

Data Science Pipelines with {targets} [Dr. Hanjo Odendaal]



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ABOUT ME

With over 10 years experience working throughout Africa, Hanjo leads 71point4's analytics, statistics and engineer aspects on projects. His expertise focuses on how data can be employed to answer impactful and meaningful questions that always aim to put people at the center.

He has worked in banking, insurance, housing, agriculture, telecommunications and credit; focusing mostly on designing and managing the implementation of analytical solutions to deliver real-time insights for the client.

Hanjo holds a PhD in Economics from the University of Stellenbosch. His thesis focused on the application of natural language processing and statistical learning applied within traditional economic frameworks.



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Software Engineering



Production Machine Learning



Web Scraping



High Performance Cloud Computing





Defence against the dark arts a.k.a code entropy

Why pipelines?



When writing code, there are always two people involved: You and Future You... and about 90% of the time Future You is going to hate how you structured your project 6 months ago.

Without a build automation tool, a pipeline is nothing but a series of scripts that get called one after the other. In most cases when students start out, the pipeline tends to be one very long script that is usually a 4 and hope as scaffolding that ensures operations successfully.

- Scripts can, and will, be executed out of order.
 - if you make a small change in one of your scripts, a logic execution needs to follow to ensure everything is executed in the correct order.
- Pipelines written as scripts are usually quite difficult to read and understand.
 - Comments help, but has to also be updated as you update your scripts.

These are just some of the basic issues... BUT luckily you have heard of the term "functional" programming. This part of the course will show you how to supercharge this concept 🚀!

Learning outcomes



targets

Now and then, the bright and shiny objects that we stumble across can turn out to be very useful.

We want to take you on a high-level tour of {targets}. Getting you started with the basics! The rest of the journey is up to you:

- What is {targets}?
- Integrating databases into your {targets} pipeline
- NLP and {targets}



r Reviewed 1,055 10.21105/joss.02959 DOI 10.5281/zenodo.10357140 R Targetopia memi 15.1 repo status Active () check passing (? codecov 100% () lint passing

Pipeline tools coordinate the pieces of computationally demanding analysis projects. The targets package is a <u>Make-like</u> pipeline tool for statistics and data science in R. The package skips costly runtime for tasks that are already up to date, orchestrates the necessary computation with implicit parallel computing, and abstracts files as R objects. If all the current output matches the current upstream code and data, then the whole pipeline is up to date, and the results are more trustworthy than otherwise.

Philosophy

A pipeline is a computational workflow that does statistics, analytics, or data science. Examples include forecasting customer behavior, simulating a clinical trial, and detecting differential expression from genomics data. A pipeline contains tasks to prepare datasets, run models, and summarize results for a business deliverable or research paper. The methods behind these tasks are user-defined R functions that live in R scripts, ideally in a folder called "##" in the project. The tasks themselves are called "targets", and they run the functions and return R objects. The targets package orchestrates the targets and stores the output objects to make your pipeline efficient, painless, and reproducible.

Prerequisites

familiarity with the <u>R programming language</u>, covered in <u>R for Data Science</u>.
 <u>Data science workflow management techniques</u>.
 <u>How to write functions</u> to prepare data, analyze data, and summarize results in a data analysis project

What is {targets}?



It is a workflow management package for the R programming language developed and maintained by Will Landau. If some of you program python / bash you might be familiar or have heard about snakemake or make files. {targets} is the R equivalent of make files as you will see when we run tar_make() to run our pipeline.

The major features of targets include:

- Automation of workflow
- Caching of workflow steps
- Batch creation of workflow steps
- Parallelization at the level of the workflow

Why is this amazing?

- Return to a project after working on something else and immediately pick up where you left off without confusion or trying to remember what you were doing
- If the workflow changes in any way, only re-run the parts that that are affected by the change
- Massively scale up the workflow without changing individual functions through dynamic branching and parallelization (advanced)

All of these characteristics of pipelines give us a high probability of \neq reproducibility \neq !

