

https://edcarp.github.io/introduction-to-conda-for-data-scientists

```
# Working with Environments
## Creating environments
conda create -- name python3-env python
conda create --name python36-env python=3.6
conda search scikit-learn
conda create --name basic-scipy-env ipython=7.13 matplotlib=3.1 numpy=1.18 scipy=
## Exercise: Creating a new environment
conda create -- name machine-learning-env \
 ipython=7.19 \
 matplotlib=3.3 \
 pandas=1.2 \
 python=3.8 \
 scikit-learn=0.23 \
 numba=0.51
## Activating an existing environment
conda activate basic-scipy-env
## Deactivate the active environment
conda deactivate
## Exercise: Activate an existing environment by name
conda activate machine-learning-env
conda deactivate
## Installing a package into an existing environment
conda activate basic-scipy-env
conda install numba
conda install scikit-learn=0.22
## Remove a package from an environment
conda uninstall scikit-learn -n basic-scipy-env
## Exercise: Installing a package into a specific environment
conda install --name machine-learning-env dask=2020.12
## Where do Conda environments live?
conda config -- show envs dirs
## How do I specify a location for a Conda environment?
conda create --prefix ./env ipvthon=7.13 matplotlib=3.1 pandas=1.0 pvthon=3.6
conda activate ./env
```

https://raw.githubusercontent.com/edcarp/introduction-to-conda-for-data-scientists/gh-pages/files/commands.txt

Introduction to Conda for Biologists: Timetable

Episode	Time	Website
Getting Started with Conda	9:30-10:00	https://edcarp.github.io/introduction-to-conda-for-data-scientists/01-getting-started-with-conda
Working with Environments	10:00-10:45	https://edcarp.github.io/introduction-to-conda-for-data-scientists/02-working-with-environments
Break	10:45-11:00	
Using Packages and Channels	11:00-12:00	https://edcarp.github.io/introduction-to-conda-for-data-scientists/03-using-packages-and-channels/index.html
Break	12:00-13:00	
Sharing Environments	13:00-13:45	https://edcarp.github.io/introduction-to-conda-for-data-scientists/04-sharing-environments
Configuring Conda	13:45-14:30	https://edcarp.github.io/introduction-to-conda-for-data-scientists/05-configuration/index.html

Getting Started with Conda

Getting Started with Conda: Questions

• What is conda?

Why use conda?

What is Conda?

Conda is an open source package management system and environment management system

Getting Started with Conda: Questions

 Why should I use a package and environment management system as part of my research workflow?

Packages

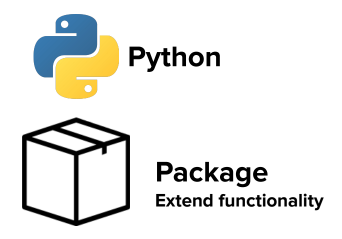
What are Packages?



What is Python used for?

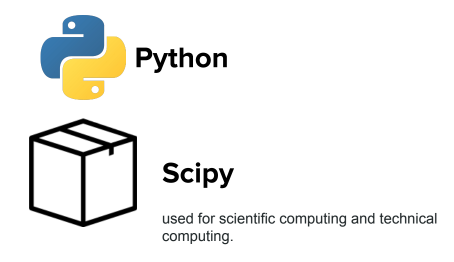
- Al and machine learning. ...
- Bioinformatics analysis ...
- Data visualisation. ...
- Programming applications. ...
- Web development. ...
- Game development. ...
- Finance.

What are Packages?

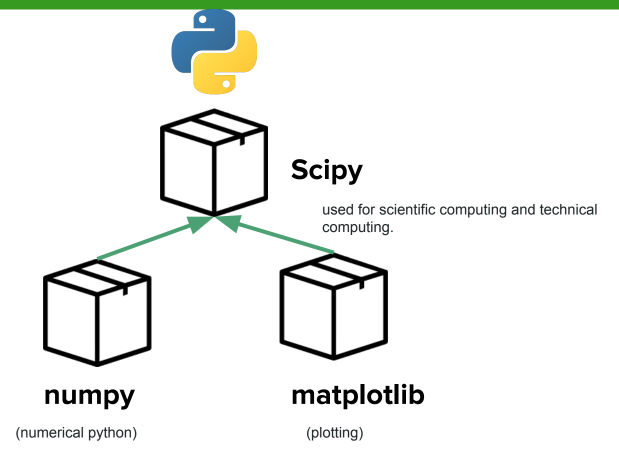


- Package: a collection of modules in a directory
 - a. Module: a collection of functions and variables, as in a script

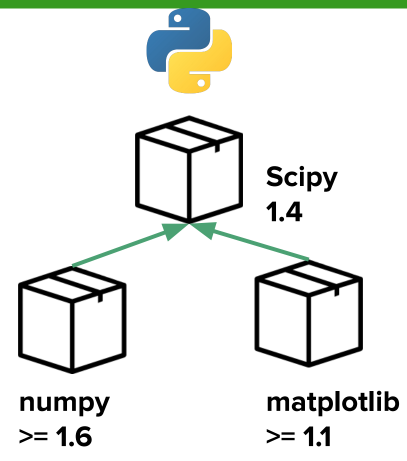
What are Packages?



Dependencies



Package Dependencies



Environment

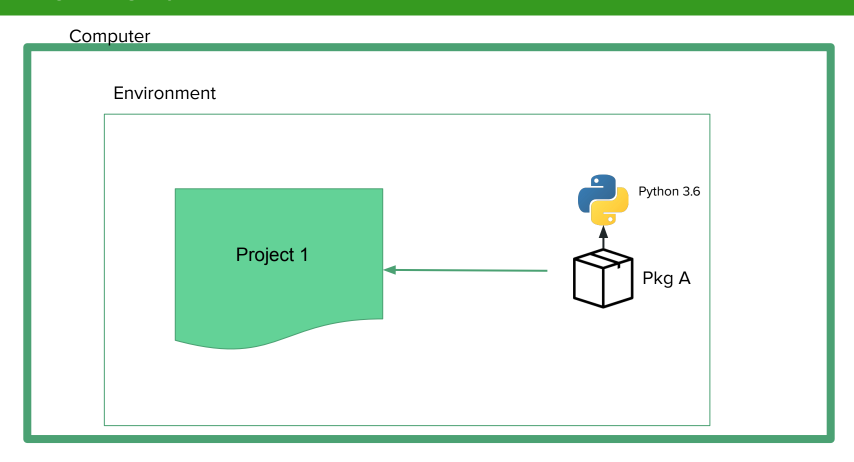
Environment

Computer

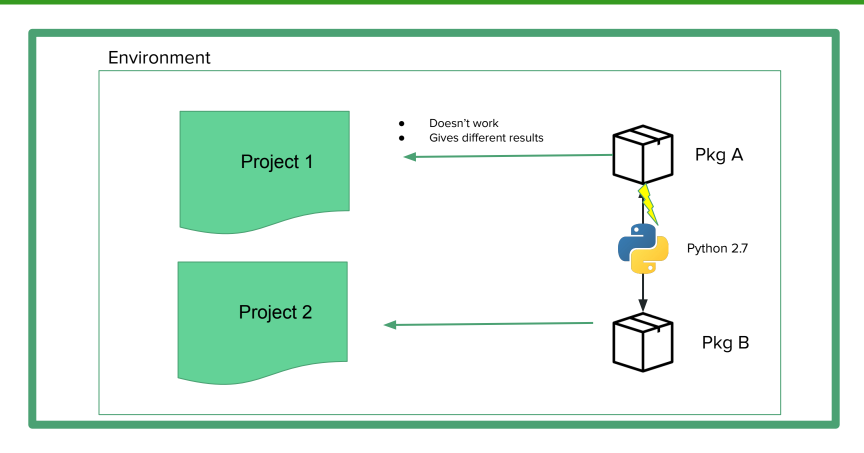
Environment

- All the Software
 - R, Python, C/C++, Java, sequence aligners ...
- Packages/Libraries
 - DESeq2, pysam, htslib
- Environment variables
 - PATH, RSTUDIO_WHICH_R

