

The NiPreps ecosystem for reproducible neuroimaging

Russ Poldrack
Stanford University



The standard design (anti-)pattern for fMRI preprocessing

- Pick a single software package
 - Usually based on considerations other than performance



The standard design (anti-)pattern for fMRI preprocessing

- Pick a software package
 - Usually based on considerations other than performance
- String together the tools from that package into a script to run the preprocessing workflow

```
#!/bin/bash

# Generate the subject list to make modifying this script
# to run just a subset of subjects easier.

for id in `seq -w 1 26` ; do
    subj="sub-$id"
    echo "====> Starting processing of $subj"
    echo
    cd $subj

    # If the brain mask doesn't exist, create it
    if [ ! -f anat/${subj}_T1w_brain_f02.nii.gz ] ; then
        echo "Skull-stripped brain not found, using bet with a fractional intensity thresh
# Note: This fractional intensity appears to work well for most of the subjects in
# Flanker dataset. You may want to change it if you modify this script for your own
bet2 anat/${subj}_T1w.nii.gz \
anat/${subj}_T1w_brain_f02.nii.gz -f 0.2
    fi

    # Copy the design files into the subject directory, and then
    # change "sub-08" to the current subject number
    cp ..design_run1.fsf .
    cp ..design_run2.fsf .

    # Note that we are using the | character to delimit the patterns
    # instead of the usual / character because there are / characters
    # in the pattern.
    sed -i '' "s|sub-08|${subj}|g" \
        design_run1.fsf
    sed -i '' "s|sub-08|${subj}|g" \
        design_run2.fsf

    # Now everything is set up to run feat
    echo "====> Starting feat for run 1"
    feat design_run1.fsf
    echo "====> Starting feat for run 2"
    feat design_run2.fsf
    echo

    # Go back to the directory containing all of the subjects, and repeat the loop
    cd ..
done
```

