

# From Docker to Kubernetes: Scaling Machine Learning Algorithms

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## Important links

DSSV 2019 Talk: <https://github.com/jharner/DSSV2019rc2>

RSpark: <https://github.com/jharner/rspark>

RSpark Tutorial: <https://github.com/jharner/rspark-tutorial>

Rc<sup>2</sup> Server: <https://github.com/rc2server>

Rc<sup>2</sup> Swift Client: <https://github.com/mlilback/rc2SwiftClient>

# DevOps for Data Science

Data Science Platforms are complex! How do we solve the data analysis issues of:

- ▶ collaboration,
- ▶ sharing,
- ▶ reliability
- ▶ scalability, and
- ▶ reproducibility?

DevOps (development + operations) has been revolutionized by containers and methods of orchestrating containers. Specifically, we will examine two technologies which are changing how complex software systems are built, deployed, and maintained. These are:

- ▶ Docker: a technology for building, deploying, and running container images;
- ▶ Kubernetes: a technology for deploying, scaling, and managing containerized applications.

Docker Swarm is not as scalable or feature complete as Kubernetes.

# Docker

Docker containers are becoming essential for software development and for the deployment of complex computing environments. This talk presents Docker solutions to address these and other use cases relevant to statisticians and data scientists.

- ▶ Rocker: R-based images built by and hosted on Docker Hub
- ▶ Rc<sup>2</sup>: R-based image hosted on GitHub.

Perhaps the most important reason for using Docker by statisticians and data scientists is the complexity of their computing environments, e.g., aligning version numbers for R and its packages, drivers, SQL and NoSQL databases, Hadoop, Spark and its packages, TensorFlow and GPUs, etc.

## What is Docker?

Docker allows developers, devops, and sysadmins to develop, deploy, and run applications using containers. We call this containerization.

Containers are:

- ▶ Flexible
- ▶ Lightweight
- ▶ Portable
- ▶ Scalable

A container runs natively on Linux and shares the kernel of the host machine with other containers. A container runs as a discrete process and thus its memory requirements are nearly equivalent to other executables, i.e., it is lightweight. On the other hand, a virtual machine runs a guest OS which is built on a hypervisor, through which host resources are accessed. Running multiple VMs is very heavy.

## Why Use Docker?

A Container Platform provides a complete solution, e.g., the components needed for teaching a data science program.  
Increasingly these platforms are being built on Kubernetes.

Example: JupyterHub

JupyterHub allows instances of a single-user Jupyter notebook to be spawned and managed. Two distributions are available:

- ▶ The Littlest JupyterHub for 0–50 users on a single machine
- ▶ Zero to JupyterHub on Kubernetes for a large number of users and machines

Example: RStudio Server Pro with Launcher and Kubernetes

RStudio Server is not supported, i.e., Launcher is a Pro product.

## Images and Containers

The two principal Docker entities are:

- ▶ Image: an executable package that includes everything needed to run an application
- ▶ Container: a runtime instance of an image

The image contains the code, configuration files, environmental variables, libraries, and the runtime. You can see the images by running the following command in a terminal:

```
docker images
```

A container is an image with state, i.e., a user process. You can view the running containers by executing:

```
docker ps
```

or you can see all containers by executing:

```
docker ps -a
```

## Running an Image

You can run an image by:

```
docker run hello-world
```

If the `hello-world` image is not found locally, it is automatically downloaded and launched from Docker Hub, the default image registry.

## Rocker Project

Rocker project: a widely-used suite of Docker images with customized R environments for particular tasks.

Currently, there are 17 repos in rocker-org.

The most important to us are:

- ▶ rocker-versioned
  - ▶ r-ver: versioned base R
  - ▶ rstudio: adds rstudio
  - ▶ tidyverse: adds tidyverse and devtools
  - ▶ verse: adds tex and publishing packages
- ▶ shiny
- ▶ ml

These and other available images provide a large number of use cases.