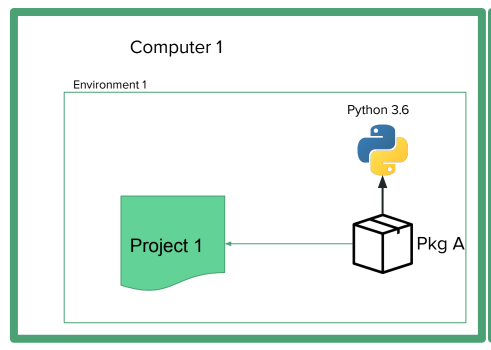
Environment

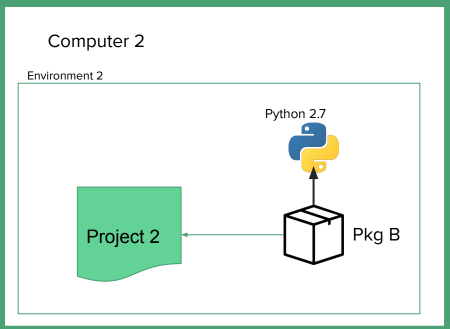


Environment: Update issues



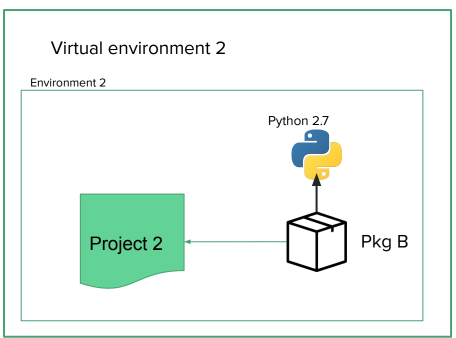
Solution: Separate Computers /VMs





Solution: Virtual Environments

Computer 1 Virtual environment 1 **Environment 1** Python 3.6 Pkg A Project 1



Environment Managers

Environment Management System

- Allow you to use different versions of a package for different projects.
- Make your projects self-contained and reproducible.
 - Capture packages and dependencies in a single file
- Can allow you to install packages on a host on which you do not have admin privilege

Solution: Environments Management

 Language specific
 Any Language

 Python
 R

 Venv, pipenv
 renv,packrat

 modules

Package management

Why should I use a Package Management System?

- Installing software can be hard.
 - Dependency hell

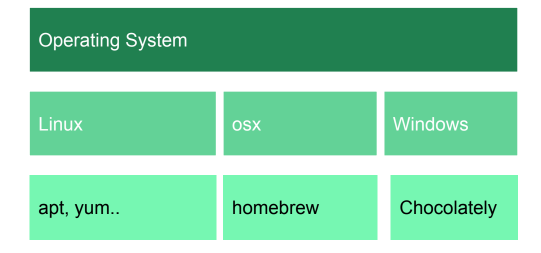
Installing bioinformatic software can take weeks!

Good Package Management tools

 Identifies and installing compatible versions of packages (software and all required dependencies).

 Handling the process of updating packages as more recent versions become available.

Package Management Tools



Packagement tools limitations

Installing software system wide has issues;

- It can be difficult to figure out what software is required for any particular research project.
- It is often impossible to install different versions of the same software package at the same time.
- Updating software required for one project can often "break" the software installed for another project.

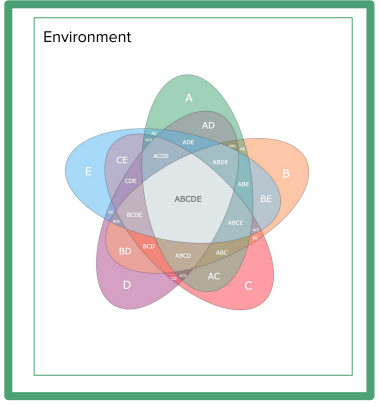
Packagement tools limitations

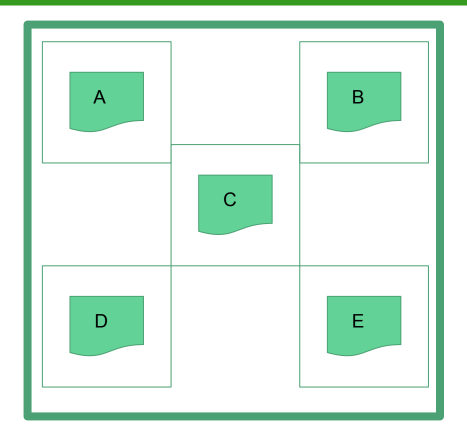
- As researcher you typically multiple scripting languages
 - o R, Python, SQL

- A project may require a different version of the scripting language than the one installed on your system.
 - Python v2 or v3

Project dependencies

Computer





Discussion

 In the etherpad write a potential benefits from installing software separately for each project?

• And write a potential costs?

Conda: Package & Environment Manager

Conda: Package Manager

- Package Manager:
 - o find,
 - o install,
 - o run,
 - update packages and their dependencies.

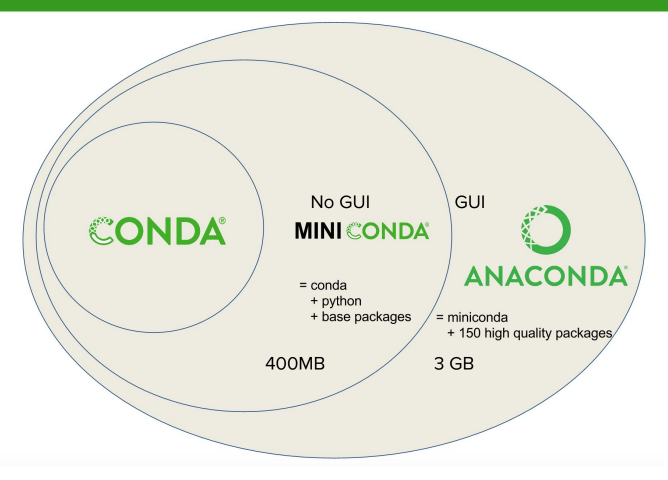
Conda: Environment Manager

- Environment Manager:
 - create,
 - o save,
 - load,
 - switch between project specific software environments on your local computer or HPC.

Conda: Package any software

- Although Conda was created for Python programs, Conda can package and distribute software for any language such as
 - R,
 - Ruby,
 - Lua,
 - Scala,
 - Java,
 - JavaScript,

Conda, Anaconda & Miniconda



Why use Conda?

- Conda solves both a package and environment manager and explicitly targeted at science use cases.
- Conda is an open source and free
- Conda is cross platform
- Conda can access dedicated repositories (channels) with thousands of prebuilt packages packages (no need to compile software).
 - Including a dedicated Bioinformatics package repository.

Why use Conda?

 Unlike some package managers e.g. pip, conda provides prebuilt packages don't need to be compiled.

 Conda makes sure all packages dependencies installed in an environment are met simultaneously before installing them.

Conda: Reproducibility

- Conda environments support reproducibility via a single environment files that describe their installation state.
- Conda is tightly integrated into popular solutions for reproducible data analysis such as
 - Snakemake
 - Nextflow
 - Galaxy,
 - bcbio-nextgen

Key Points

- Conda is a platform agnostic, open source package and environment management system.
- Using a package and environment management tool facilitates portability and reproducibility of data science workflows.
- Conda solves both the package and environment management problems and targets multiple programming languages.
 - Other open source tools solve either one or the other, or target only a particular programming language.

Working with Environments

What is a conda environment?

- What is a conda environment?
- How do I create (delete) an environment?

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- How do I find out what environments that exist on my machine?
- How do I delete an environment that I no longer need?

A Conda environment is a directory that contains a specific collection of Conda packages that you have installed.

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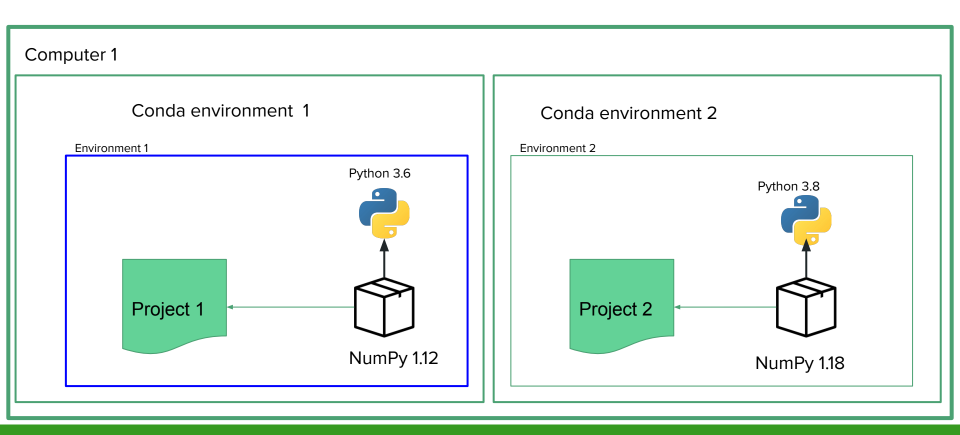
Conda can activate/deactivate different environments to give different versions of software

