

ARCify your research project

Hands-on ARC introduction @ IRTG 2843

Dominik Brilhaus, [CEPLAS](#)

October 29th, 2025



Interactive pad for the course

- Please use the pad to raise questions and feedback
- I'll provide links for hands-on there



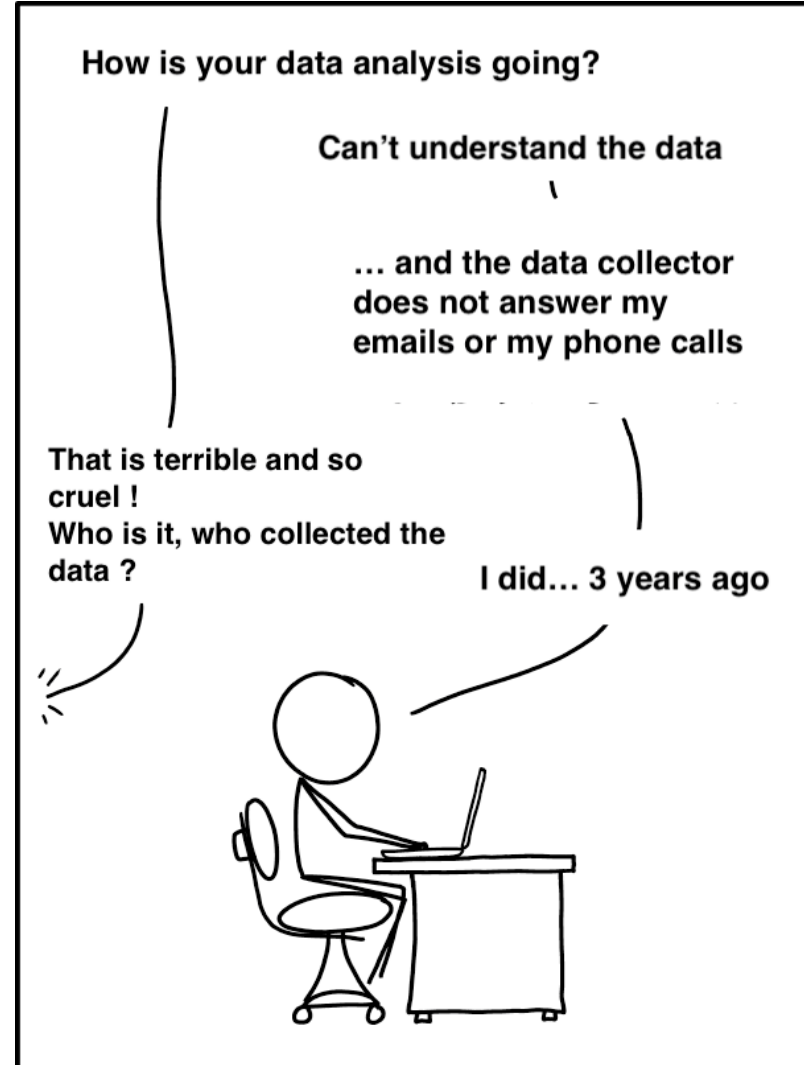
<https://t1p.de/4bp9i>

Introduce yourself

- My motivation / expectation
- Operating system: Linux / Windows / Mac?
- Used code / programming language before?
- Experience with Git / GitLab / GitHub?
- Have an ORCID
- My favorite experiment (lab / field / computational)

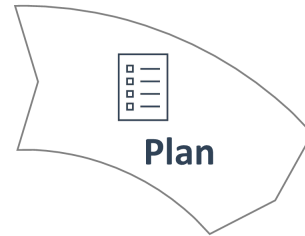
Why Research Data Management (RDM)?

- Increase transparency
- Make data accessible
- Save time (writing, reusing)
- Reduce risk of data loss
- Optimize costs
- Facilitate future reuse and sharing
- Improve citations

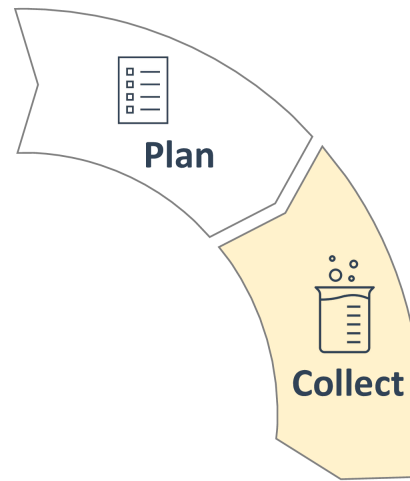


Your first collaborators
are your future selves,
be nice to them !