The standard design (anti-)pattern for fMRI preprocessing

- Pick a software package
 - Usually based on considerations other than performance
- String together the tools from that package into a script to run the preprocessing workflow
 - Generally written by a student or postdoc with little software engineering experience
- Hope it keeps working over time...
- Trust that it does the right thing...

```
#!/bin/bash

# Generate the subject list to make modifying this script
# to run just a subset of subjects easier.

for id in `seq -w 1 26` ; do
    subj="sub-$id"
    echo "====> Starting processing of $subj"
    echo
    cd $subj

    # If the brain mask doesn't exist, create it
    if [ ! -f anat/${subj}_T1w_brain_f02.nii.gz ] ; then
        echo "Skull-stripped brain not found, using bet with a fractional intensity thresh
# Note: This fractional intensity appears to work well for most of the subjects in
# Flanker dataset. You may want to change it if you modify this script for your own
bet2 anat/${subj}_T1w.nii.gz \
anat/${subj}_T1w_brain_f02.nii.gz -f 0.2
    fi

    # Copy the design files into the subject directory, and then
    # change "sub-08" to the current subject number
    cp ..//design_run1.fsf .
    cp ..//design_run2.fsf .

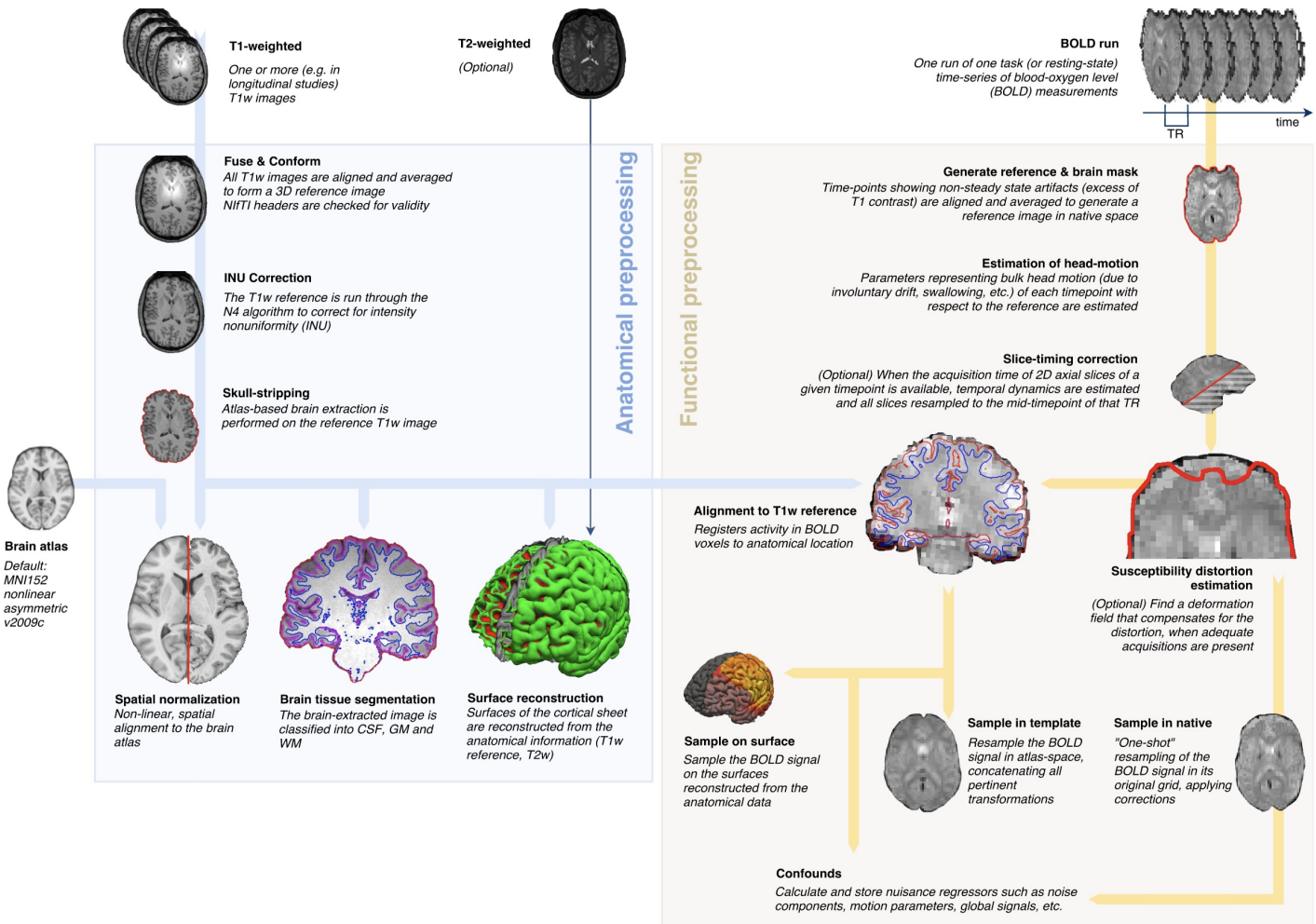
    # Note that we are using the | character to delimit the patterns
    # instead of the usual / character because there are / characters
    # in the pattern.
    sed -i '' "s|sub-08|${subj}|g" \
        design_run1.fsf
    sed -i '' "s|sub-08|${subj}|g" \
        design_run2.fsf

    # Now everything is set up to run feat
    echo "====> Starting feat for run 1"
    feat design_run1.fsf
    echo "====> Starting feat for run 2"
    feat design_run2.fsf
    echo

    # Go back to the directory containing all of the subjects, and repeat the loop
    cd ..
done
```