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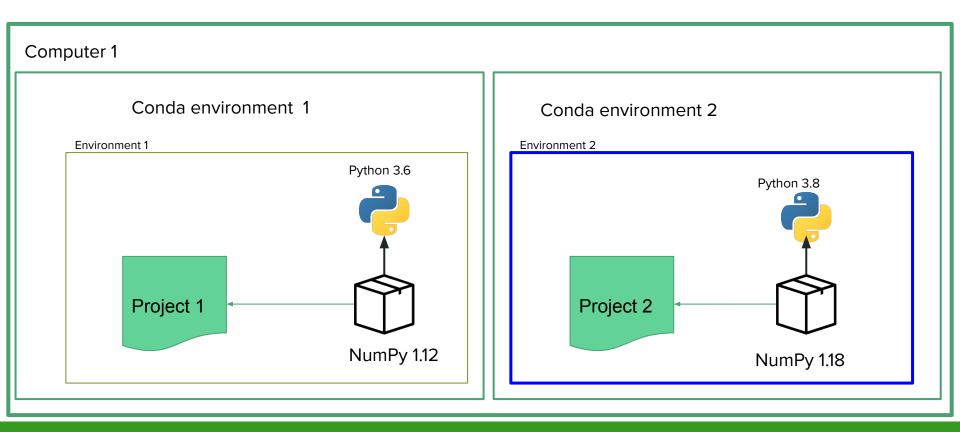
Conda can activate/deactivate different environments to give different versions of software

If you change one environment, your other environments are not affected

Project 1 requires NumPy 1.12 and Python 3.6



Deactivate Project 1 and activate Project for project that requires NumPy 1.18 and Python 3.8



To create a new environment for Python development using conda you can use the conda create command.

\$ conda create

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Add an environment name using --name option

\$ conda create --name python3-env

It is a good idea to give your environment a meaningful name in order to help yourself remember the purpose of the environment.

While naming things can be difficult, **PROJECTNAME**—env is a good convention to follow.

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\$ conda create --name python3-env

Sometimes also the specific version of a package why you had to create a new environment is a good name

To create a new environment for Python development using conda you can use the conda create command.

Add a package name

\$ conda create --name python3-env python

Working with Environments: Specify version

The previous command above will create a new Conda environment called "python3-env" and install the most recent version of Python.

If you wish, you can specify a particular version of packages for conda to install when creating the environment =<major.minor.patch>.

\$ conda create --name python36-env python=3.6

You can create a Conda environment and install multiple packages by listing the packages that you wish to install.

New environment named basic-scipy-env

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New environment named basic-scipy-env

\$ conda create --name basic-scipy-env ipython=7.13 matplotlib=3.1 numpy=1.18 scipy=1.4

Working with Environments: Dependencies

When conda installs a package into an environment it also installs any required dependencies.

For example, even though Python is not listed as a packaged to install into the **basic-scipy-env** environment above, conda will still install Python into the environment because it is a required dependency of at least one of the listed packages.

Exercise: Creating a new environment

Create a new environment named **machine-learning-env** with Python and the most current versions of

- IPython
- Matplotlib
- Pandas
- Numba
- Scikit-Learn

Hint to install latest compatible version do not specify a version number

https://edcarp.github.io/introduction-to-conda-for-data-scientists/02-working-with-environments/index.html

Activating environments is essential to making the software in environments work

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- 1. Adds entries to PATH for the environment.
- 2. Runs any activation scripts that the environment may contain.

Activating environments is essential to making the software in environments work

Activation of an environment does two things.

- 1. Adds entries to \$PATH for the environment.
- 2. Runs any activation scripts that the environment may contain.

Step 2 is particularly important as activation scripts are how packages can set arbitrary environment variables that may be necessary for their operation.

You activate the basic-scipy-env environment by name using the activate command.

\$ conda activate basic-scipy-env

You can see that an environment has been activated because the shell prompt will now include the name of the active environment.

```
(basic-scipy-env) $
```

Working with Environments: Deactivating Envs

To deactivate the currently active environment use the Conda deactivate command as follows.

(basic-scipy-env) \$ conda deactivate

Working with Environments: Deactivating Envs

You can see that an environment has been deactivated because the shell prompt will no longer include the name of the previously active environment.

```
(base) $
```

Working with Environments: Deactivating Envs

To return to the base Conda environment, it's better to call conda activate with no environment specified, rather than to use deactivate.

If you run conda deactivate from your base environment, you may lose the ability to run conda commands at all

```
$ conda activate basic-scipy-env
```

```
$ (basic-scipy-env) $ conda activate
```

Exercise: Activate an existing environment by name

1. Activate the machine-learning-env environment created in the previous challenge by name.

2. Deactivate the machine-learning-env environment that you activated in the previous challenge.

You can install a package into an existing environment using the **conda install** command.

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This command accepts a list of package specifications (i.e., numpy=1.18) and installs a set of packages consistent with those specifications and compatible with the underlying environment.

(my_active-env) \$ conda install numpy=1.18

You can install a package into an existing environment using the **conda install** command.

This command accepts a list of package specifications (i.e., numpy=1.18) and installs a set of packages consistent with those specifications and compatible with the underlying environment.

If full compatibility cannot be assured, an error is reported and the environment is not changed.

By default the conda install command will install packages into the current, active environment.

The following would activate the **basic-scipy-env** we created above and install

```
$ conda activate basic-scipy-env
```

(basic-scipy-env) \$ conda install numba

As was the case when listing packages to install when using the conda create command, if version numbers are not explicitly provided, Conda will attempt to install the newest versions of any requested packages.

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To accomplish this, Conda may need to update some packages that are already installed or install additional packages.

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To accomplish this, Conda may need to update some packages that are already installed or install additional packages.

It is always a good idea to explicitly provide version numbers when installing packages with the conda install command.

The following would install a particular version of scikit-learn, into the current, active environment.

(basic-scipy-env) \$ conda install scikit-learn=0.22

Working with Environments: Remove a package from an environment

To remove a package from an environment you can run the command.

\$ conda uninstall PKGNAME --name ENVNAME

Working with Environments: Remove a package from an environment

To remove the scikit-learn package from the basic-scipy-env environment run

\$ (basic-scipy-env) conda uninstall scikit-learn -n basic-scipy-env

Exercise: Installing a package into a specific environment

Dask provides advanced parallelism for data science workflows enabling performance at scale for the core Python data science tools such as Numpy Pandas, and Scikit-Learn.

Run

\$ conda install --help

And figure mout how to Install dask into the machine-learning-env that you created in the previous challenge.

Working with Environments: Where do Conda environments live?

Environments created with conda, by default, live in the **envs/** folder of your miniconda3 (or anaconda3) directory the absolute path to which will look something the following:

/home/training/miniconda3/envs

C:\Users\\$USERNAME\Anaconda3

Working with Environments: Where do Conda environments live?

You can see the location of your conda environments by running the command.

\$ conda config --show envs_dirs

Working with Environments: Where do Conda environments live?

Running **ls** (linux) / **dir** (Windows) on your anaconda envs/ directory will list out the directories containing the existing Conda environments.

\$ ls /home/training.miniconda3/envs

You can control where a Conda environment lives by providing a path to a target directory when creating the environment.

The following command will create a new environment in a sub-directory of the current working directory called **env**

```
$ pwd
$ conda create --prefix ./env
```

You can control where a Conda environment lives by providing a path to a target directory when creating the environment.

For example the following command will create a new environment in a sub-directory of the current working directory called **env**

```
$ pwd
```

^{\$} conda create --prefix ./env ipython=7.13 matplotlib=3.1 pandas=1.0 python=3.6

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For example the following command will create a new environment in a sub-directory of the current working directory called **env**

```
$ conda create --prefix ./env ipython=7.13 matplotlib=3.1 pandas=1.0 python=3.6

In unix the dot-slash, ./, is a relative path a file or directory in the current directory.
```

It is often a good idea to specify a path to a sub-directory of your project directory when creating an environment. Why?

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Makes it easy to tell if your project utilizes an isolated environment by including the environment as a sub-directory.

Makes your project more self-contained as everything including the required software is contained in a single project directory.

An additional benefit of creating your project's environment inside a sub-directory is that you can then use the same name for all your environments; if you keep all of your environments in your $\sim/\miniconda3/env/$ folder, you'll have to give each of them a different name.

Exercise: Creating a new environment as a sub-directory within a project directory environment

First create a project directory called project-dir using the following command.

```
$ mkdir project-dir
```

\$ cd project-dir

Next, create a new environment inside the newly created project-dir in a sub-directory called env an install

- Python 3.6,
- Matplotlib 3.1
- TensorFlow 2.0

Placing Conda environments outside of the default ~/miniconda3/envs/ folder comes with a couple of minor drawbacks.

1. conda can no longer find your environment with the --name flag; you'll generally need to pass the --prefix flag along with the environment's full path to find the environment.

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1. conda can no longer find your environment with the --name flag; you'll generally need to pass the --prefix flag along with the environment's full path to find the environment.