The Research Data Lifecycle



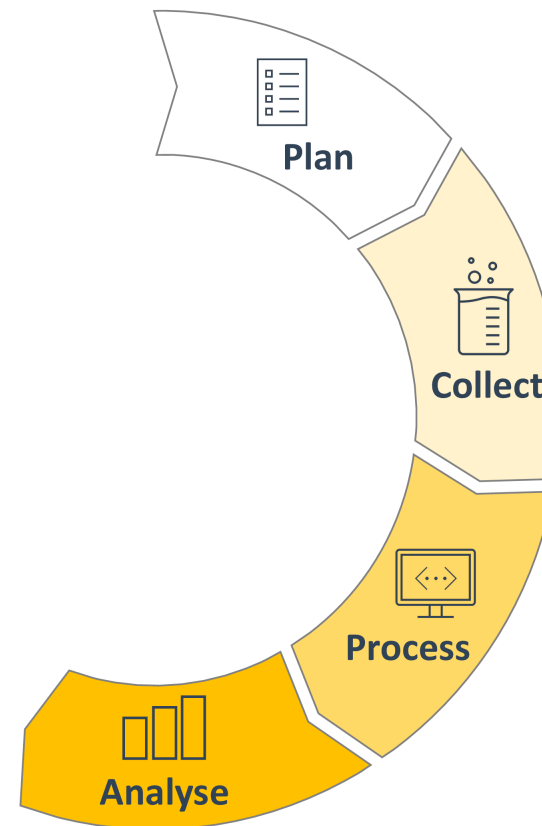
The Research Data Lifecycle



The Research Data Lifecycle



The Research Data Lifecycle



The Research Data Lifecycle



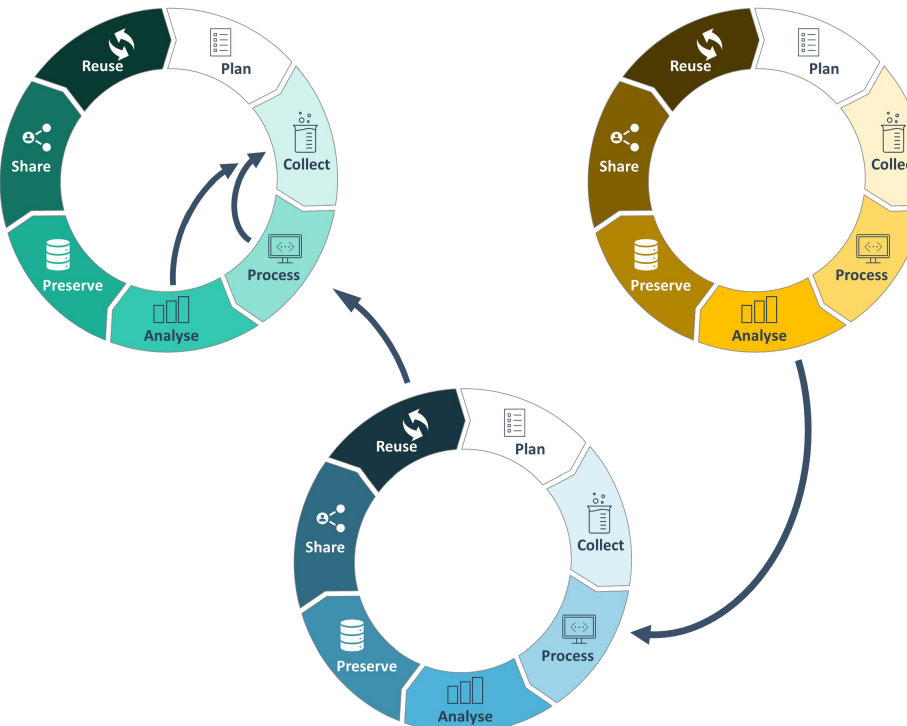
The Research Data Lifecycle



The Research Data Lifecycle



The Research Data Lifecycle *is mutable*



FAIR data stewardship

- Findable
- Accessible
- Interoperable
- Reusable

[nature](#) > [scientific data](#) > [comment](#) > [article](#)

[Open Access](#) | Published: 15 March 2016

The FAIR Guiding Principles for scientific data management and stewardship

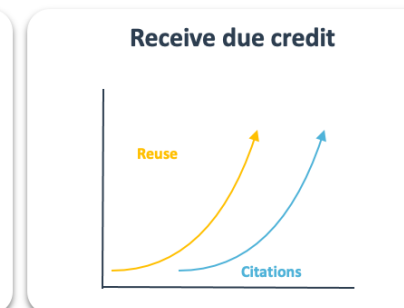
[Mark D. Wilkinson](#), [Michel Dumontier](#), [IJsbrand Jan Aalbersberg](#), [Gabrielle Appleton](#), [Myles Axton](#), [Arie Baak](#), [Niklas Blomberg](#), [Jan-Willem Boiten](#), [Luiz Bonino da Silva Santos](#), [Philip E. Bourne](#), [Jildau Bouwman](#), [Anthony J. Brookes](#), [Tim Clark](#), [Mercè Crosas](#), [Ingrid Dillo](#), [Olivier Dumon](#), [Scott Edmunds](#), [Chris T. Evelo](#), [Richard Finkers](#), [Alejandra Gonzalez-Beltran](#), [Alasdair J.G. Gray](#), [Paul Groth](#), [Carole Goble](#), [Jeffrey S. Grethe](#), [Jaap Heringa](#), [Peter A.C. 