Towards a new design pattern for fMRI preprocessing: fMRIPrep

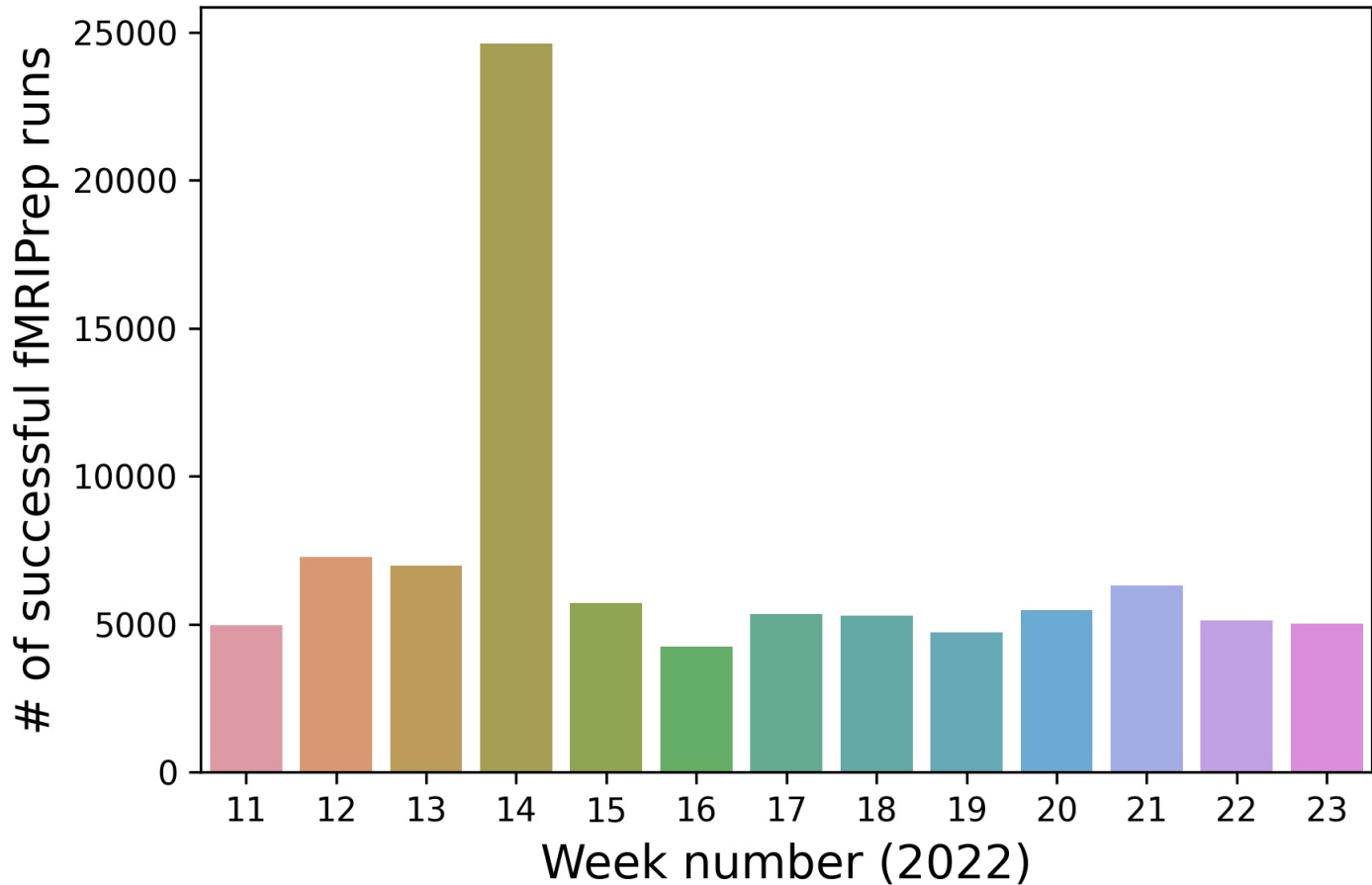
- A robust workflow for preprocessing fMRI data
 - Tested on a large number of fMRI datasets from OpenNeuro
 - Adapts to each dataset to process it in the best way possible
 - Provides a common interface to a variety of tools
 - Provide visualization to help identify problems