# Running R/Rstudio in Containers

A terminal version of R can be run in `r-base` or `r-ver`.

```
docker run --rm -ti rocker/r-base bash
```

The `rocker/r-base` image is downloaded from Docker Hub (if not already in the local filesystem) and run. The `-it` flags cause a bash session to start. Typing R into the terminal starts R (currently version 3.5). The `-rm` flag causes the container to be removed when the session is exited.

Or just:

```
docker run --rm -ti rocker/r-base
```

RStudio can be run by:

```
docker run --rm -p 8787:8787 rocker/rstudio
```

Open a browser with the URL `localhost:8787` and sign in with user/password: `rstudio/rstudio`.

## Extending Images

A developer or user may find that an image needs to be extended, e.g., a critical R package is missing or needs to be updated. This can be done in one of two ways:

- ▶ install the R package inside the container using `install.packages`, or
- ▶ rebuild the image by modifying the Dockerfile.

The first method results in mutable infrastructure, i.e., changes are made incrementally. The state of the infrastructure is thus dependent on incremental updates. This method should only be used in critical situations and should be viewed as a temporary fix.

The second method results on an immutable infrastructure, i.e., the state does not change based on actions by the user. Rather, the image is completely rebuilt by the developer.

## Rocker Deployment

Any of the Rocker images can be deployed in a single container. This container can be run on your laptop, on your departmental server, or on a cloud platform such as AWS. Although the Rocker images can be built from their GitHub sources, typically the pre-built images on Docker Hub are simply downloaded and run.

Why go to this trouble? Why not just run RStudio locally or on a server?

Suppose you are writing a paper and you want it to be truly reproducible. At this point you want to capture and maintain not only the Rmd or Rnw files, but also the synchronized R and required package versions. Further, the R environment can be synchronized with Linux distro and version along with required storage platform for your data.

But the reasons go well beyond reproducibility.

# Using Docker for Continuous Integration Deployment [DevOps]

Docker is a platform for containerized networking, compute, and storage for distributed applications. Docker improves:

- ▶ Velocity: increase flow by continuous integration (CI) and continuous delivery (CD);
- ▶ Variation: converge artifacts with immutable binaries;
- ▶ Visualization: visualize (elevate) services in the pipeline by bounding context and using containers for isolation.

Reference paper on CI/CD

Use Kubernetes for container orchestration!

# Kubernetes

Kubernetes is a platform for running and coordinating related, containerized applications across a cluster of (typically virtual) machines. It manages the complete lifecycles of containerized applications and services to provide:

- ▶ predictability
- ▶ reliability
- ▶ scalability
- ▶ availability

## Kubernetes Principles

Container and Kubernetes developers should adhere to certain principles:

- ▶ immutable infrastructure: update an image with a newer image in a single operation, i.e., no incremental updates (except in emergencies);
- ▶ declarative configuration: describe the desired state of your application.

In mutable infrastructures changes are applied incrementally, e.g., `apt-get` in Linux or `install.packages` in R. This should only be done as a temporary fix. An imperative declaration defines the state as a series of actions rather than a declaration of the desired state as is done with Kubernetes.

Kubernetes makes sure the actual state of your environment is aligned with the desired state. Not only that, Kubernetes is an online, self-healing system, i.e., it continuously takes action to ensure the current state matches the desired state.

# Kubernetes Architecture

The machines in a Kubernetes cluster are classified as:

- ▶ a master server: the primary point of contact with the cluster.  
It exposes the Kubernetes API, aligns the actual state to the desired state, schedules work, orchestrates communication, etc.
- ▶ nodes: servers responsible for running workloads using local and external resources.

The API server is the management point of the cluster, e.g., for configuring Kubernetes workloads and organizational units. The API interface is RESTful. A CLI client called `kubectl` is the default method of interacting with the Kubernetes cluster from a local computer.

The basic Kubernetes unit is the pod. Containers are assigned to pods—not hosts. Pods should contain a single application or related applications, e.g., ones that need to share the same filesystem.