't Hoen](#), [Rob Hooft](#), [Tobias Kuhn](#), [Ruben Kok](#), [Joost Kok](#), [Scott J. Lusher](#), [Maryann E. Martone](#), [Albert Mons](#), [Abel L. Packer](#), [Bengt Persson](#), [Philippe Rocca-Serra](#), [Marco Roos](#), [Rene van Schaik](#), [Susanna-Assunta Sansone](#), [Erik Schultes](#), [Thierry Sengstag](#), [Ted Slater](#), [George Strawn](#), [Morris A. Swertz](#), [Mark Thompson](#), [Johan van der Lei](#), [Erik van Mulligen](#), [Jan Velterop](#), [Andra Waagmeester](#), [Peter Wittenburg](#), [Katherine Wolstencroft](#), [Jun Zhao](#) & [Barend Mons](#)✉

— Show fewer authors

[Scientific Data](#) 3, Article number: 160018 (2016) | [Cite this article](#)

<https://doi.org/10.1038/sdata.2016.18>

The FAIR principles



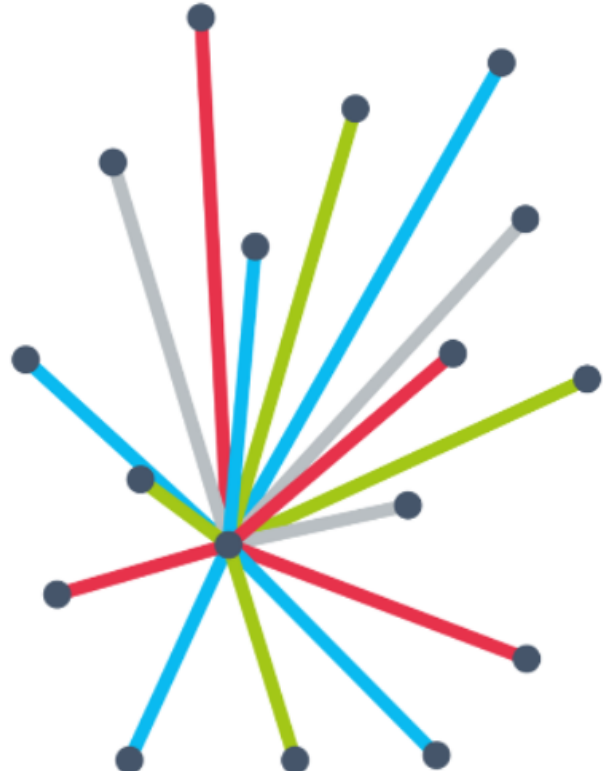
Is your data FAIR?

Findable | Accessible | Interoperable | Reusable

- Where do you store your data?
- How do you annotate your data?
- How do you share your data?
- What tools do you use to analyse your data?
- How do you reuse other people's data?



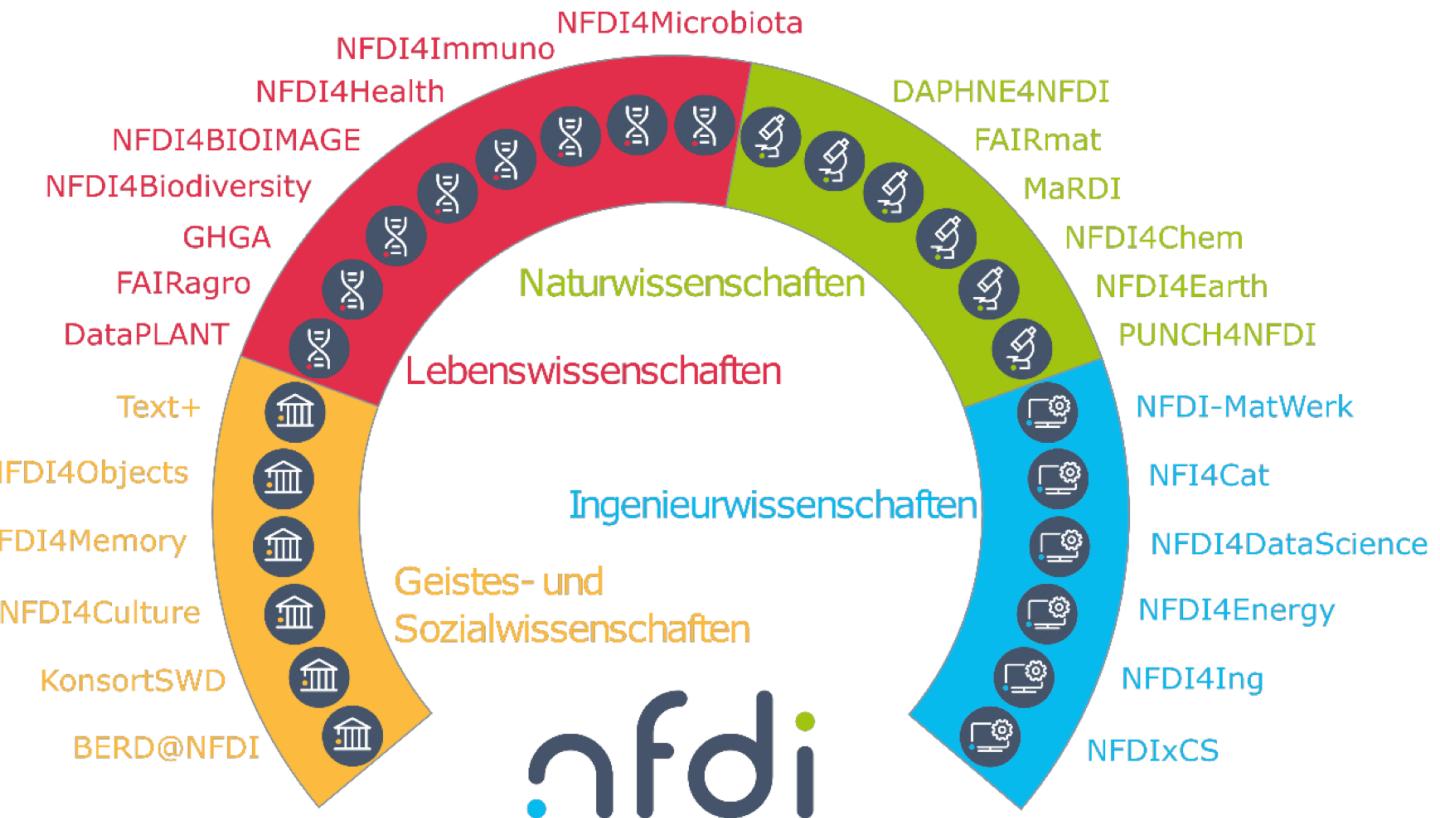
NFDI – Nationale Forschungsdaten Infrastruktur