2. Second, an annoying side-effect of specifying an install path when creating your Conda environments is that your command prompt is now prefixed with the active environment's absolute path rather than the environment's name. After activating an environment using its prefix your prompt will look similar to the following.

(/absolute/path/to/env) \$

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2. Second, an annoying side-effect of specifying an install path when creating your Conda environments is that your command prompt is now prefixed with the active environment's absolute path rather than the environment's name. After activating an environment using its prefix your prompt will look similar to the following.

If (you find this long prefix to your shell prompt annoying, then there is a quick fix: modify the env_prompt setting in your .condarc file, which you can do with the following command.

```
$ conda config --set env_prompt '({name})'
```

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```
$ conda config --set env_prompt '({name})'
```

This will either edit your ~/.condarc file if you already have one or create a ~/.condarc file if you do not.

Now your command prompt will display the active environment's generic name.

```
$ cd project-directory
$ conda activate ./env
(env) project-directory $
```

Exercise: Activate an existing environment by path

- 1. Activate the environment created in a previous challenge. project-dir. using the path to the environment directory.
- 2. Create an environments for R project

First create a project directory called r-project-dir using the following command.

```
cd ~/
mkdir r-project-dir
cd r-project-dir
```

Create a new environment inside the newly created r-project-dir in a sub-directory called env and install

- r-base
- r-tidyverse

Working with Environments: Listing existing environments

Now that you have created a number of Conda environments on your local machine you have probably forgotten the names of all of the environments and exactly where they live.

This a conda command lists all of your existing environments together with their locations.

\$ conda env list

Working with Environments: Listing the contents of environments

You will probably forget exactly what has been installed in a particular Conda environment.

To list the contents of the basic-scipy-env that you created above, run the following command.

```
$ conda list --name basic-scipy-env
```

Working with Environments: Listing the contents of environments

If you created your Conda environment using the **--prefix** option to install packages into a particular directory, then you will need to use that prefix in order for conda to locate the environment on your machine.

```
$ conda list --prefix /path/to/conda-env
```

Exercise: Listing the contents of a particular environment.

List the packages installed in the machine-learning-env environment that you created in a previous challenge.

Working with Environments: Deleting entire environments

Occasionally, you will want to delete an entire environment. The command to delete an environment is the following.

```
$ conda remove --name python36-env --all --dry-run
```

```
$ conda remove --name python36-env --all
```

- --all Remove all packages, i.e., the entire environment.
- --dry-run Only display what would have been done.

Working with Environments: Deleting entire environments

If you wish to delete and environment that you created with a --prefix option, then you will need to provide the prefix again when removing the environment.

```
$ conda remove --prefix /path/to/conda-env/ --all
```

Exercise: Delete an entire environment

Delete the entire machine-learning-env environment.

Hint try --dry-run first

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- You install packages into environments using conda install;
- You should install each environment as a sub-directory inside its corresponding project directory
- Use the conda env list command to list existing environments and their respective locations.

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 commands.
- You activate (deactivate) an environment using the conda activate (conda deactivate) commands.
- You install packages into environments using conda install;
- You should install each environment as a sub-directory inside its corresponding project directory
- Use the conda env list command to list existing environments and their respective locations.
- Use the conda list command to list all of the packages installed in an environment.



Using Packages and Channels: Questions

- What are Conda packages?
- What are Conda channels?
- Why should I be explicit about which channels my research project uses?

Using Packages and Channels: What are Packages?

A conda package is a compressed archive file (.tar.bz2) that contains:

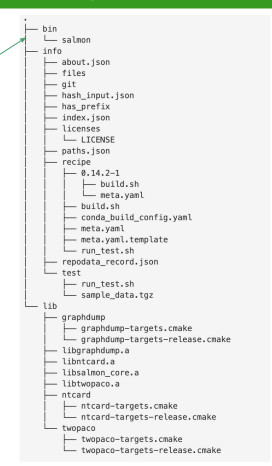
- system-level libraries
- Python or other modules
- executable programs and other components
- metadata under the info/ directory
- a collection of files that are installed directly into an install prefix.

Conda keeps track of the dependencies between packages and platforms; the conda package format is identical across platforms and operating systems.

Using Packages and Channels: Package Structure?

All conda packages have a specific sub-directory structure inside the tarball file.

There is a bin directory that contains any binaries for the package;



Using Packages and Channels: Package Structure?

All conda packages have a specific sub-directory structure inside the tarball file.

There is a bin directory that contains any binaries for the package;

a lib directory containing the relevant library files (i.e., the .py files);

and an info directory containing package metadata.

```
∟ salmon
  about.ison
 files
  ait
  - hash_input.json
  has prefix
  index.json
   licenses
   LICENSE
   paths. ison
      - 0.14.2-1
         build.sh
         meta.yaml
       build.sh
       conda_build_config.yaml

    meta.vaml

     meta.yaml.template
     run test.sh
   repodata record.json
  test
    — run test.sh

    sample data.tgz

    graphdump-targets.cmake

    graphdump-targets-release.cmake

  libgraphdump.a
  libntcard.a
  libsalmon core.a
  libtwopaco.a
  ntcard
    — ntcard-targets.cmake
    — ntcard-targets-release.cmake
    — twopaco-targets.cmake

    twopaco-targets-release.cmake
```

Using Packages and Channels: Package Structure?

All conda packages have a specific sub-directory structure inside the tarball file.

There is a bin directory that contains any binaries for the package;

a lib directory containing the relevant library files (i.e., the .py files);

and an info directory containing package metadata.

```
∟ salmon
    about.ison
    - files
    ait
    hash_input.json
    has_prefix
    index.json
     licenses
     LICENSE
     paths.ison
       - 0.14.2-1
           build.sh
           meta.yaml
         build.sh
         conda_build_config.yaml
        meta.vaml
       meta.yaml.template
       run test.sh
     repodata record.json
    test
      run test.sh

    sample data.tgz

- lib
     graphdump
       graphdump-targets.cmake

    graphdump-targets-release.cmake

    libgraphdump.a
    libntcard.a
    libsalmon core.a
    libtwopaco.a
    ntcard
      — ntcard-targets.cmake