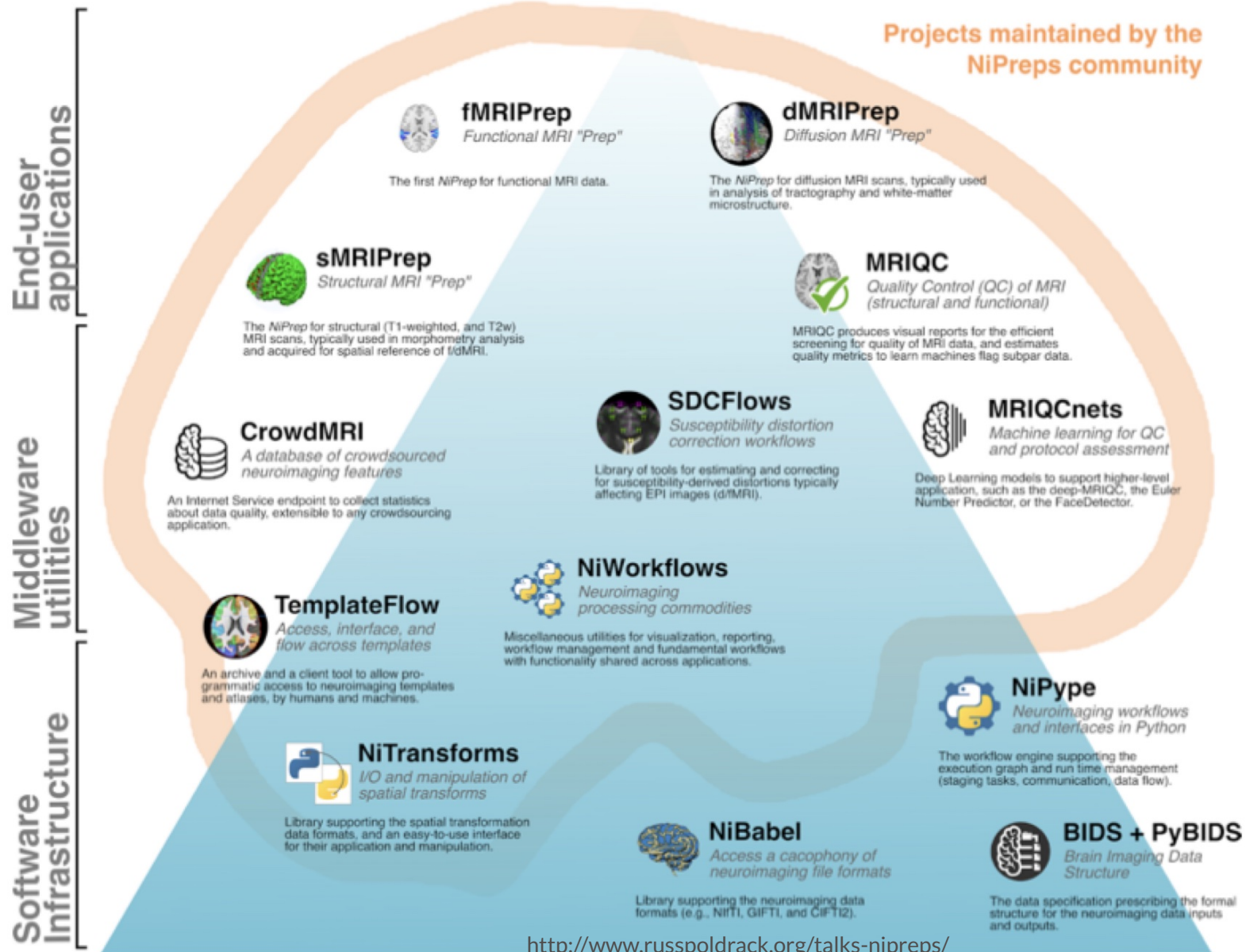
!

fMRIPrep usage

- Usage tracked using an opt-out telemetry system
 - Allows quick identification of bugs and usage patterns



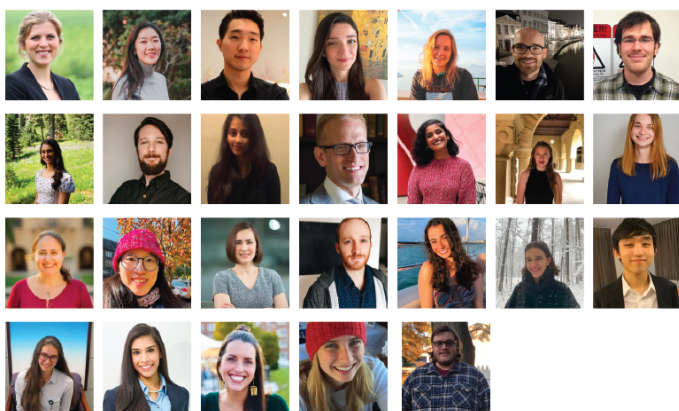
NiPreps: An ecosystem of Neuroimaging Preprocessing Tools



The NiPreps community



Poldracklab (Stanford)
Esteban Lab (CHUV Lausanne)



Satterthwaite Lab (Penn)
Milham Lab (Child Mind Inst)



Ariel Rokem
Research Associate Professor
email: arokem@uw.edu
[Website](#)