The aim of the national research data infrastructure (NFDI) is to systematically manage scientific and research data, provide long-term data storage, backup and accessibility, and network the data both nationally and internationally.

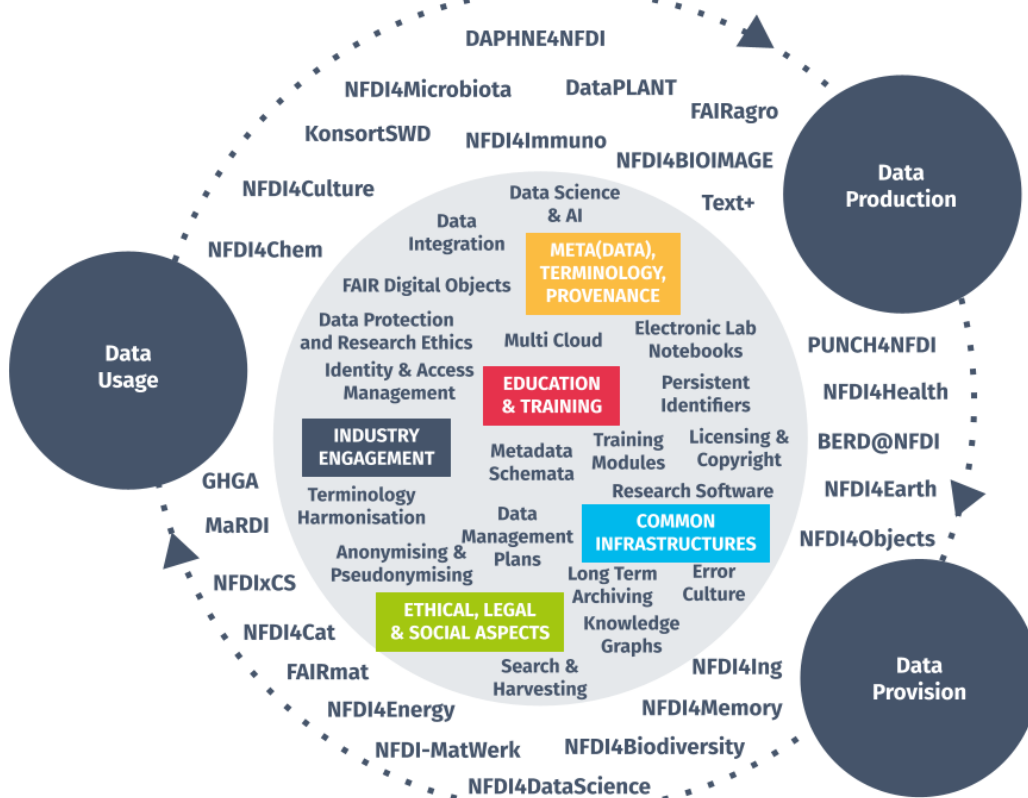
nfdi.de

NFDI – A community–targeted approach for RDM



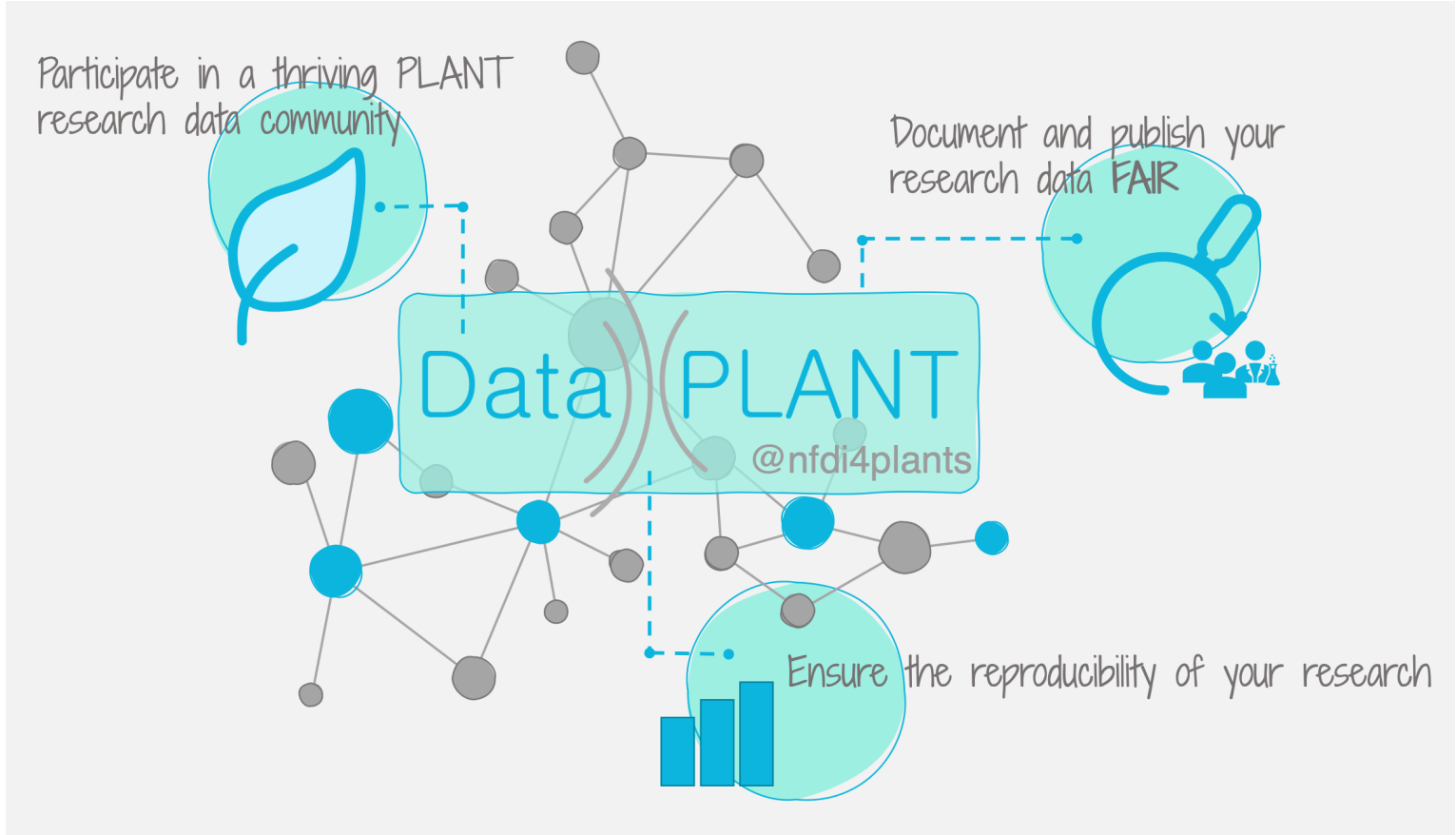
nfdi.de

NFDI – A community-targeted approach for RDM



<https://base4nfdi.de/>

DataPLANT – NFDI4plants



<https://nfdi4plants.org>

Data Stewardship between DataPLANT and the community

Community



CEPLAS

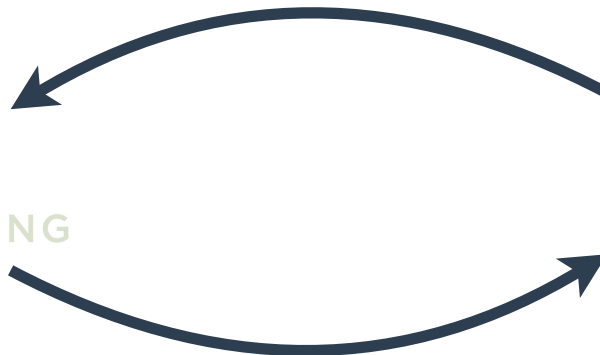


Domain experts
User experience
Training

nfdi4plants



Service provider
Developers
Tech experts



Resources



Info & materials

- DataPLANT Website: <https://nfdi4plants.org/>
- ARC website: <https://arc-rdm.org>
- Knowledge Base:
<https://nfdi4plants.org/nfdi4plants.knowledgebase/>

Tools and Services

- ARCitect: <https://github.com/nfdi4plants/arcitect>
- DataHUB: <https://git.nfdi4plants.org>

Continuous support

- HelpDesk: <https://helpdesk.nfdi4plants.org>
- Matrix for ad hoc support: <https://matrix.to/#/%23arc-user-support:matrix.org>
- User Support Meeting (2nd Friday of the month | 1 – 2pm):
<https://nfdi4plants.github.io/events/arc-user-support/>
- User Support Mailing List: [Click here to subscribe](#)

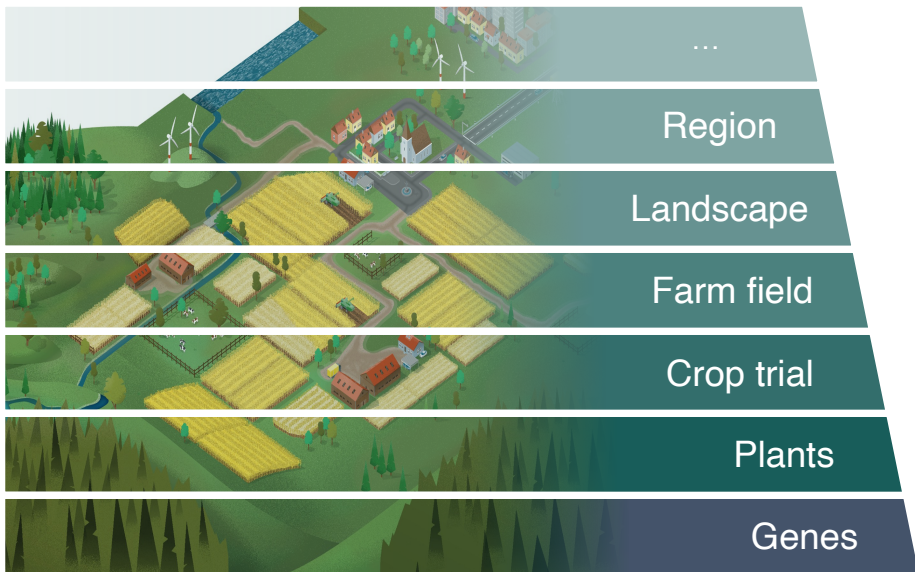
Open Source Development

- GitHub: <https://github.com/nfdi4plants>

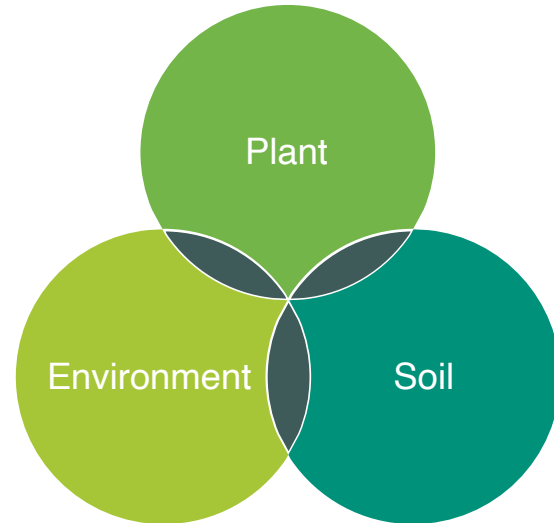
What is FAIRagro?!