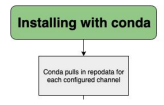
    ntcard-targets-release.cmake

      — twopaco-targets.cmake

    twopaco-targets-release.cmake
```

Installing with conda

\$ conda install super-fun-package=0

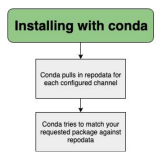


Conda pulls in repodata for each configured channel

- https://repo.anaconda.com/pkgs/main/linux-64
- https://repo.anaconda.com/pkgs/main/noarch
- https://repo.anaconda.com/pkgs/r/linux-64
- https://repo.anaconda.com/pkgs/r/noarch

https://repo.anaconda.com/pkgs/main/linux-64/repodata.json

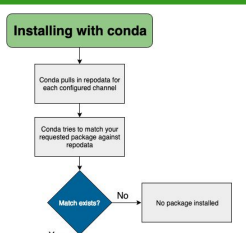
\$ conda install super-fun-package=0

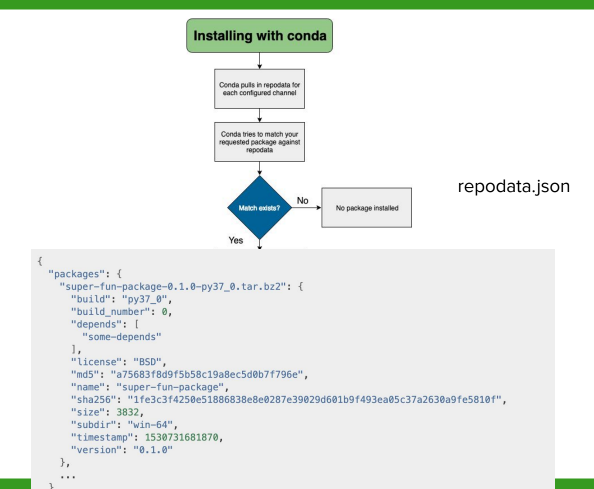


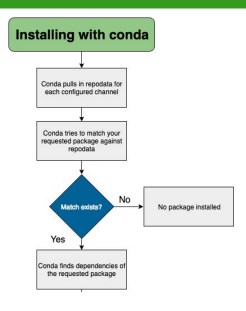
- https://repo.anaconda.com/pkgs/main/lenux-64
- https://repo.anaconda.com/pkgs/main/norch
- https://repo.anaconda.com/pkgs/r/linux-64
- https://repo.anaconda.com/pkgs/r/noarch

repodata.json

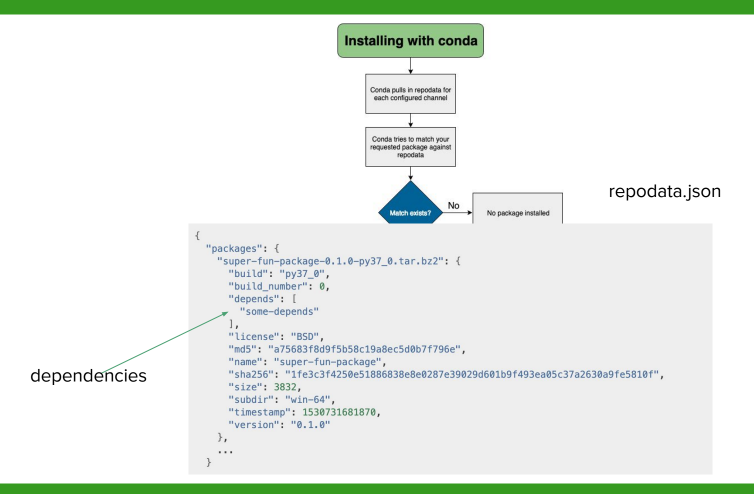
Conda tries to match your requested package against repodata

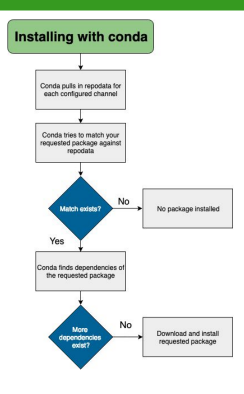


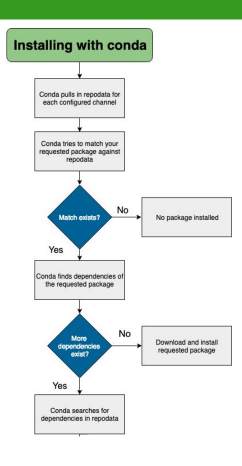


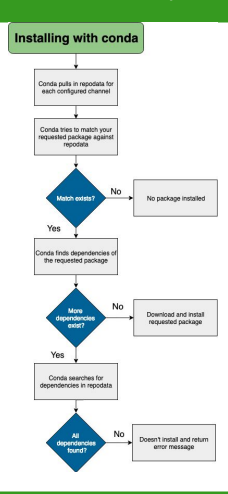


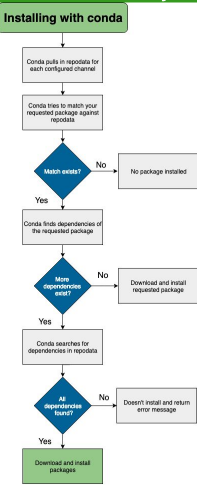
Conda find dependencies of requested package











Using Packages and Channels: Installing salmon package

Let's create a new environment <code>basic-rnaseq-env</code> and install a transcript quantification package, salmon , for an RNA-Seq analysis project.

\$ conda create --name basic-rnaseq-env salmon

Using Packages and Channels: Installing salmon package

```
Loading channels: done
No match found for: salmon, Search: *salmon*
PackagesNotFoundError: The following packages are not available from current channels:
  salmon
Current channels:
  - https://repo.anaconda.com/pkgs/main/linux-64
  - https://repo.anaconda.com/pkgs/main/noarch
  - https://repo.anaconda.com/pkgs/r/linux-64
  - https://repo.anaconda.com/pkgs/r/noarch
To search for alternate channels that may provide the conda package you're
looking for, navigate to
    https://anaconda.org
and use the search bar at the top of the page.
```

When you install or search for a package in conda it searches for it in remote repositories called channels.

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These remote channel are URLs to directories containing conda packages.

Current channels:

- https://repo.anaconda.com/pkgs/main/linux-64
- https://repo.anaconda.com/pkgs/main/noarch
- https://repo.anaconda.com/pkgs/r/linux-64
- https://repo.anaconda.com/pkgs/r/noarch

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- https://repo.anaconda.com/pkgs/r/noarch

```
Current channels:
    - https://repo.anaconda.com/pkgs/main/linux-64
    - https://repo.anaconda.com/pkgs/main/noarch
    - https://repo.anaconda.com/pkgs/r/linux-64
```

By default the conda search command searches a set of channels defined here. Anaconda Cloud channels.

• main: The majority of all new Anaconda, Inc. package builds are hosted here. Included in conda's defaults channel as the top priority channel.

When you install or search for a package in conda it searches for it in remote repositories called channels.

These remote channel are URLs to directories containing conda packages.

Current channels:

- https://repo.anaconda.com/pkgs/main/linux-64
- https://repo.anaconda.com/pkgs/main/noarch
- https://repo.anaconda.com/pkgs/r/linux-64
- https://repo.anaconda.com/pkgs//r/noarch

By default the conda search command searches a set of channels defined here. Anaconda Cloud channels.

- main: The majority of all new Ariaconda, Inc. package builds are hosted here. Included in conda's defaults channel as the top priority channel.
- r: Microsoft R Open conda packages and Anaconda, Inc.'s R conda packages. This channel is included in conda's defaults channel. When creating new environments, MRO is now chosen as the default R implementation.

Collectively, the Anaconda managed channels are referred to as the defaults channel because, unless otherwise specified, packages installed using conda will be downloaded from these channels.

Using Packages and Channels: My package isn't available on the defaults

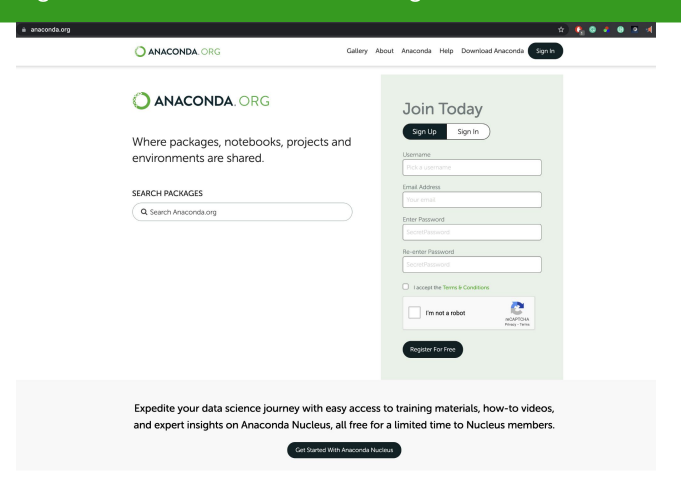
What should I do?

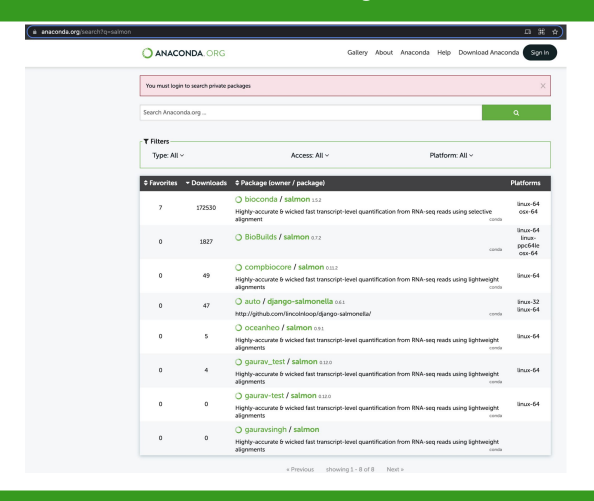
Using Packages and Channels: My package isn't available on the defaults

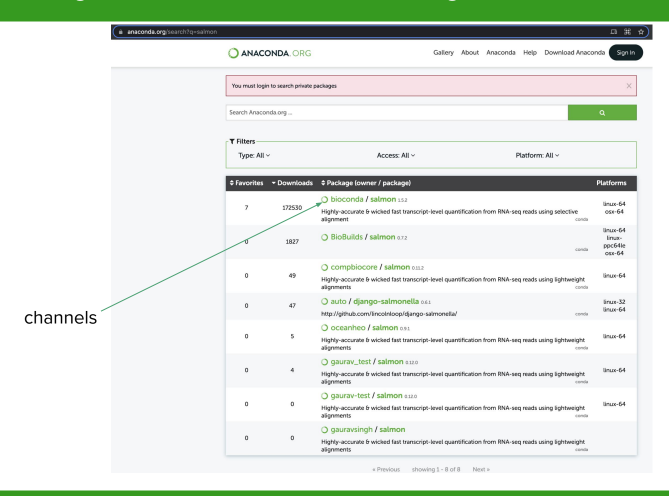
```
Loading channels: done
No match found for: salmon, Search: *salmon*
PackagesNotFoundError: The following packages are not available from current channels:
  salmon
Current channels:
  - https://repo.anaconda.com/pkgs/main/linux-64
  - https://repo.anaconda.com/pkgs/main/noarch
  - https://repo.anaconda.com/pkgs/r/linux-64
  - https://repo.anaconda.com/pkgs/r/noarch
To search for alternate channels that may provide the conda package you're
```

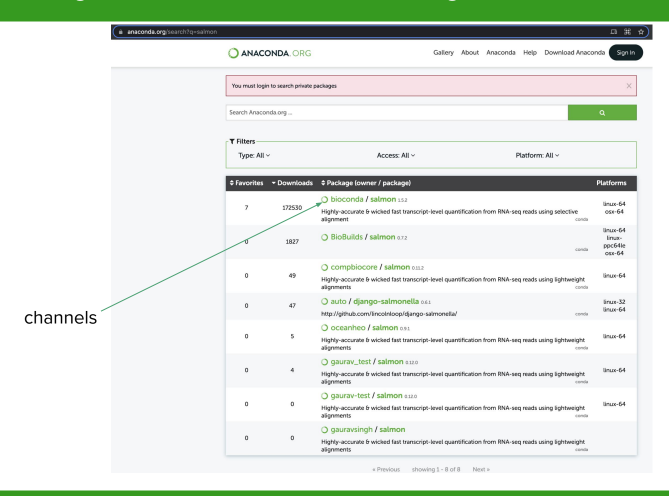
https://anaconda.org
and use the search bar at the top of the page.

looking for, navigate to



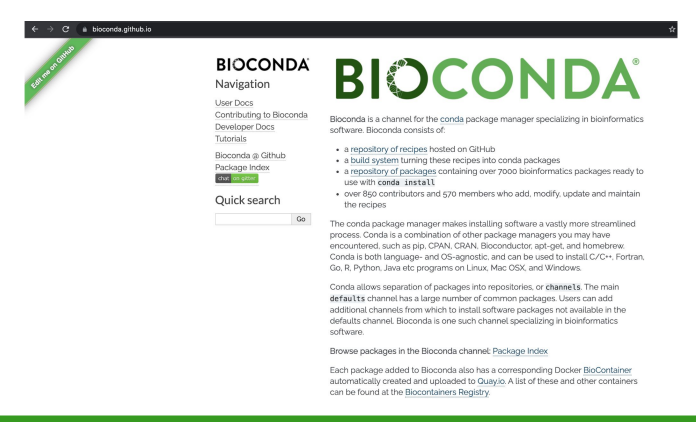






Using Packages and Channels: bioconda

Bioconda is a channel, maintained by the <u>Bioconda project</u>, specialising in bioinformatics software. Bioconda contains 1000's of bioinformatics packages ready to use with conda install.

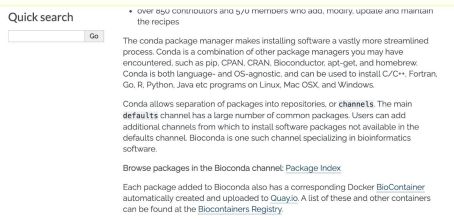


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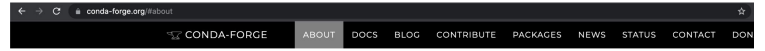


R and Bioconductor packages Most R packages on CRAN should be submitted at Conda-Forge. However, if the CRAN package has a Bioconductor, a repository for bioinformatics R packages, dependency, it belongs in Bioconda. If the CRAN package does not have a Bioconductor package dependency, it belongs in Conda-Forge.



In addition to the default channels that are managed by Anaconda Inc., there is another channel called Conda-Forge that also has a special status.

The Conda-Forge project "is a *community* led collection of recipes, build infrastructure and distributions for the conda package manager."



ABOUT CONDA-FORGE

conda-forge is a GitHub organization containing repositories of conda recipes. Thanks to some awesome continuous integration providers (AppVeyor, Azure Pipelines, CircleCl and TravisCl), each repository, also known as a feedstock, automatically builds its own recipe in a clean and repeatable way on Windows, Linux and OSX.

There are a number of reasons that you may wish to use the <code>conda-forge</code> channel instead of the <code>defaults</code> channel maintained by Anaconda:

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1. Packages on conda-forge may be more up-to-date than those on the defaults channel.

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- 2. There are packages on the conda-forge channel that aren't available from defaults.

There are a number of reasons that you may wish to use the conda-forge channel instead of the defaults channel maintained by Anaconda:

- 1. Packages on conda-forge may be more up-to-date than those on the defaults channel.
- 2. There are packages on the conda-forge channel that aren't available from defaults.
- 3. You may wish to use a dependency such as openblas (from conda-forge) instead of mkl(from defaults).

Using Packages and Channels: Search Channel

If you know the channel your package is likely to be located on, you can can use the conda search command with the --channel option and the name of the channel.

\$ conda search --channel bioconda salmon

Using Packages and Channels: Install from Channel

If you know the channel your package is available from you can install a package from a specific channel into the currently activate environment by passing the --channel or -c option to the conda install command as follows.

- \$ conda create --name basic-rnaseq-env
- \$ conda activate basic-rnaseq-env
- \$ conda install --channel bioconda salmon

Using Packages and Channels: Install from Channel

You can also install a package from a specific channel into a named environment (using --name or -n)

Or into an environment installed at a particular prefix (using --prefix or -p).