## Rc<sup>2</sup> on Kubernetes

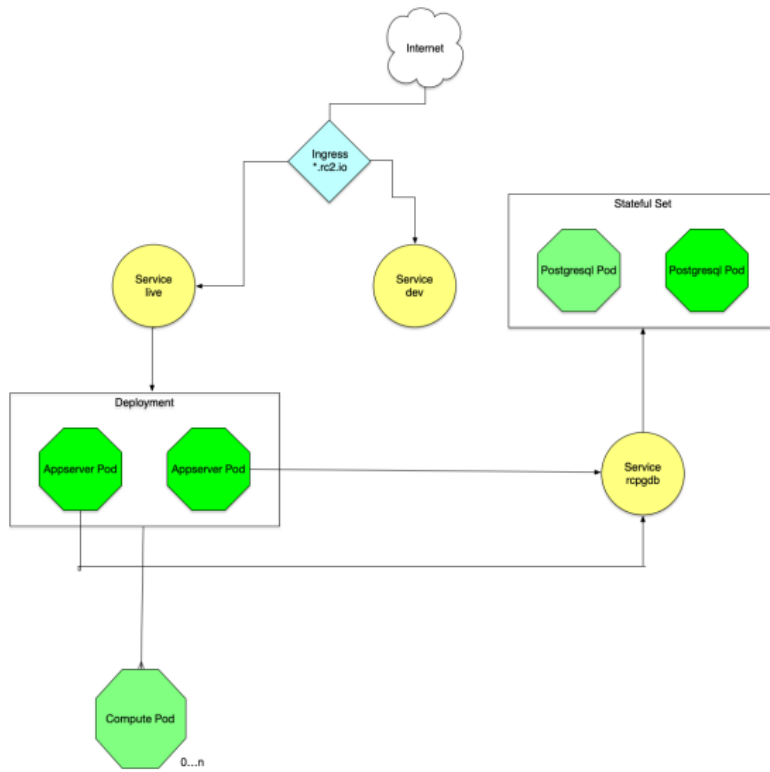
Rc<sup>2</sup> is a client-server environment for easily scaling R in a Kubernetes cluster. Other languages will be supported, e.g., Python and bash. Various services beyond R can be invoked, e.g., Spark and TensorFlow.

Rc<sup>2</sup> has two components:

- ▶ Rc<sup>2</sup> Server: a modern architecture for running R projects designed to interface with other environments at scale, e.g., Spark and Tensorflow
- ▶ Rc<sup>2</sup> Client: written in Swift currently running on macOS and iOS.

Rc<sup>2</sup> is being deployed to Digital Ocean Kubernetes.

# Rc<sup>2</sup> Architecture



# Rc<sup>2</sup> Client with an R File

The screenshot shows the Rc<sup>2</sup> ClientR interface. On the left, a sidebar lists 'Source Files' including dframe.R, mdtest.Rmd, newton.Rmd, and puromycin.R. The puromycin.R file is currently selected. The main area displays the R code and its execution results.

```
default
Source Files
dframe.R
mdtest.Rmd
newton.Rmd
puromycin.R
Other
mdtest.html
newton.html

1 conc <- c(.02, .02, .06, .06, .11, .11, .22, .22, .56, .56, 1.1, 1.1)
2 velocity <- c(76L, 47L, 97L, 107L, 123L, 139L, 159L, 152L, 191L,
201L, 207L, 208L)
3
4 puromycin.nls <- nls(velocity ~ (beta0 * conc) / (beta1 + conc),
+ start=c(beta0=200, beta1=0.1))
5 summary(puromycin.nls)
6
7 plot(conc, velocity, xlab="Concentration", ylab="Velocity")
8 conc.seq <- seq(min(conc), max(conc), length=100)
9 lines(conc.seq, predict(puromycin.nls, list(conc=conc.seq)), col =
"red", lwd = 2)
10
11 puromycin.df <- data.frame(conc=conc, velocity=velocity)
12 puromycin.df
13 str(puromycin.df)
14
15 ls()
16 rm(conc, velocity)
17 ls()
18
19 puromycin.nls <- nls(velocity ~ (beta0 * conc) / (beta1 + conc),
20 data=puromycin.df, start=c(beta0=200,
beta1=0.1))
21 puromycin.nls$data
22 puromycin.nls
```