FAIRagro is the NFDI consortium for the **agrosystem sciences**

Scales

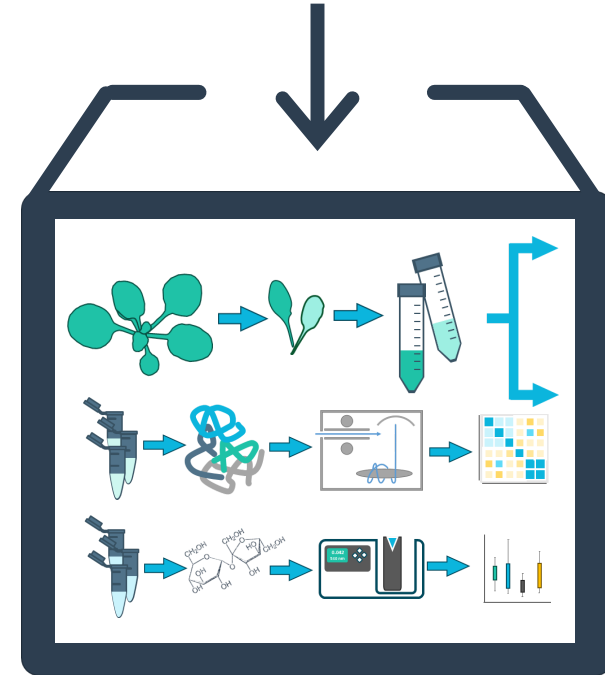
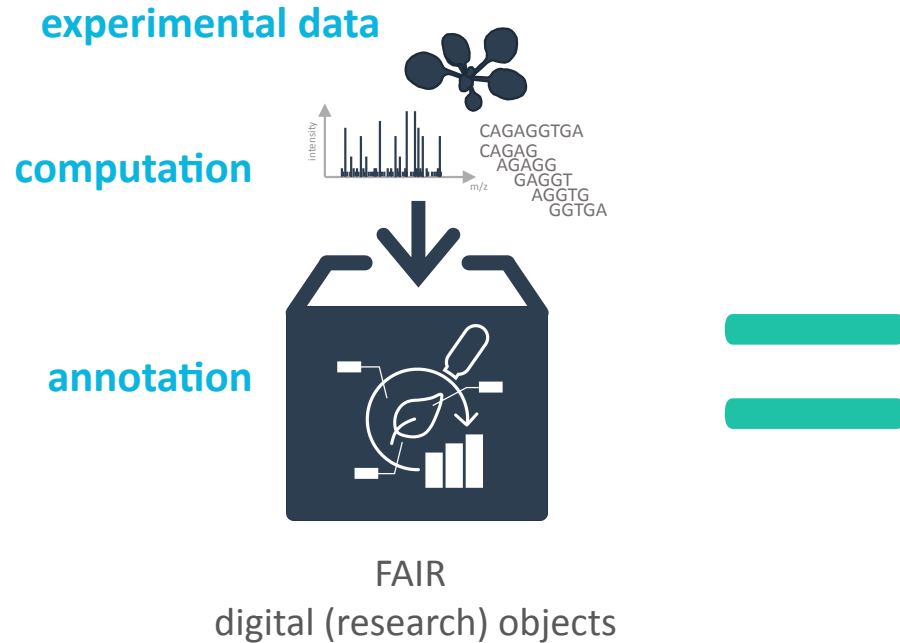


Subjects and disciplines

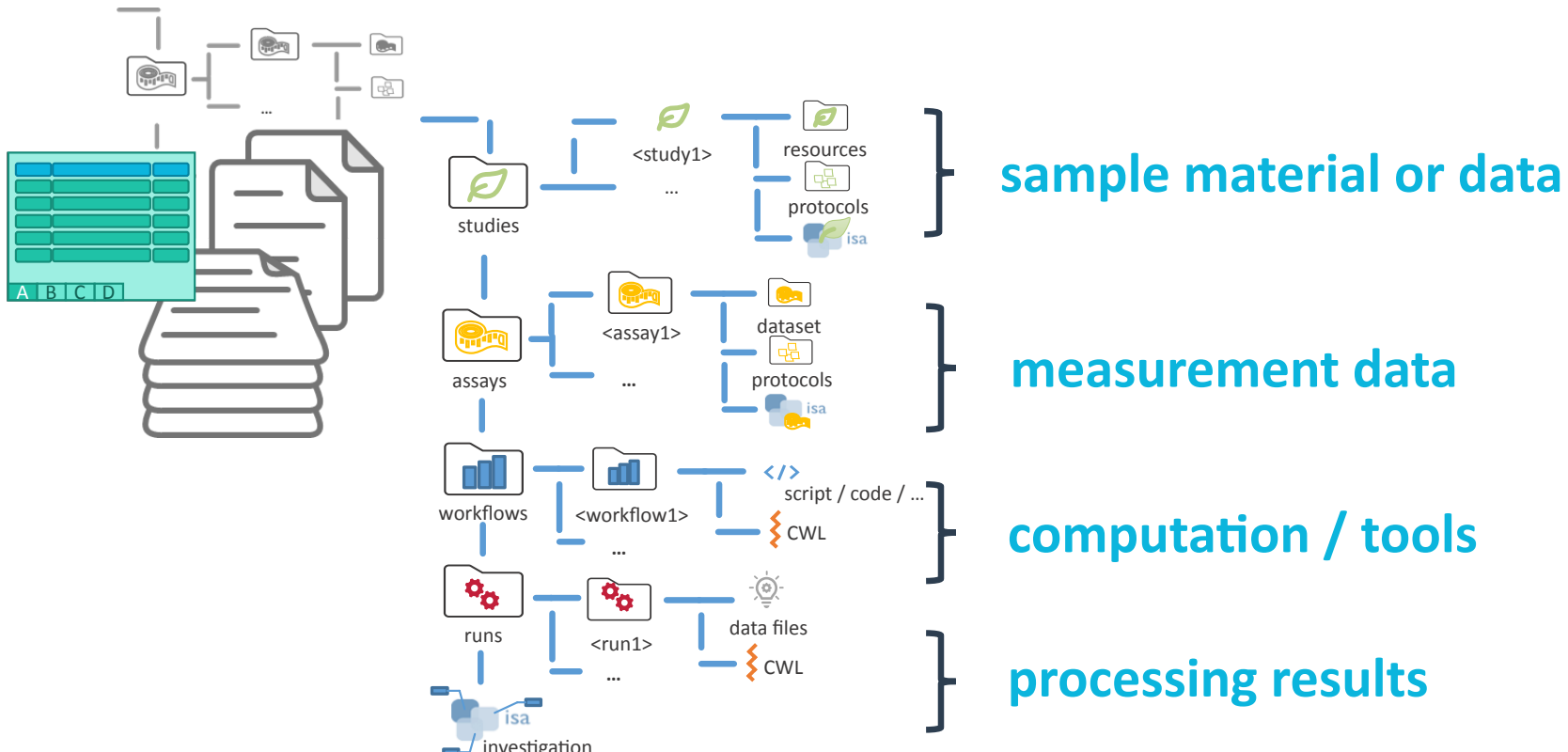


- Agronomy
- Crop physiology/ecology
- Plant pathology
- Plant nutrition
- Crop breeding
- Crop functional genetics
- Crop bioinformatics
- Soil biology/chem./physics
- Soil microbiology
- Soil hydrology
- Agroecology
- Agrobiodiversity
- Remote sensing
- Modelling
- ...