What is a DAG?



DAG or Directed acyclic graph, is a mathematical concept. This "DAG" consists of vertices and edges (also called arcs), with each edge directed from one vertex to another, such that following those directions will never form a closed loop.

You are constantly thinking about 3 things: *Input > function > output*.





What is a DAG?



DAG or Directed acyclic graph, is a mathematical concept. This "DAG" consists of vertices and edges (also called arcs), with each edge directed from one vertex to another, such that following those directions will never form a closed loop.

penguin_csv_file >

penguins_data_raw(penguin_csv_file) >

penguins_data(penguins_data_raw)



Ready for the basics!





Time for the setup



Please take a couple of minutes to install these packages (Ncpu parameter for parallel installation):

Pipeline

```
install.packages("igraph", Ncpus = 4)
install.packages("visNetwork", Ncpus = 4)
install.packages("targets", Ncpus = 4)
install.packages("tarchetypes", Ncpus = 4)
```

Utilies

```
install.packages("tidyverse", Ncpus = 4)
install.packages("glue", Ncpus = 4)
install.packages("logger", Ncpus = 4)
install.packages("lubridate", Ncpus = 4)
```

Modeling

install.packages("ranger", Ncpus = 4)

NLP

install.packages("quanteda", Ncpus = 4)
install.packages("topicmodels", Ncpus = 4)



Time for the setup



You always have to work in a Project when working with {targets} otherwise its not going to work! We will use the Texevier to help us setup that folder.

(base) hanjo@optimus:~/Downloads/targets_example\$ tree

- code
 Data_Create.R
 Example_Plot_Scatter.R
 data
 targets_example.Rmd
 targets_example.Rproj
 Tex
 EconEJ.bst
 elsevier-harvard.csl
 harvard1.csl
 harvard-stellenbosch-university.csl
 packages.txt
 ref.bib
 - TexDefault.txt

Sidequest 💎 🕉 👑 👕



If you want your prompt to show you what Github branch you are on, add the following to your .Rprofile. Easiest is usethis::edit_r_profile():

```
if(requireNamespace("prompt", quietly = TRUE)){
# devtools::install_github("gaborcsardi/prompt@x")
prompt_git ← function( ... ){
    paste0(
        "[", prompt::git_branch(), "]", ">"
    )
    }
    prompt::set_prompt(prompt_git)
    rm(prompt_git)
}
```

Time for the setup



After you have ensured that you can knit the file called: targets_example.Rmd, create a file called _targets.R in the parent folder. Take special note of the _ in the filename. Also create a README.qmd !

#_targets.R
<pre># 1.0 Setup options(tidyverse.quiet = TRUE)</pre>
library (targets) library (tarchetypes)
* Libraries
<pre>tar_option_set(packages = c(</pre>
data
"tidyverse",
"lubridate",
"glue",
"logger",
modeling
"plm",
reporting
"huxtable"
<pre># Source all of your functions into environment sapply(list.files("code", full.names = T), source)</pre>

README.qmd

```
title: "My readme"
format: html
editor_options:
   chunk_output_type: console
execute:
   echo: false
   eval: false
---
```

```
```{r}
library(tidyverse)
library(targets)
```

Execute the pipeline using `tar\_make`:

```
```{r}
tar_make()
```
```



# Building first pipeline





The most obvious step in our data analysis is obviously getting our data into R.

Golden rule:

- If it can be avoided, never transform your data in Excel
- Use R to showcase how you extracted, loaded and transformed (ELT) your data

In your \_targets.R file, add the following "targets":

c(target\_data\_load)

#### Data target breakdown



l assign an object of list() class to my first grouping of targets.