Execution results:

```
> conc <- c(.02, .02, .06, .06, .11, .11, .22, .22, .56, .56, 1.1, 1.1)
> velocity <- c(76L, 47L, 97L, 107L, 123L, 139L, 159L, 152L, 191L, 201L, 207L, 208L)

> puromycin.nls <- nls(velocity ~ (beta0 * conc) / (beta1 + conc),
+ start=c(beta0=200, beta1=0.1))

> summary(puromycin.nls)

Formula: velocity ~ (beta0 * conc)/(beta1 + conc)

Parameters:
Estimate Std. Error t value Pr(>|t|)
beta0 2.127e+02 6.947e+00 30.615 3.24e-11 ***
beta1 6.412e-02 8.281e-03 7.743 1.57e-05 ***

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 10.93 on 10 degrees of freedom

Number of iterations to convergence: 6
Achieved convergence tolerance: 6.085e-06

> plot(conc, velocity, xlab="Concentration", ylab="Velocity")
> conc.seq <- seq(min(conc), max(conc), length=100)
> lines(conc.seq, predict(puromycin.nls, list(conc=conc.seq)), col =
"red", lwd = 2)

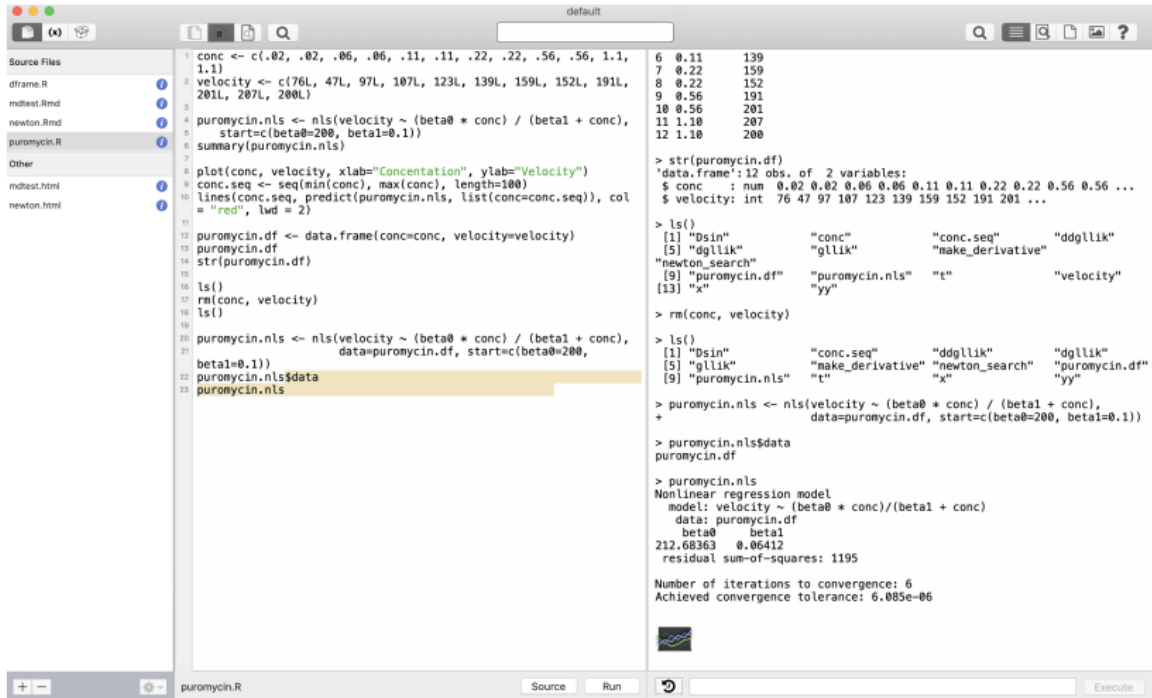
> puromycin.df <- data.frame(conc=conc, velocity=velocity)

> puromycin.df
  conc velocity
1 0.02      76
2 0.02      47
3 0.06      97
4 0.06     187
5 0.11     123
6 0.11     139
7 0.22     159
8 0.22     152
9 0.56     191
10 0.56     201
11 1.18     287
12 1.18     288
```

Bottom navigation bar: Source, Run, Execute.