McKenzie Hagen
Graduate student, started 2021



Ezgi Yucel
Graduate student (with Ione Fine), started 2017



John Kruper
Post-bacc student, started July 2020



Teresa Gomez
Post-bacc student, started January 2022

Rokem Lab (UW)

Nipreps principles

- **Openness and community engagement**
 - All work done in the open
 - Community-driven decision making

Nipreps principles

- **Openness and community engagement**
 - All work done in the open
 - Community-driven decision making
- **Reproducibility and transparency**
 - Containerized applications following the BIDS-Apps specification
 - Detailed visual reporting
 - Citation boilerplate to ensure standards-compliant reporting

Nipreps principles

- **Openness and community engagement**
 - All work done in the open
 - Community-driven decision making
- **Reproducibility and transparency**
 - Containerized applications following the BIDS-Apps specification
 - Detailed visual reporting
 - Citation boilerplate to ensure standards-compliant reporting
- **Generality, consistency, and robustness**
 - Fully based on the BIDS and BIDS-Derivatives standards
 - Tested using continuous integration

Nipreps principles

- **Openness and community engagement**
 - All work done in the open
 - Community-driven decision making
- **Reproducibility and transparency**
 - Containerized applications following the BIDS-Apps specification
 - Detailed visual reporting
 - Citation boilerplate to ensure standards-compliant reporting
- **Generality, consistency, and robustness**
 - Fully based on the BIDS and BIDS-Derivatives standards
 - Tested using continuous integration
- **Modularity and longevity potential**
 - Reusable middleware components

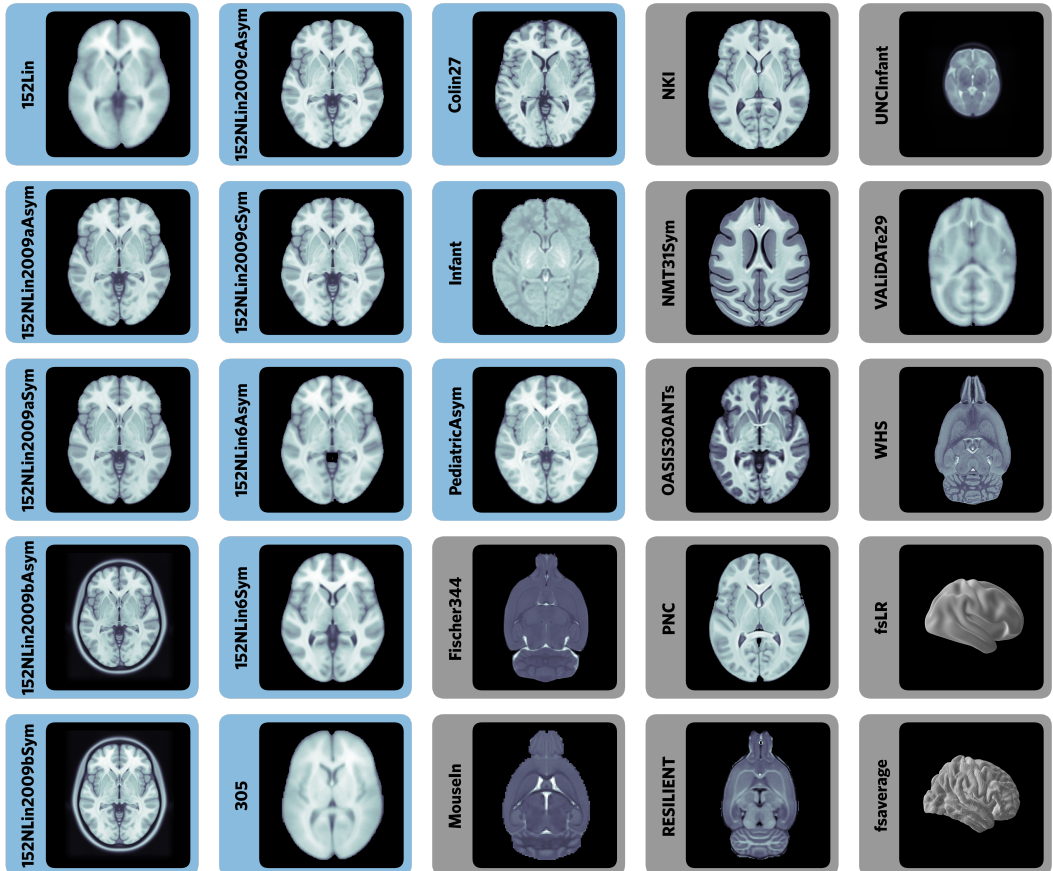
RF1MH121867: NiPreps: integrating neuroimaging preprocessing workflows across modalities, populations, and species