```
target_data_load \leftarrow list()
```

In my first target I say that the input of "data/Example\_data.rds" should be assigned to input\_file:

tar\_target(input\_file,"data/Example\_data.rds")

Secondly I then use input\_file as my input in the function read\_rds and call that output, example\_df:

tar\_target(examples\_df, read\_rds(input\_file)

Lastly I concatenate all my groups together:

c(target\_data\_load)

^ I always use a suffix to help me know what the output type is: \*\_df, \*\_list, \*\_plot, \*\_write





Once you have completed the input into your \_targets.R file, go to your Readme.qmd and run the tar\_make() command:

[main]>tar\_make()

- start target input\_file
- built target input\_file [1.188 seconds]
- start target examples\_df
- built target examples\_df [0 seconds]
- ▶ end pipeline [1.249 seconds]

Ok, so what happened and what now? Well, congrats on your running your first pipeline!! 🎉

To see your pipeline, add the command tar\_visnetwork(targets\_only = TRUE, label = "time")
to your Readme.qmd ... you should see the your DAG once
you have run it.



## Accessing the objects



Loading data into the pipeline is all good, but how can we use it? There is primarily two commands: tar\_read and tar\_load.

In the *first* instance you can imagine its like reading in a csv:

```
[main]>out \leftarrow tar read(examples df)
[main]>print(out)
A tibble: 100 × 3
 Height_Score Weight_Score Agility_Score
 <dbl>
 <dbl>
 <dbl>
 5.75
 6.11
 3
1
2
 3.84 3.37
 3
3
 5.71
 2.86
 3
 6.02
 5.59
 3
4
5
 6.56
 6.48
 3
```

Where in the case of tar\_load the object is loaded into your environment and you can then use it:

```
[main]>tar_load(examples_df)
[main]>print(examples df)
```

## How much a target should do?



The {targets} package automatically skips targets that are already up to date, so it is best to define targets that maximize time savings and breaks down the analytical work into succinct problem definitions: (1) Read, (2) Augement, (3) Analyse etc. Good targets usually:

- Invoke no side effects such as modifications to the global environment!
- Are large enough to subtract a decent amount of runtime when skipped
- Are small enough that some targets can be skipped even if others need to run
- Return a single value that is:
  - $\circ\,$  Easy to understand and introspect.
  - Meaningful to the project.
  - $\circ~\mbox{Easy}$  to save as a file

Think of organising your pipeline as a package. 📦

## Adding modeling component to pipeline



I would bet most pipelines dont just want to read data, but analyse them. So, how do we do that? Well lets start by creating a new script for our "modeling" functions: code/models.R:

Basic Model:

```
model_lm ← function(examples_df){
 lm(Agility_Score ~., data = examples_df)
}
```

Interaction Model:

<sup>\*</sup> Take note of how I specify the input name the same as my previous output. You will thank me later ;-)

## Adding modeling component to pipeline



Once you have saved the R file, we need to now add the functions to our pipeline in \_targets.R:

```
2.0 Data -----
target data load \leftarrow list(
 tar target(input_file,
 "data/Example data.rds"),
 tar target(examples df,
 read rds(input file))
3.0 Model -----
target_model \leftarrow list(
 tar target(model lm model,
 model lm(examples df)),
 tar target(model lm interactions model,
 model lm interactions(examples df))
c(target data load,
 target model)
```

## Adding modeling component to pipeline



Now that you have added in the new targets. Have a look at your DAG:



Then, you have guessed it, time to run tar\_make():

- [main]>tar\_make()
- ✓ skip target input\_file
- skip target examples\_df
- start target model\_lm\_model
- built target model\_lm\_model [0.012 seconds]
- start target model\_lm\_interactions\_model
- built target model\_lm\_interactions\_model [0.001 seconds]
- ▶ end pipeline [1.213 seconds]



Models in themselves arent that useful. Lets extract the RMSE using yardstick to compare which model was a better fit and then compare them visually. Because we want to apply the same action on the model, we only going to need a single function! In the code/models.R file lets add a function to extract the models' RMSE.