```
$ conda install salmon --channel bioconda --name basic-rnaseq-env
```

```
$ conda install salmon --channel bioconda --prefix ./env
```

Where would this command would install salmon package? What version of salmon is installed?

Using Packages and Channels: Install from Channel

You may have noticed that we didn't manage to install the latest version for salmon, why?

The bioconda channel contains bioinformatics packages (salmon, STAR, samtools, DESeq2, etc.), however the channel conda-forge has most of the dependencies (numpy, scipy, zlib, CRAN packages, etc.) needed.

We need to specify multiple channels to install the latest version.

Using Packages and Channels: Multiple Channels

To specify multiple channels for installing packages by passing the --channel argument multiple times.

```
$ conda install salmon=1.5 --channel conda-forge --channel bioconda --name basic-rnaseq-env
```

Using Packages and Channels: Multiple Channels

To specify multiple channels for installing packages by passing the --channel argument multiple times.

This also works when install multiple packages.

```
$ conda install salmon=1.5 --channel conda-forge --channel bioconda --name basic-rnaseq-env
$ conda install fastqc=0.11 multiqc=1.10 --channel conda-forge --channel bioconda --name
basic-rnaseq-env
```

Exercise: Specifying channels when installing packages

Create a new directory called rnaseq-project and then create an environment in a sub-directory called env/ with the the packages

- salmon=1.5
- fastqc=0.11
- multiqc=1.11

Hint: For the lazy typers: the --channel argument can also be shortened to -c

Exercise: Alternative syntax for installing packages from specific channels

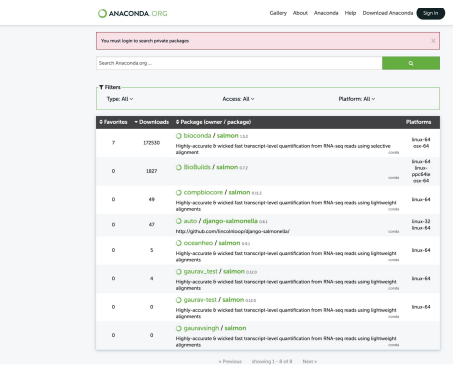
There exists an alternative syntax for installing conda packages from specific channels that more explicitly links the channel being used to install a particular package.

```
channel::<package_name>
$ conda install biconda::multiqc --prefix ./env
```

Install the latest version of the workflow manager **nextflow** using this alternative syntax

Different channels can have the same package, so conda must decide which channel to install the package from.

Conda channels have a priority hierarchy.



Different channels can have the same package, so conda must decide which channel to install the package from. Conda channels have a priority hierarchy.

By default, conda prefers packages from a higher priority channel over any version from a lower priority channel.

Channel priority listed on the command line decreases from left to right.

So if you were to install base R r-base using the command below.