- **Aim 1: Solidify and extend NiPreps infrastructure and middleware**
 - TemplateFlow: FAIR Sharing and management of neuroimaging templates and atlases
 - SDCflows: Integrating susceptibility distortion correction
 - NiTransforms: Integrating spatial transforms

Middleware: TemplateFlow

- Templates and atlases are commonly used in neuroimaging research
- There is significant lack of clarity in the use of these templates
 - There are numerous versions of the widely used “MNI template”
- Templateflow provides programmatic access to a database of templates and mappings between them
- Python API client:

```
>>> from templateflow import api as tf
>>> tf.get("MNI152NLin2009cAsym",
...         suffix="T1w")
```



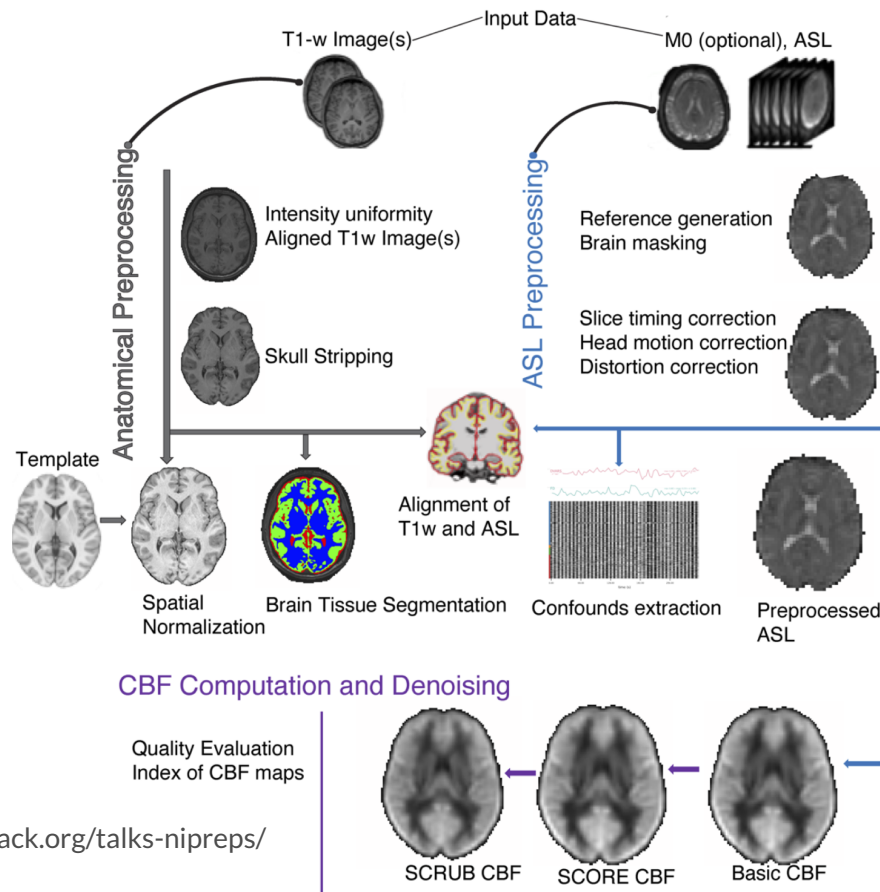
Middleware: SDCFlows

RF1MH121867: NiPreps: integrating neuroimaging preprocessing workflows across modalities, populations, and species

- Aim 2: Expand the portfolio of end-user NiPreps
 - ASLPrep
 - dMRIPrep
 - PETPrep
 - NiBabies
 - NiRodents

Workflows: ASLPrep (cerebral blood flow quantification)