Figure 2: Rc<sup>2</sup> ClientR

# Rc<sup>2</sup> Client with an R Image Icon



The screenshot shows the Rc<sup>2</sup> Client interface running on a Mac OS X desktop. The window title is "default". The left sidebar lists "Source Files" including dframe.R, mdtest.Rmd, newton.Rmd, and puromycin.R. The main pane displays R code for fitting a non-linear regression model to puromycin data. The code includes data loading, model definition, parameter estimation, and diagnostic output. A small R logo icon is visible in the bottom right corner of the client window.

```
1 conc <- c(.02, .02, .06, .06, .11, .11, .22, .22, .56, .56, 1.1, 1.1)
2 velocity <- c(76L, 47L, 97L, 107L, 123L, 139L, 159L, 152L, 191L, 261L, 267L, 260L)
3
4 puromycin.nls <- nls(velocity ~ (beta0 * conc) / (beta1 + conc),
5   start=c(beta0=200, beta1=0.1))
6 summary(puromycin.nls)
7
8 plot(conc, velocity, xlab="Concentration", ylab="Velocity")
9 conc.seq <- seq(min(conc), max(conc), length=100)
10 lines(conc.seq, predict(puromycin.nls, list(conc=conc.seq)), col = "red", lwd = 2)
11
12 puromycin.df <- data.frame(conc=conc, velocity=velocity)
13 puromycin.df
14 str(puromycin.df)
15
16 ls()
17 rm(conc, velocity)
18 ls()
19
20 puromycin.nls <- nls(velocity ~ (beta0 * conc) / (beta1 + conc),
21   data=puromycin.df, start=c(beta0=200,
22   beta1=0.1))
23 puromycin.nls$data
24 puromycin.nls
> str(puromycin.nls)
'data.frame': 12 obs. of  2 variables:
 $ conc : num  0.02 0.02 0.06 0.06 0.11 0.11 0.22 0.22 0.56 0.56 ...
 $ velocity: int  76 47 97 107 123 139 159 152 191 201 ...
> ls()
[1] "Dsin"           "conc"          "conc.seq"       "ddgllik"
[5] "dglilik"        "gllik"         "make_derivative"
[9] "newton_search"  "puromycin.df"  "puromycin.nls"  "t"
[13] "x"              "yy"
> rm(conc, velocity)
> ls()
[1] "Dsin"           "conc.seq"       "ddgllik"        "dglilik"
[5] "gllik"          "make_derivative" "newton_search"  "puromycin.df"
[9] "puromycin.nls"  "t"              "x"              "yy"
> puromycin.nls <- nls(velocity ~ (beta0 * conc) / (beta1 + conc),
+   data=puromycin.df, start=c(beta0=200, beta1=0.1))
> puromycin.nls$data
puromycin.nls
> puromycin.nls
Nonlinear regression model
  model: velocity ~ (beta0 * conc)/(beta1 + conc)
    data: puromycin.df
      beta0      beta1
  212.68363  0.06412
 residual sum-of-squares: 1195
Number of iterations to convergence: 6
Achieved convergence tolerance: 6.085e-06
```