```
$ conda create --name rproject-env --channel defaults --channel conda-forge r-base
```

The first channel defaults would have a higher priority than the second conda-forge.

This is true even, if the version number of the package is higher in the second channel.

Different channels can have the same package, so conda must decide which channel to install the package from. Conda channels have a priority hierarchy.

By default, conda prefers packages from a higher priority channel over any version from a lower priority channel.

Higher Priority

\$ conda install salmon=1.5 --channel conda-forge --channel bioconda --name
basic-rnaseq-env

The bioconda team suggests that the conda-forge channel be a higher priority than the bioconda channel.

Different channels can have the same package, so conda must decide which channel to install the package from. Conda channels have a priority hierarchy.

By default, conda prefers packages from a higher priority channel over any version from a lower priority channel.

Conda collects all of the packages with the same name across all listed channels and processes them as follows:

1. Sorts packages from highest to lowest channel priority.

- Sorts packages from highest to lowest channel priority.
- 2. Sorts tied packages—packages with the same channel priority—from highest to lowest version number.
 - a. For example, if channelA contains NumPy 1.12.0 and 1.13.1, NumPy 1.13.1 will be sorted higher.

Sorts packages from highest to lowest channel priority.

- Sorts tied packages—packages with the same channel priority—from highest to lowest version number. For example, if channelA contains NumPy 1.12.0 and 1.13.1, NumPy 1.13.1 will be sorted higher.
- 3. Sorts still-tied packages, packages with the same channel priority and same version, from highest to lowest build number.
 - a. For example, if channelA contains both NumPy 1.12.0 build 1 and build 2,
 - b. build 2 is sorted first.
 - c. Any packages in channelB would be sorted below those in channelA.

- 1. Sorts packages from highest to lowest channel priority.
- 2. Sorts tied packages—packages with the same channel priority—from highest to lowest version number. For example, if channelA contains NumPy 1.12.0 and 1.13.1, NumPy 1.13.1 will be sorted higher.
- 3. Sorts still-tied packages, packages with the same channel priority and same version, from highest to lowest build number. For example, if channelA contains both NumPy 1.12.0 build 1 and build 2, build 2 is sorted first. Any packages in channelB would be sorted below those in channelA.

4. Installs the first package on the sorted list that satisfies the installation specifications.

If a Python package that you need isn't available on any Conda channel, then you can use the default Python package manager Pip to install this package from PyPI.



If a Python package that you need isn't available on any Conda channel, then you can use the default Python package manager Pip to install this package from PyPI.

However, there are a few potential issues that you should be aware of when using Pip to install Python packages when using Conda.



Pip is sometimes installed by default on operating systems where it is used to manage any Python packages needed by your OS.

You do not want to use this pip to install Python packages when using Conda environments.

Need to deactivate first

```
(base) $ conda deactivate*
$ which python
/usr/bin/python
$ which pip # sometimes installed as pip3
/usr/bin/pip
```



Second, Pip is also included in the Miniconda installer where it is used to install and manage OS specific Python packages required to setup your base Conda environment.

You do not want to use this pip to install Python packages when using Conda environments.

```
$ conda activate
(base) $ which python
~/miniconda3/bin/python
$ which pip
~/miniconda3/bin/pip
```

If you find yourself needing to install a Python package that is only available via Pip, then you should first install pip into your Conda environment and then use that pip to install the desired package.

python -m pip install <package_name>

Note: Prefer python -m pip install over pip install; never use pip with the --user argument.

Exercise: Installing packages into Conda environments using pip

pandas is a fast, powerful, flexible and easy to use open source data analysis and manipulation too

- Activate the conda environment python3-env
- Use python -m pip install to install the python package pandas.
- Use conda list to see which Channel pandas was installed from.

Using Packages and Channels: Key Points

• A package is a tarball containing system-level libraries, Python or other modules, executable programs and other components, and associated metadata.

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- A package is a tarball containing system-level libraries, Python or other modules, executable programs and other components, and associated metadata.
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- A package is a tarball containing system-level libraries, Python or other modules, executable programs and other components, and associated metadata.
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- You can specific a conda channel using the option --channel or add it to your .condarc

Using Packages and Channels: Key Points

- A package is a tarball containing system-level libraries, Python or other modules, executable programs and other components, and associated metadata.
- A Conda channel is a URL to a directory containing a Conda package(s).
- You can specific a conda channel using the option --channel or add it to your .condarc
- If a python package isn't available on a conda channel you can install it into your environment using the python package installer pip.



Why should I share my Conda environment with others?

- Why should I share my Conda environment with others?
- How do I share my Conda environment with others?

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- How do I create an environment file that can be read by Windows, Mac OS, or Linux.
- How do I specifying the package version in a Conda environment file.

Conda environments are useful when making bioinformatics projects reproducible.

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Full reproducibility requires the ability to recreate the system that was originally used to generate the results.

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This can, to a large extent, be accomplished by using a Conda environment file to make an environment with specific versions of the packages that are needed in the project.

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Full reproducibility requires the ability to recreate the system that was originally used to generate the results.

This can, to a large extent, be accomplished by using a Conda environment file to make an environment with specific versions of the packages that are needed in the project.

This environment file can then be shared with others users to reproduce your analysis environment containing software with the same version number.

Conda uses YAML ("YAML Ain't Markup Language") for writing its environment files.

YAML is a human-readable language that is commonly used for configuration files and that that uses Python-style indentation to indicate nesting.

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name: rnaseq-env

a key-value pair **Key** name **Value** rnaseq-env

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YAML is a human-readable language that is commonly used for configuration files and that that uses Python-style indentation to indicate nesting.

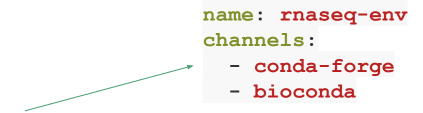
```
name: rnaseq-env
channels:
- conda-forge
- bioconda
```

The second item is an array channels with 2 elements, each denoted by an opening dash -.

https://edcarp.github.io/introduction-to-conda-for-data-scientists/04-sharing-environments/index.html

Conda uses YAML ("YAML Ain't Markup Language") for writing its environment files.

YAML is a human-readable language that is commonly used for configuration files and that that uses Python-style indentation to indicate nesting.



I indented the elements in channels with two spaces. Indentation is how YAML denotes nesting.

The number of spaces can vary from file to file, but tabs are not allowed

https://edcarp.github.io/introduction-to-conda-for-data-scientists/04-sharing-environments/index.html

Creating your project's Conda environment from a single environment file is a Conda "best practice".

Not only do you have a file to share with collaborators but you also have a file that can be placed under version control (Git) to further enhancing the reproducibility of your research project and workflow.

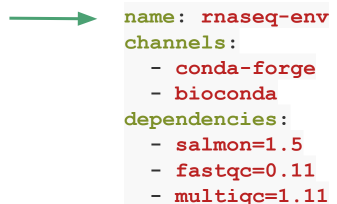
```
name: rnaseq-env
channels:
    - conda-forge
    - bioconda
dependencies:
    - salmon=1.5
```

- fastqc=0.11
- multiqc=1.11

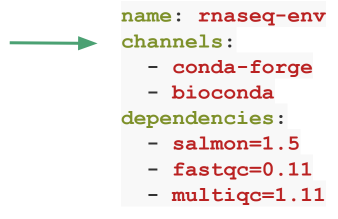
The first line specifies a default name rnaseq-env for the environment

```
name: rnaseq-env
channels:
- conda-forge
- bioconda
dependencies:
- salmon=1.5
- fastqc=0.11
- multiqc=1.11
```

The environment would be installed inside the conda environment directory e.g. ~/miniconda3/envs/ directory, unless we specified a different path using conda create command line option --prefix or -p.



The second line specifies a list of channels, listed in priority order, that packages may need to be installed from.



The dependencies lists the most current and mutually compatible versions of the listed packages (including all required dependencies) to download.

name: rnaseq-env
channels:
 - conda-forge
 - bioconda
dependencies:
 - salmon=1.5
 - fastqc=0.11
 - multiqc=1.11

Explicit versions numbers for all packages should be preferred a better environment file would be the following.

```
name: rnaseq-env
channels:
   - conda-forge
   - bioconda
dependencies:
   - salmon=1.5
   - fastqc=0.11
   - multiqc=1.11
```

Note that we are only specifying the major and minor version numbers and not the patch or build numbers.

4.2.1
MAJOR Minor patch

```
name: rnaseq-env
channels:
```

- conda-forge
- bioconda

dependencies:

- salmon=1.5
- fastqc=0.11
- multiqc=1.11

Defining the version number by fixing only the major and minor version numbers while allowing the patch version number to vary allows us to use our environment file to update our environment to get any bug fixes whilst still maintaining significant consistency of our Conda environment across updates.

```
$ cd ~/
$ mkdir rnaseq-project-2
$ cd rnaseq-project-2
```

Once your project folder is created, create an environment.yml file using your favourite editor for instance nano.

```
name: rnaseq-env
channels:
  - conda-forge
  - bioconda
dependencies:
  - salmon=1.5
  - fastqc=0.11
  - multiqc=1.11
```

Sharing Environments: Create a new conda environment:

Create a new conda environment: conda env create

```
$ conda env create --prefix ./env --file environment.yml
```

\$ conda activate ./env

Sharing Environments: Create a new conda environment:

We can automatically generate the contents of an environment file using the conda env export command.

```
conda env export --name basic-rnaseq-env
```

Sharing Environments: Export a conda environment:

To export this list into an **environment.yml** file, you can use **--file** option to directly save the resulting YAML environment into a file.