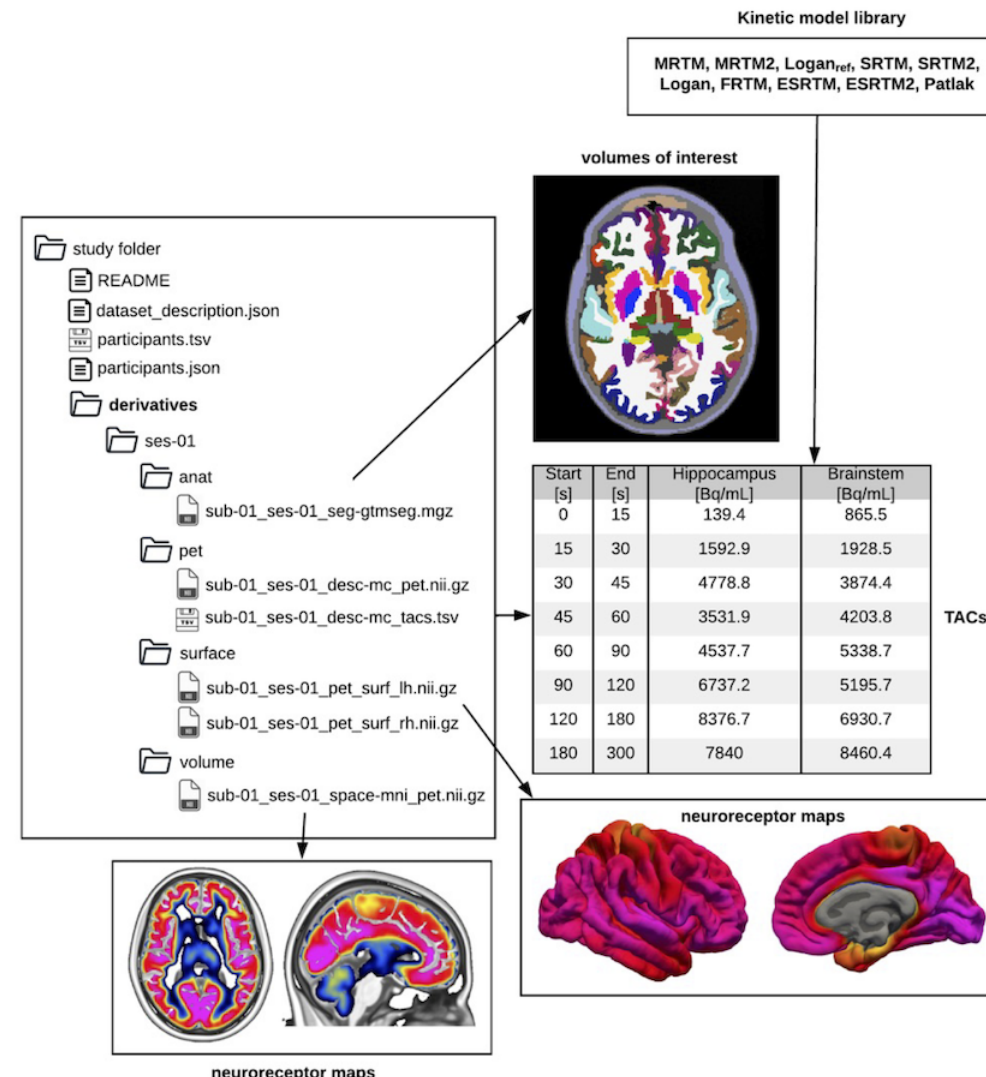
- A robust workflow for preprocessing arterial spin labeling (ASL) data
 - Including cerebral blood flow (CBF) quantification
 - Provides quality evaluation for CBF maps
 - Provides CBF quantification at the regional level using atlases



Workflows: dMRIPrep (diffusion MRI)

Workflows: PETPrep (positron emission tomography)

- A NiPreps workflow for PET preprocessing
 - Successfully merged petsurfer into nipype (1.8.0)
 - Incorporated nipype implementation of a robust head motion correction workflow (petprep_hmc)
 - Developing a BIDS-Derivatives standard for PET derivatives



Workflows: NiBabies (infant human MRI)



Workflows: NiRodents (rodent MRI)

RF1MH121867: NiPreps: integrating neuroimaging preprocessing workflows across modalities, populations, and species

- Aim 3: Consolidate the NiPreps community
 - Project monitoring infrastructure: MIGAS
 - Evaluation of cross-workflow reproducibility
 - Hackathons and documentation

Project monitoring: MIGAS

Reproducibility: Cross-workflow evaluation

Hackathons and documentation

Thank you!