Figure 3: Rc<sup>2</sup> ClientR2

# Rc<sup>2</sup> Client with an R Plot

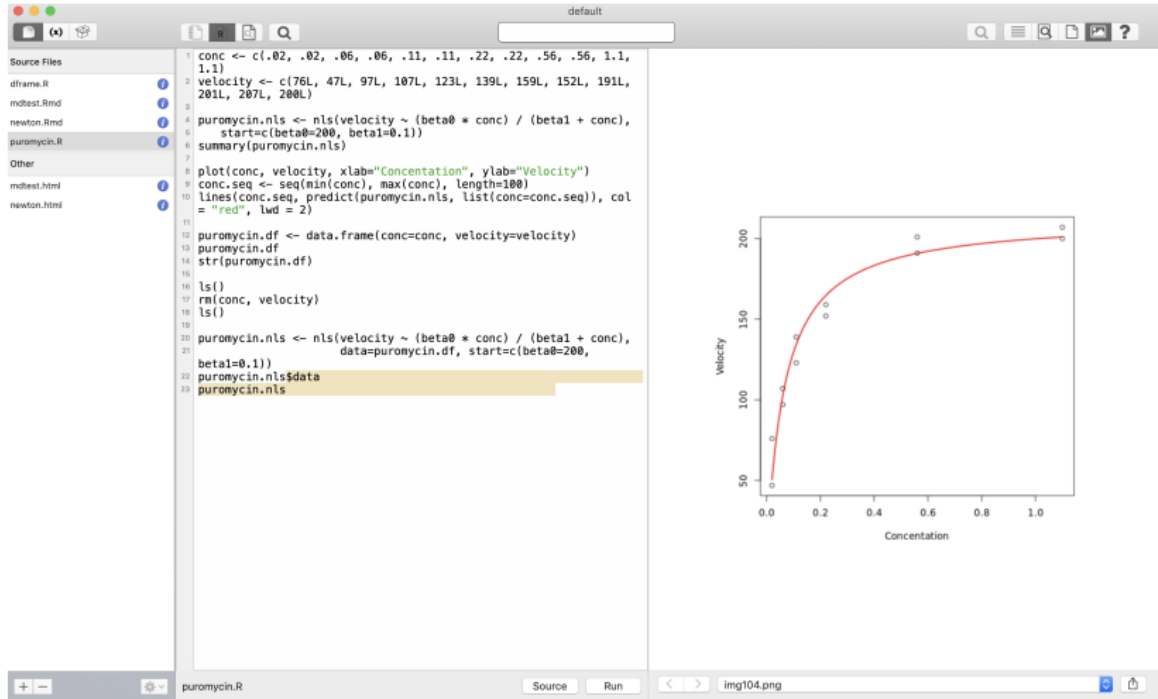


Figure 4: Rc<sup>2</sup> ClientR3

# Rc<sup>2</sup> Client with an Rmd File

The screenshot shows the Rc<sup>2</sup> Client interface with the following details:

- Source Files:** A sidebar listing files: dframe.R, ndtest.Rmd, newton.Rmd (selected), puromycin.R, Other, ndtest.html, and newton.html.
- Code Editor:** The main area displays R code for Newton's method. The code defines a function `newton_search` that takes four arguments: `f`, `df`, `guess`, and `conv`. It uses a while loop to iteratively improve the guess until the absolute value of the function at the guess is less than `conv`. It also includes a local function `improve` to calculate the next guess based on the current guess and the derivative at that point.
- Output Area:** To the right of the code editor, the output of the R code is shown. It includes the function definition, a note about potential infinite loops if `f` does not have a root, and the result of testing the function with `f = sin` and `df = cos`.
- Console:** At the bottom, there are buttons for Source and Run, and a navigation bar with back, forward, and search icons.
- File Status:** The status bar at the bottom indicates the file is "newton.Rmd" and the output is "newton.html".