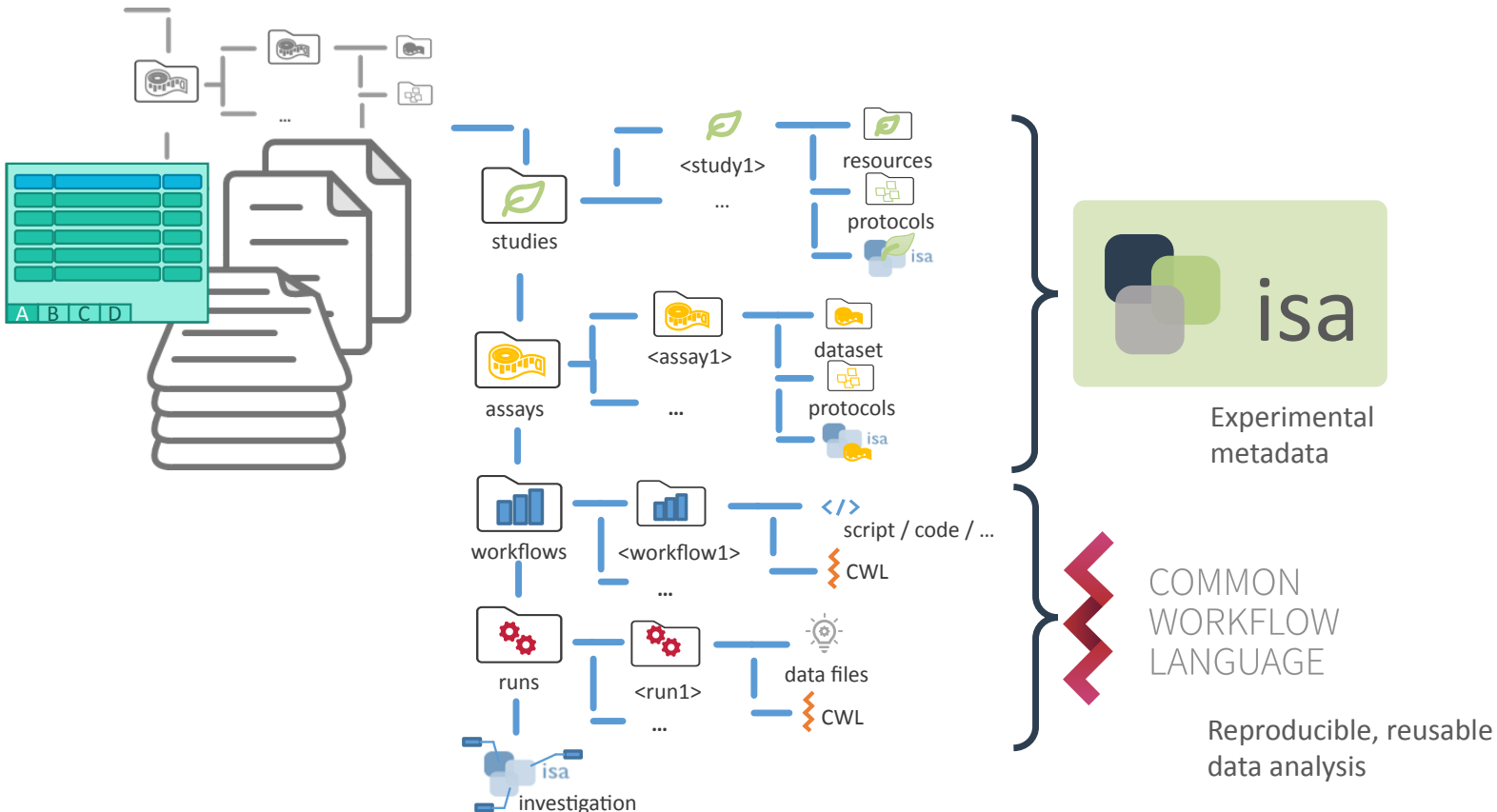
Annotated Research Context (ARC)



The ARC scaffold structure



ARC builds on standards



<https://isa-tools.org> | <https://www.commonwl.org> | <https://www.researchobject.org/ro-crate> | <https://git-scm.com>

ISA abstract model in a nutshell



isa

Investigation
administrative (meta)data

- Summary
 - Titel
 - Description
- Person
- Organisation
- Publication reference

Study
descriptive (meta)data
information on the subject

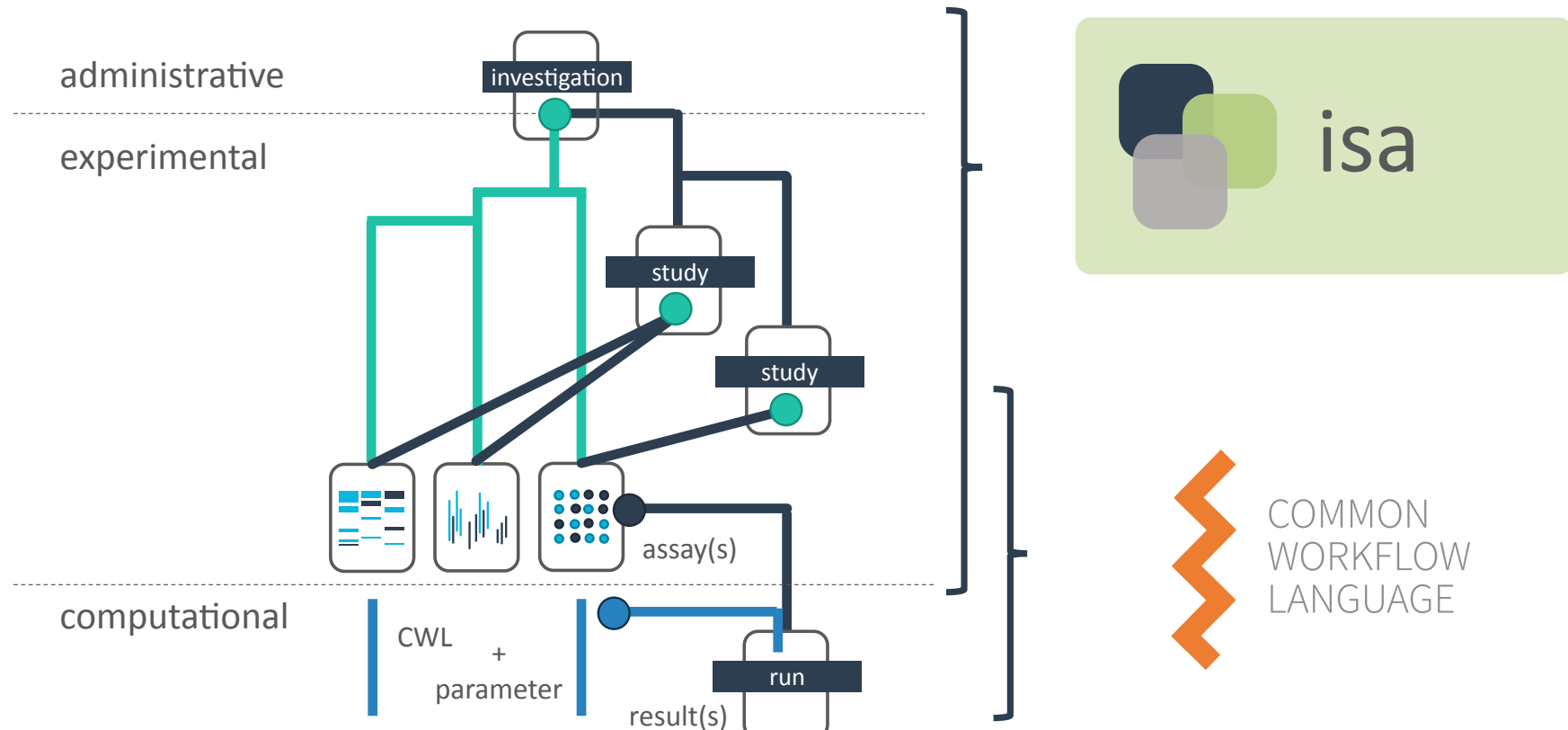
- Characteristics
- Parameters
- Components
- Factors

