">highcharter</a> is a wrapper around javascript library <a href="highcharts">highcharter</a> .

```
library(highcharter)
h <- iris %>%
 hchart("scatter",hcaes(x="Sepal.Length",y="Sepal.Width",group="Species")) %>%
 hc_xAxis(title=list(text="Sepal Length"),crosshair=TRUE) %>%
 hc_yAxis(title=list(text="Sepal Width"),crosshair=TRUE) %>%
 hc_chart(zoomType="xy",inverted=FALSE) %>%
 hc_legend(verticalAlign="top",align="right") %>% hc_size(height=320)

htmltools::tagList(list(h))
```

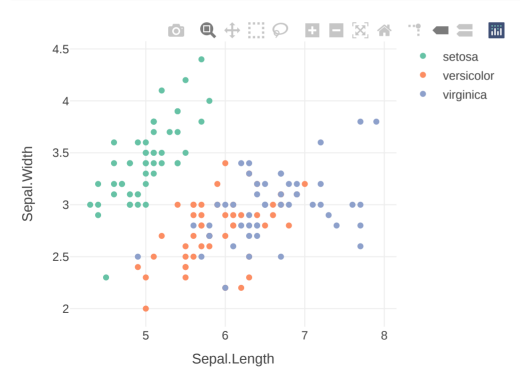


# Interactive plots • plotly



R package plotly provides R binding around javascript plotting library plotly.

```
library(plotly)
p <- iris %>%
 plot_ly(x=~Sepal.Length,y=~Sepal.Width,color=~Species,width=500,height=350) %>%
 add_markers()
p
```



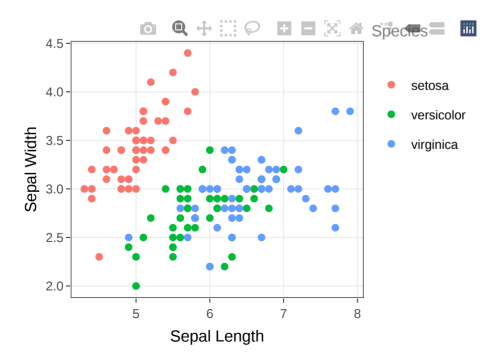
# Interactive plots • ggplotly



plotly also has a function called ggplotly which converts a static ggplot2 object into an interactive plot.

```
library(plotly)
p <- ggplot(iris,aes(x=Sepal.Length,y=Sepal.Width,col=Species))+
 geom_point()+
 labs(x="Sepal Length",y="Sepal Width")+
 theme_bw(base_size=12)

plotly::ggplotly(p,width=460,height=330)</pre>
```



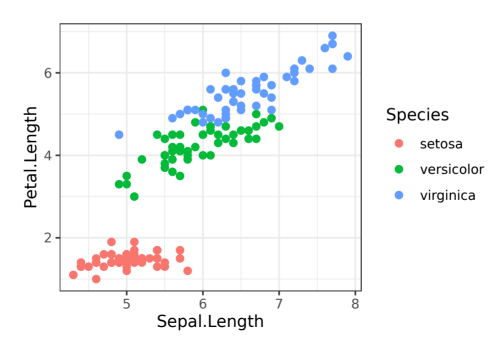
# Interactive plots • ggiraph



R package ggiraph converts a static ggplot2 object into an interactive plot.

```
library(ggiraph)
p <- ggplot(iris,aes(x=Sepal.Length,y=Petal.Length,colour=Species))+
 geom_point_interactive(aes(tooltip=paste0("Petal Length: ",Petal.Length,"\n<
 theme_bw()

tooltip_css <- "background-color:#f8f9f9;padding:10px;border-style:solid;border-width:2px
ggiraph(code=print(p),hover_css="cursor:pointer;stroke:black;fill-opacity:0.3",zoom_max=5</pre>
```