Figure 5: Rc<sup>2</sup> ClientRmd

# Rc<sup>2</sup> Client with a Help File

The screenshot shows the Rc<sup>2</sup> Client interface with a help file open for the `plot` function. The left pane displays the R code for the `plot` function, with the word `plot` highlighted in blue. The right pane shows the help documentation for the `plot` function, which includes the function's name, a brief description, usage examples, arguments, and type information.

```
default
plot (graphics)
Generic X-Y Plotting

Description
Generic function for plotting of R objects. For more details about the graphical parameter arguments, see par.
For simple scatter plots, plot.default will be used. However, there are plot methods for many R objects, including functions, data.frames, density objects, etc. Use methods\(plot\) and the documentation for these.

Usage
plot(x, y, ...)

Arguments
x
  the coordinates of points in the plot. Alternatively, a single plotting structure, function or any R object with a plot method can be provided.
y
  the y coordinates of points in the plot, optional if x is an appropriate structure.
...
  Arguments to be passed to methods, such as graphical parameters (see par). Many methods will accept the following arguments:
  type
    what type of plot should be drawn. Possible types are
    • "p" for points,
    • "l" for lines,
    • "b" for both,
    • "c" for the lines part alone of "b",
    • "o" for both 'overplotted'.
```

Q Filter puromycin.R Source Run < > R: Generic X-Y Plotting

Figure 6: Rc<sup>2</sup> Help

# Rc<sup>2</sup> Client with a Search File



The screenshot shows a search interface with a search bar containing "coordinates". Below the search bar, there is a section titled "Arguments" which lists the following:

- x the **coordinates** of points in the plot. Alternatively, a single plotting structure, function or *any R object with a plot method* can be provided.
- y the y **coordinates** of points in the plot, *optional* if x is an appropriate structure.
- ... Arguments to be passed to methods, such as [graphical parameters](#) (see [par](#)). Many methods will accept the following arguments:

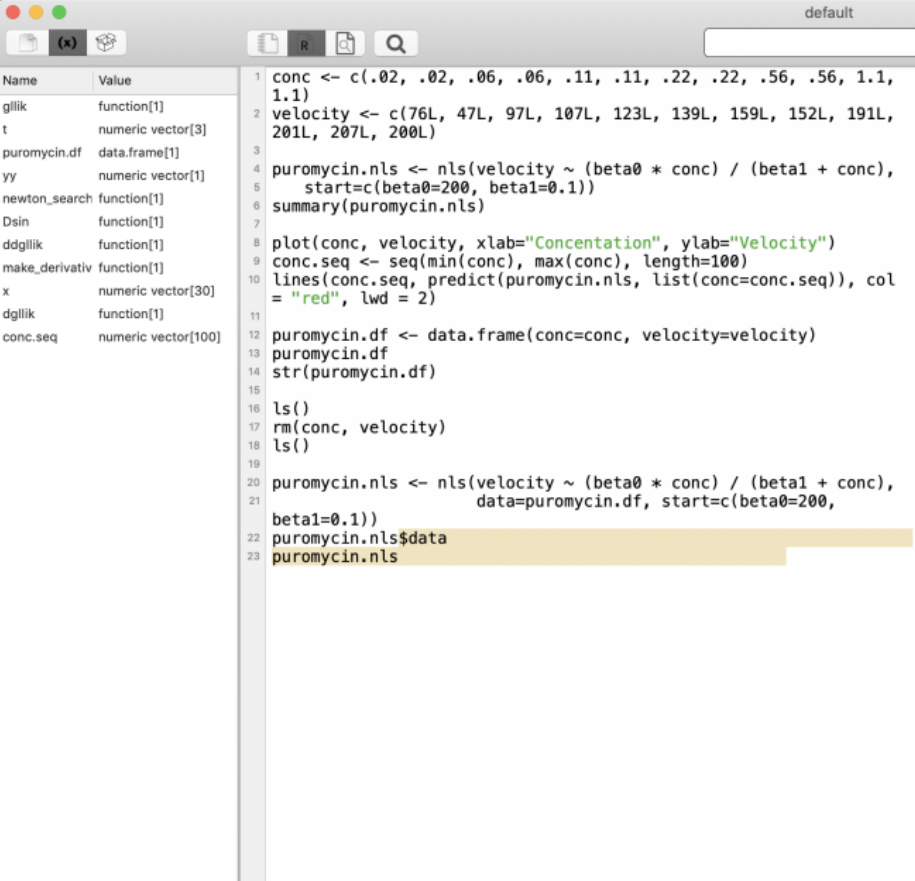
**type**

what type of plot should be drawn. Possible types are

- "p" for points,
- "l" for lines,
- "b" for both,
- "c" for the lines part alone of "b",
- "o" for both 'overplotted',
- "n" for 'histogram' like (or 'high-density') vertical lines,
- "s" for stair steps,
- "s" for other steps, see 'Details' below,
- "n" for no plotting.