Assay

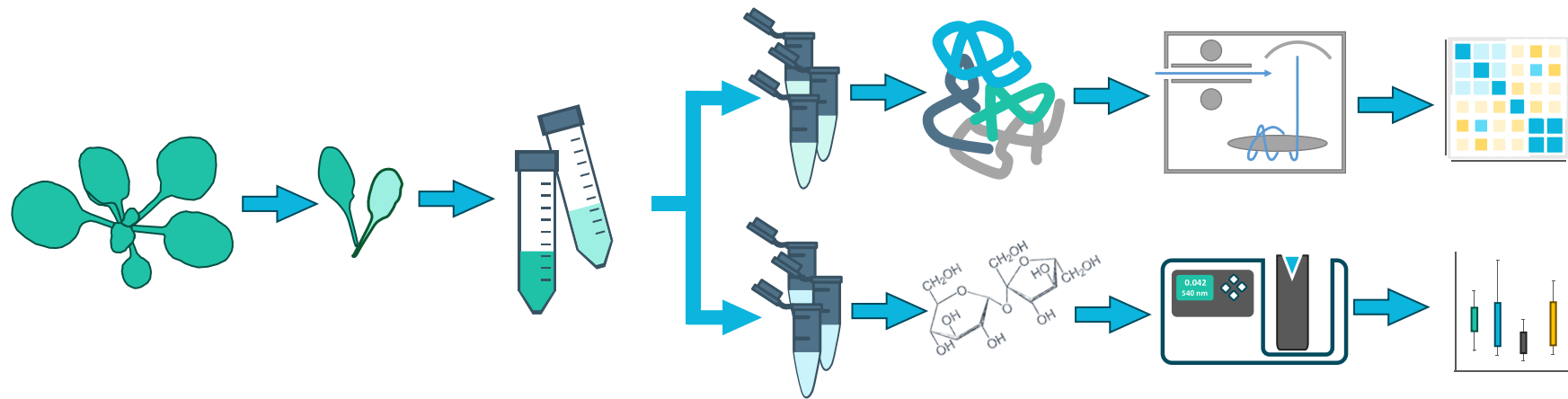
descriptive (meta)data
information on the measurement

- Characteristics
- Parameters
- Components
- Factors

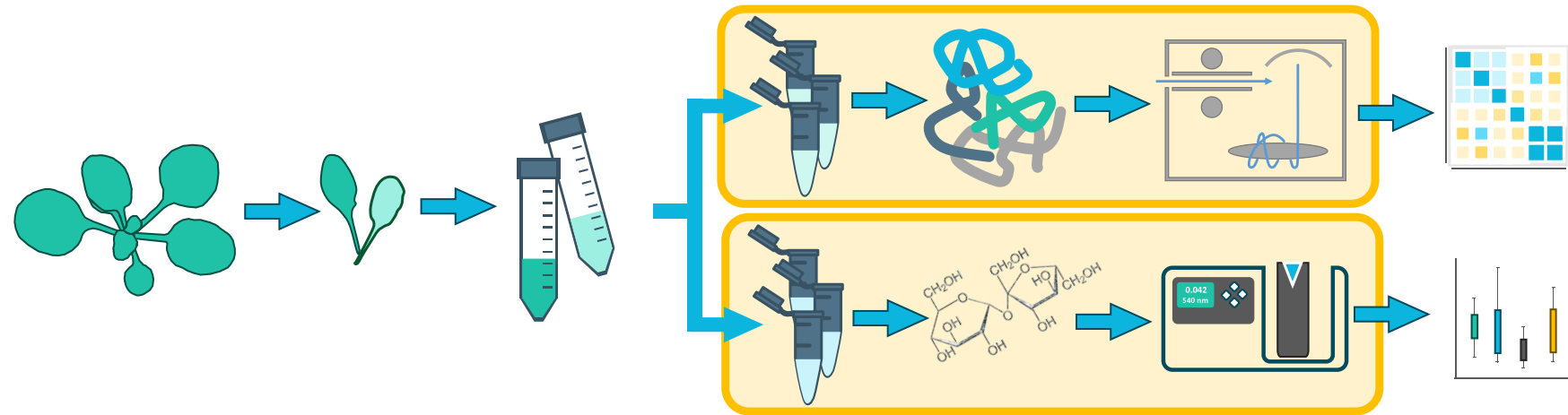
ISA and CWL – Connected by similarity



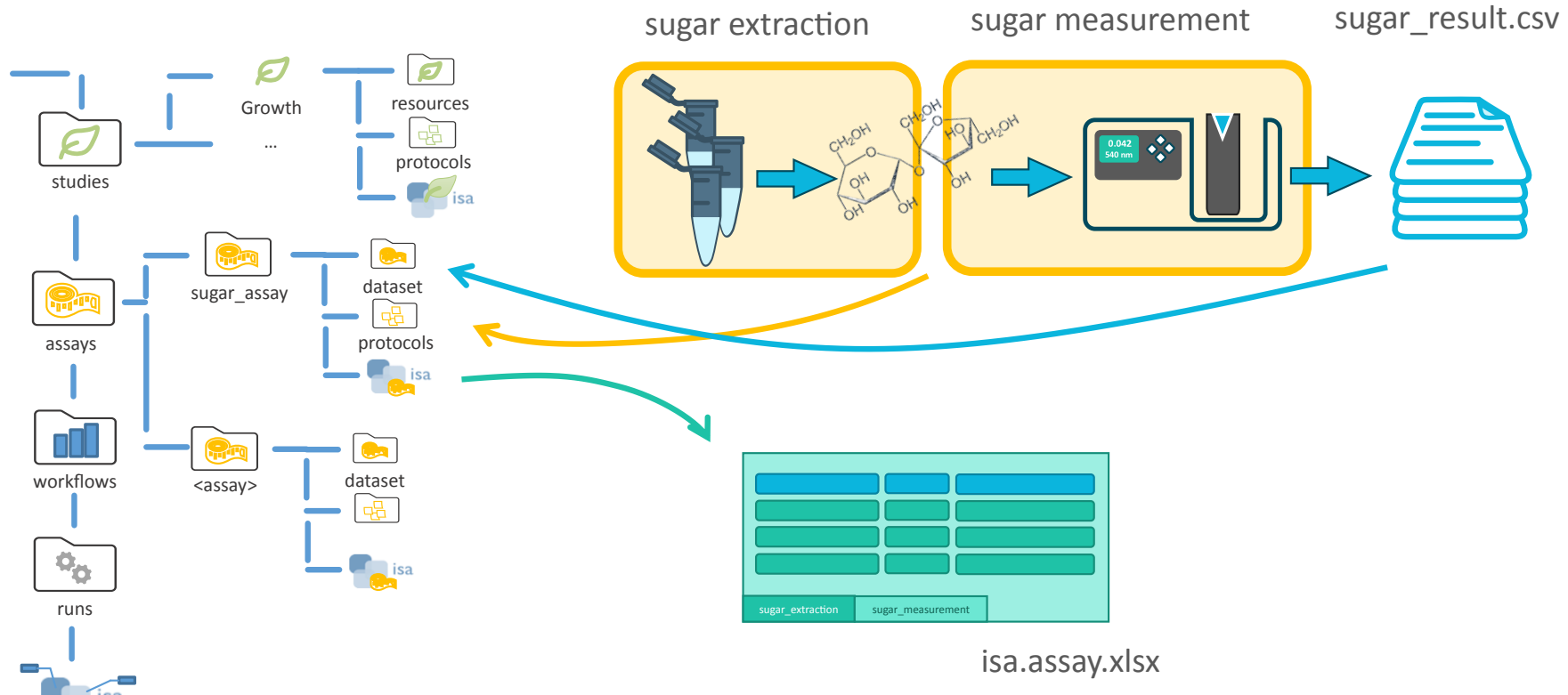
Metadata annotation – from sample to data



