How to Create an R Kernel for Jupyter Lab

Course: DS 5001
Module: 11 Lab
Topc: R Kernel
Author: R.C. Alvarado
Date: 8 April 2024

Motivation

You want to run R in a Jupyter notebook. This means having available an R kernel in the list of kernels you see in the Launcher window (in Jupyter Lab).

1. Create and activate an R environment with Conda

From the command line in Rivanna, create and activate an R environment.

Here we create one with the essentials and base packages for R.

```
conda create \neg n r_env r-essentials r-base conda activate r_env
```

You are now in an environment that runs R.

2. Run the R shell

Now, run R from the command line.

R

3. Register the kernel

Once in the R interactive shell, register the kernel with the following commands.

```
install.packages("IRkernel") # Not really needed since included in r-essentials
IRkernel::installspec(name = 'r_env', displayname = 'R Environment')
quit()
```

4. Deactivate the environment

Once you are done with this, you can deactivate the environment.

conda deactivate

5. Run Jupyter Lab

Start Jupyter Lab from the OpenOnDemand webpage.

If you already have a session running, close it down and restart it.

You should see a kernel tile like the following in the launcher window:



6. Going Forward

If everything worked, you can close down the command line terminal and work in Jupyter Lab.

Note that you can add more packages to your R environment by issuing the command from within the R environment:

```
conda install -c r <r-package>
```

<r-package> can be anything from the following list:

• https://repo.anaconda.com/pkgs/r/

Or, if you are not in the R environment, you can specify the target environment with the argument \neg n:

conda install -n r_env -c r <r-package>