```
extract_rmse ← function(model, examples_df, model_name){
 out ← tibble(truth = examples_df$Agility_Score,
 prediction = predict(model, examples_df))
 model_summary ← yardstick::rmse(out, truth, prediction) %>%
 janitor::clean_names() %>%
 mutate(model_name = model_name)
 return(model summary)
```



#### In \_targets.R:



#### Now lets plot the RMSE!

```
model lm interactions model rmse){
 bind rows(model lm interactions model rmse,
 model_lm_model_rmse) %>%
 mutate(model name = snakecase::to title case(model name)) %>%
 ggplot(., aes(model name, estimate,
 color = model_name, pch = model_name)) +
 geom point(size = 5) +
 scale_color_grey(
 start = 0.2,
 end = 0.6
) +
 labs(
 y = "RMSE",
 x = "Model Specification",
 color = ""
) +
 ggthemes::theme stata(base size = 20) +
 theme(plot.background = element rect(fill = "white")) +
 theme(legend.position = "bottom") +
 guides(pch = "none")
```

```
target model \leftarrow list(
 tar target(model lm model,
 model lm(examples df)),
 tar_target(model_lm_interactions model,
 model lm interactions(examples df)),
 tar_target(model lm model rmse,
 extract_rmse(model_lm_model,
 examples df,
 model_name = "model_lm")),
 tar target(model lm interactions model rmse,
 extract rmse(model lm interactions model,
 examples df,
 model name = "model lm interactions")),
 tar_target(plot_rmse_gg,
 plot_rmse(model_lm_model_rmse,
 model lm interactions model rmse))
```

Now lets plot the RMSE!

```
model lm interactions model rmse){
 bind rows(model lm interactions model rmse,
 model_lm_model_rmse) %>%
 mutate(model name = snakecase::to title case(model name)) %>%
 ggplot(., aes(model_name, estimate,
 color = model_name, pch = model_name)) +
 geom_point(size = 5) +
 scale_color_grey(
 start = 0.2,
 end = 0.6
) +
 labs(
 y = "RMSE",
 x = "Model Specification",
 color = ""
) +
 ggthemes::theme_stata(base_size = 20) +
 theme(plot.background = element rect(fill = "white")) +
 theme(legend.position = "bottom") +
 guides(pch = "none")
```







#### Now lets plot the RMSE!

```
model lm interactions model rmse){
 bind rows(model lm interactions model rmse,
 model_lm_model_rmse) %>%
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 ggplot(., aes(model_name, estimate,
 color = model_name, pch = model_name)) +
 geom_point(size = 5) +
 scale_color_grey(
 start = 0.2,
 end = 0.6
) +
 labs(
 y = "RMSE",
 x = "Model Specification",
 color = ""
) +
 ggthemes::theme stata(base size = 20) +
 theme(plot.background = element rect(fill = "white")) +
 theme(legend.position = "bottom") +
 guides(pch = "none")
```



## Integrate all into report



So why did we start with Texevier? Because we want our article to update if we make a change... the last piece of the puzzle is to render the article.Rmd:

**Step 1:** Make sure you have library(targets) in your setup chunk.

Step 2: Use tar\_read Or tar\_load in article.Rmd.

**Step 3:** Build the render into your pipeline \_targets.R:



**Step 4:** View your network to see that the correct outputs go in article:



**Step 5:** Run tar\_make(), sit back and marvel at your work in glory.

#### Are you not entertained?



If its good enough for Taylor<sup>\*</sup> its good enough for you...



\*approximation





#### Homework



Using the techniques of today, transform the "script" based flow from the practical Tidy Regressions into a pipeline.

The assignment is simple:

- Create a new project
- Setup project for {targets}
- Change the "chunks" into functions
- Create a pipeline to bring in data, run models and output plot
- Run pipeline from section Load Data to Backward Selection

... in the next lesson we will be introducing the concept of "databases".