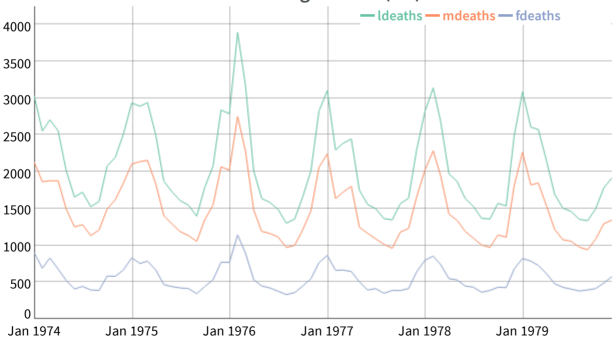
# Interactive time series • dygraphs



R package dygraphs provides R bindings for javascript library dygraphs for time series data.

```
library(dygraphs)
lungDeaths <- cbind(ldeaths, mdeaths, fdeaths)
dygraph(lungDeaths, main="Deaths from Lung Disease (UK)") %>%
 dyOptions(colors=c("#66C2A5","#FC8D62","#8DA0CB"))
```

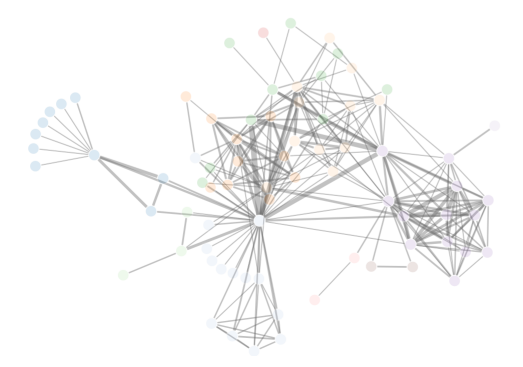




# Network graph



R package networkD3 allows the use of interactive network graphs from the D3.js javascript library.

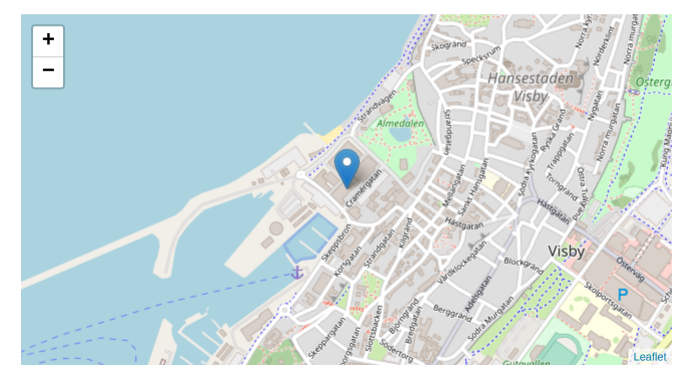


# Interactive maps • leaflet



R package leaflet provides R bindings for javascript mapping library; leafletjs.

```
library(leaflet)
leaflet(height=350,width=650) %>%
 addTiles(urlTemplate='http://{s}.tile.openstreetmap.org/{z}/{x}/{y}.png') %>%
 #addProviderTiles(providers$Esri.NatGeoWorldMap) %>%
 addMarkers(lat=57.639327,lng=18.288534,popup="RaukR") %>%
 setView(lat=57.639327,lng=18.288534,zoom=15)
```

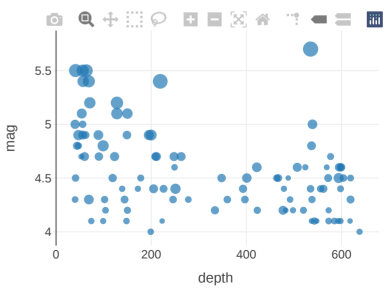


# **Linking Plots • crosstalk**



R package crosstalk allows crosstalk enabled plotting libraries to be linked. Through the shared 'key' variable, data points can be manipulated simultaneously on two independent plots.





# Thank you. Questions?



Graphics from freepik.com

Created: 12-Sep-2021 • Roy Francis • SciLifeLab • NBIS