```
$ conda env export --name basic-rnaseq-env --file environment.yml
```

Make sure you do not have any other environment.yml file from before in the same directory when running the above command.

Sharing Environments: Export a conda environment: history

This exported environment file will however not *consistently* produce environments that are reproducible across Mac OS, Windows, and Linux. The reason is, that it may include operating system specific low-level packages which cannot be used by other operating systems.

If you need an environment file that can produce environments that are reproducible across Mac OS, Windows, and Linux, then you are better off just including those packages into the environment file that your have specifically installed using the --from-history option.

\$ conda env export --name basic-rnaseq-env --from-history --file environment.yml

Sharing Environments: PExport a conda environment: history: Pip

Note: Python packages installed via pip are not exported using conda env export --from-history.

Instead add them manually to environment file under the keyword **pip**:

```
name: rnaseq-project3-env
channels:
conda-forge
- bioconda
dependencies:
- salmon=1.5
- fastqc=0.11
- multiqc=1.11
pip:

    pandas=1.4
```

Exercise: Create a new environment from a YAML file.

Create a new project directory rnaseq-project-3 and then create a new environment.yml file inside your project directory with the following contents.

```
name: rnaseq-project3-env

channels:

- conda-forge

- bioconda

dependencies:

- salmon=1.5

- fastqc=0.11

- multiqc=1.11
```

- Now use this file to create a new Conda environment.
- Where is this new environment created?

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- one of your core dependencies just released a new version (dependency version number update).
- you need an additional package for data analysis (add a new dependency).
- you have found a better visualization package and no longer need to old visualization package (add new dependency and remove old dependency).

Sharing Environments: Updating an environment:

You are unlikely to know ahead of time which packages (and version numbers!) you will need to use for your research project. For example it may be the case that

- one of your core dependencies just released a new version (dependency version number update).
- you need an additional package for data analysis (add a new dependency).
- you have found a better visualization package and no longer need to old visualization package (add new dependency and remove old dependency).

Sharing Environments: Updating an environment

If any of these occurs during the course of your research project, all you need to do is update the contents of your environment.yml file accordingly and then run the following command.

```
$ cd ~/
$ cd rnaseq-project-2
```

Change salmon version to 1.6 in environment.yml

```
$ conda env update --prefix ./env --file environment.yml
```

Note: this doesn't remove packages that you remove from environment.yml

Exercise: Update environment from environment.yml

- Update the environment file environment.yml from the previous exercise, rnaseq-project-3,
 - a. by adding the package kallisto=0.46 and
 - b. removing the salmon package.
- 2. Then rebuild the environment conda env update

Is the salmon package still present?

Sharing Environments: Rebuilding a Conda environment

When working with environment.yml files it is often just as easy to rebuild the Conda environment from scratch whenever you need to add or remove dependencies.

Sharing Environments: Rebuilding a Conda environment

When working with environment.yml files it is often just as easy to rebuild the Conda environment from scratch whenever you need to add or remove dependencies.

To rebuild a Conda environment from scratch you can pass the --force option to the conda environment.

This will remove any existing environment directory before rebuilding it using the provided environment file.

```
$ cd ~/rnaseq-project-2
```

```
$ conda env create --prefix ./env --file environment.yml --force
```

Sharing Environments: Restoring an environment:

Conda keeps a history of all the changes made to your environment, so you can easily "roll back" to a previous version.

To list the history of each change to the current environment:

```
$ conda activate basic-rnaseq-env
```

```
$ conda list --revisions
```

Sharing Environments: Restoring an environment:

To restore environment to a previous revision:

```
$ conda install --revision=REVNUM or conda install --rev REVNUM.
```

\$ conda install --revision=1

Exercise: List revisions.

- 1. Activate the environment inside the rnaseq-project-3 and list the revisions
- 2. Install revision 0

Sharing Environments: Key Points

 Sharing Conda environments with other researchers facilitates the reproducibility of your research.

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• Sharing Conda environments with other researchers facilitates the reproducibility of your research.

 Conda environment files ,environment.yml, describes your project's software environment.

Configuring Conda

Configuring Conda: Questions

How can I configure conda?

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- How can I configure conda?
- How can I see conda's configuration values?

Configuring Conda: Questions

- How can I configure conda?
- How can I see conda's configuration values?
- How can I modify conda's configuration settings?

Configuration

Conda has a number of configuration setting which control how it performs.

To display and control these setting we can use the conda config subcommand.

To display all configuration settings run the config --show subcommand:

\$ conda config --show

Configuration

As you can see conda supports a large number of configuration options.

To show a single setting add the setting name after the conda config --show command.

\$ conda config --show channels

Configuration

By default conda only searches the defaults channel, this is why we had to include conda-forge and bioconda channels via the command line option --channel in the previous episode.

To get more information about an individual conda setting and its' possible values run conda config --describe <option>.

\$ conda config --describe channels

A user's conda setting are store in the runtime configuration configuration file, .condarc. This file allows users to configure various aspects of conda including:

Where conda looks for packages channels.

- Where conda looks for packages channels.
- Where conda lists known environments envs_dirs.

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- Whether to update the Bash prompt with the currently activated environment name env prompt.

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- Whether to update the Bash prompt with the currently activated environment name env_prompt.
- What default packages or features to include in new environments create default packages.

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- Where conda lists known environments envs dirs.
- Whether to update the Bash prompt with the currently activated environment name env_prompt.
- What default packages or features to include in new environments create default packages.

Like the environment file the .condarc configuration file follows a simple YAML syntax

The .condarc file is not included by default, but it is automatically created in your home directory the first time you run the conda config command.

Configuration: Creating or modify .condarc

To create or modify a .condarc file, enter the conda config command and use the modifier options

- --add
- --set
- --append,
- --prepend
- --remove

followed by the configuration key and a value.

conda config <modifier> <KEY> <VALUE>

Configuration: config --add

Adding conda-forge

```
$ conda config --add channels conda-forge
```

Configuration: config --add

Adding conda-forge

```
$ conda config --add channels conda-forge
```

This would add the conda-forge channel to the top of the channel list.

```
$ conda config --show channels
```

Configuration: config --append

We can use the conda config modifier --append to add conda-forge to the end of the channel list,

\$ conda config --append channels conda-forge

Giving it the lowest priority.

\$ conda config --show channels

Configuration: config --prepend

To move a channel to the highest priority use the conda config --prepend modifier.

\$ conda config --prepend channels conda-forge

Giving it the highest priority.

\$ conda config --show channels

Note: It is generally best to have conda-forge as the highest priority channel as this will usually have the most up-to-date packages.

Exercise: Adding the channels bioconda and conda-forge to .condarc.

Use conda config --add, --append to add the bioconda and conda-forge channels to your .condarc file.

Give conda-forge the highest priority.

Configuration: Setting configuration settings -- describe

If our configuration setting has a single boolean or string value we can use conda config --set to set it.

For example, In a previous episode we set the command line prompt setting for conda using env_prompt.

\$ conda config --describe env_prompt

The env_prompt setting takes a value of either '{prefix}', '{name}', and '{default_env}'.

Configuration:: Setting configuration settings -- set

To set the <code>env_prompt</code> to the default value '({default_env})' we can run:.

```
$ conda config --set env_prompt '({default_env})'
```

Note: You need to deactivate then reactivate the environment for the changes in env prompt to take effect.

Configuration: Setting configuration settings -- set

To set the env prompt to the default value '({default env})' we can run:.

```
$ conda config --set env_prompt '({default_env})'
```

To change it back to the just the environment name, we can run.

```
$ conda config --set env_prompt '({name})'
```

Exercise: Set conda channel_priority

 Use the conda config --describe to investigate the setting channel_priority.

2. Set the **channel_priority** so that packages in lower priority channels are not considered if a package with the same name appears in a higher priority channel.

3. Why would you want to do change this setting?

Configuration: Editing the .condarc file manually

You can also use a text editor such as nano to directly edit the .condarc.

To show the location and contents of your . condarc file you can use

\$ conda config --show-sources

Note: If the .condarc file is in the root environment, it will override any in the home directory

Exercise: Locate and view the .condarc

- Locate your .condarc file.
- Using your favourite text editor look at the .condarc.

Configuration: Help

As with all conda commands you can use the --help option to get help.

For example, for a complete list of conda config commands run

```
$ conda config --help
```

https://conda.io/projects/conda/en/latest/commands/config.html

Configuring Conda: Key Points

 The .condarc is an optional configuration file that stores custom conda setting.

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• The .condarc is an optional configuration file that stores custom conda setting.

 You can use the conda config subcommand to add, set or remove configuration setting in the .condarc file.

You can also edit the contents of the .condarc file directly using a text editor.

https://edcarp.github.io/introduction-to-conda-for-data-scientists/05-configuration/index.html

Conda: Cheat Sheet



https://docs.conda.io/projects/conda/en/4.6.0/ downloads/52a95608c49671267e40c689e0bc00ca/conda-cheatsheet.pdf

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