All other types give a warning or an error; using, e.g., `type = "punkte"` being equivalent to `type = "p"` for S compatibility. Note that some methods, e.g. [`plot.factor`](#), do not accept this.

# Rc<sup>2</sup> Client with an Environment Display



Name	Value
gllik	function[1]
t	numeric vector[3]
puromycin.df	data.frame[1]
yy	numeric vector[1]
newton_search	function[1]
Dsin	function[1]
ddgllik	function[1]
make_derivativ	function[1]
x	numeric vector[30]
dgllik	function[1]
conc.seq	numeric vector[100]

```
1 conc <- c(.02, .02, .06, .06, .11, .11, .22, .22, .56, .56, 1.1,
2 velocity <- c(76L, 47L, 97L, 107L, 123L, 139L, 159L, 152L, 191L,
3 201L, 207L, 200L)
4 puromycin.nls <- nls(velocity ~ (beta0 * conc) / (beta1 + conc),
5 start=c(beta0=200, beta1=0.1))
6 summary(puromycin.nls)
7
8 plot(conc, velocity, xlab="Concentration", ylab="Velocity")
9 conc.seq <- seq(min(conc), max(conc), length=100)
10 lines(conc.seq, predict(puromycin.nls, list(conc=conc.seq)), col
= "red", lwd = 2)
11
12 puromycin.df <- data.frame(conc=conc, velocity=velocity)
13 puromycin.df
14 str(puromycin.df)
15
16 ls()
17 rm(conc, velocity)
18 ls()
19
20 puromycin.nls <- nls(velocity ~ (beta0 * conc) / (beta1 + conc),
21 data=puromycin.df, start=c(beta0=200,
beta1=0.1))
22 puromycin.nls$data
23 puromycin.nls
```

# Rc<sup>2</sup> Client with a Data Frame Display

The screenshot shows the Rc<sup>2</sup> Client interface with a data frame display. The interface has a top menu bar with icons for file operations and a title bar labeled "default". Below the title bar is a search bar. The main area contains three windows:

- puromycin.df**: A data frame window showing a table with two columns: "conc" and "velocity". The data consists of 12 rows of values.

	conc	velocity
1	0.02	76
2	0.02	47
3	0.06	97
4	0.06	107
5	0.11	123
6	0.11	139
7	0.22	159
8	0.22	152
9	0.56	191
10	0.56	201
11	1.1	207
12	1.1	200

- puromycin.nls**: A script editor window containing R code for non-linear least squares fitting. It includes a plot command with "conc" and "velocity" as arguments, and a call to "nls" with "start=c(beta0=200)".

```
16, .06, .11, .11, .22, .22, .56, .56, 1.1,  
.97L, 107L, 123L, 139L, 159L, 152L, 191L,  
  
velocity ~ (beta0 * conc) / (beta1 + conc),  
beta1=0.1))  
  
:lab="Concentration", ylab="Velocity")  
inc), max(conc), length=100)  
t(puromycin.nls, list(conc=conc.seq)), col  
  
frame(conc=conc, velocity=velocity)  
  
velocity ~ (beta0 * conc) / (beta1 + conc),  
data=puromycin.df, start=c(beta0=200,
```

## Rc<sup>2</sup> Deployment

Kubernetes is available on AWS, Azure, and of course Google Cloud as a service. However, these cloud providers do not have predictable price models and users/developers can incur large unexpected costs.

Digital Ocean Kubernetes gives the user far more control over these costs, i.e., limits can be set and the user is only charged for what is actually used.