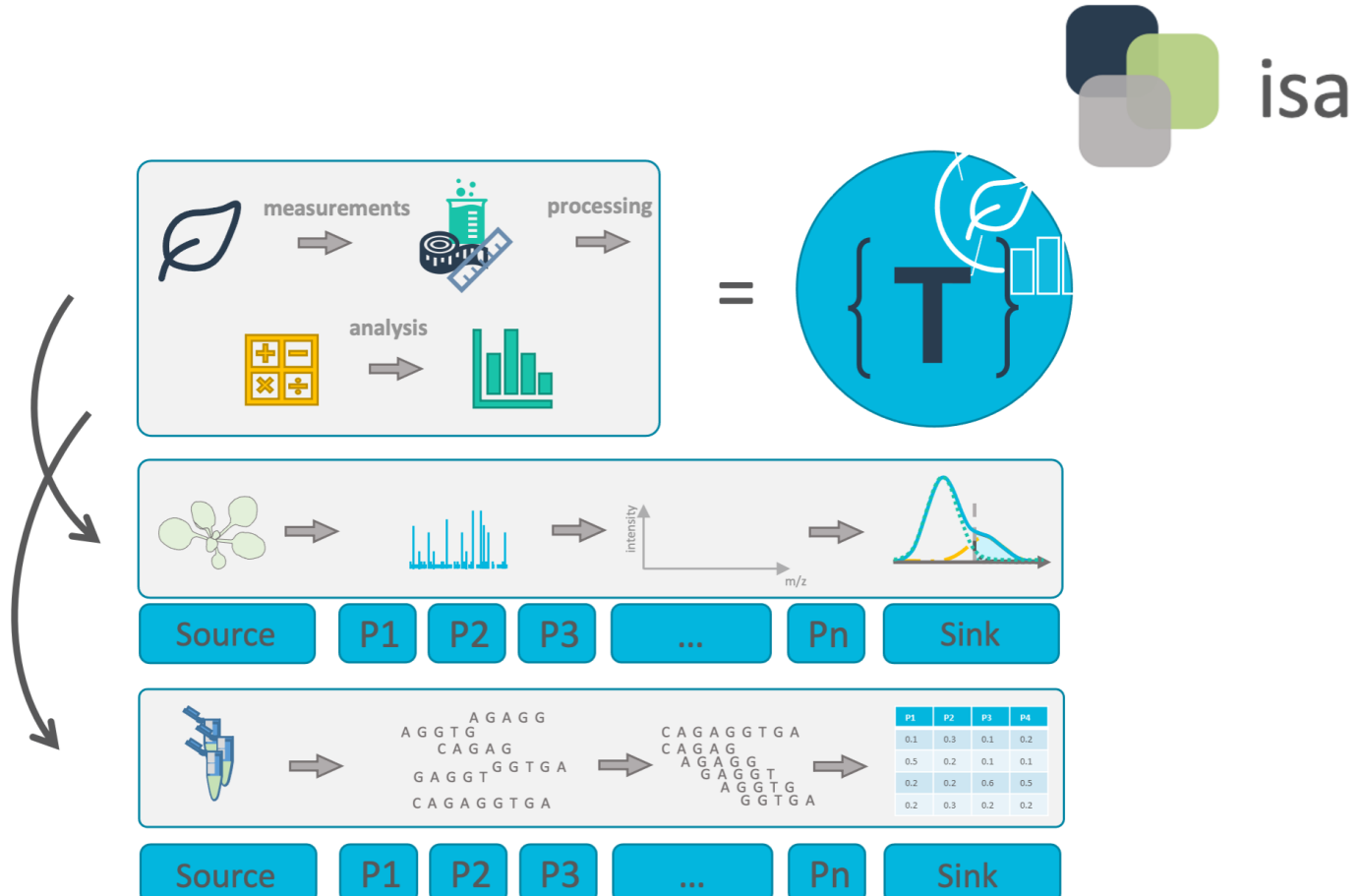
Modular separation of experimental processes



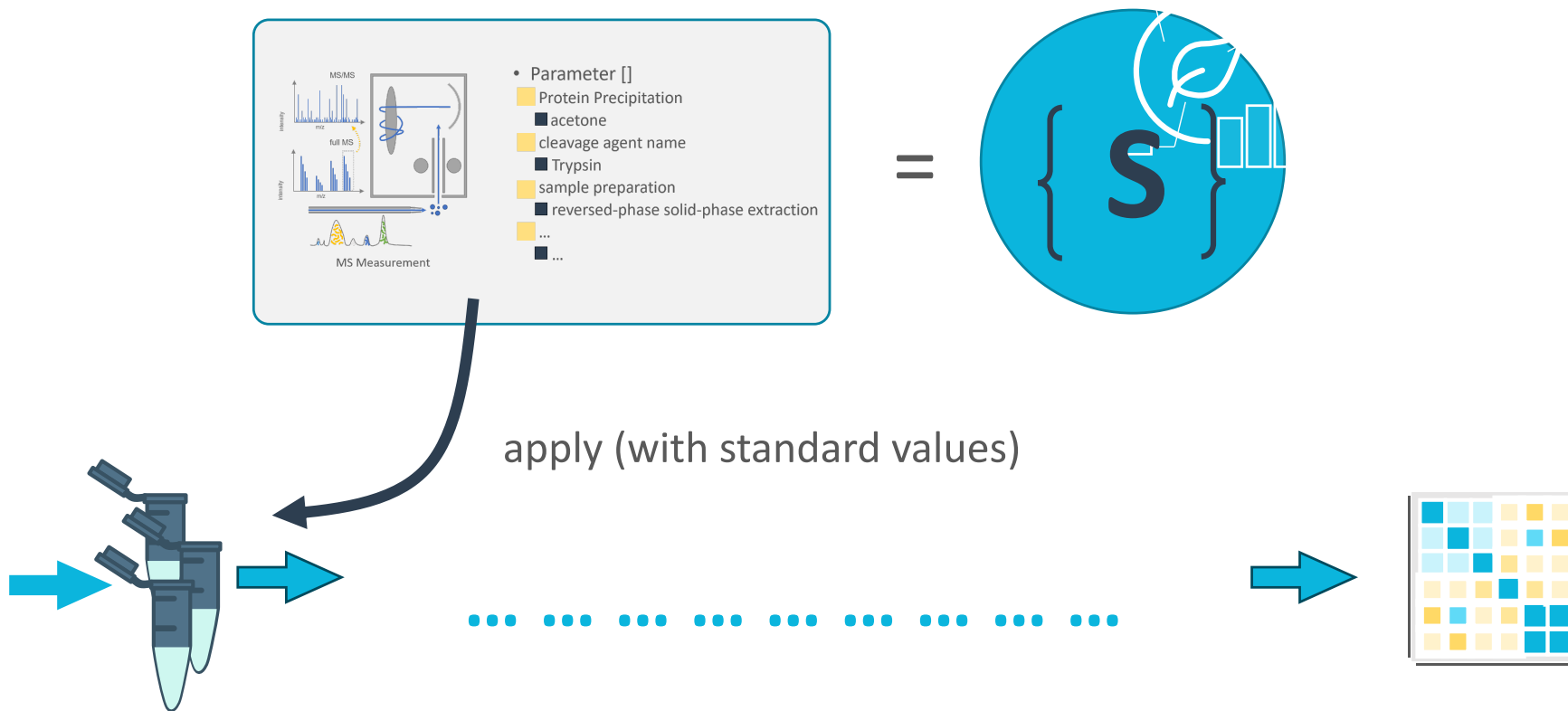
Modular separation of experimental processes



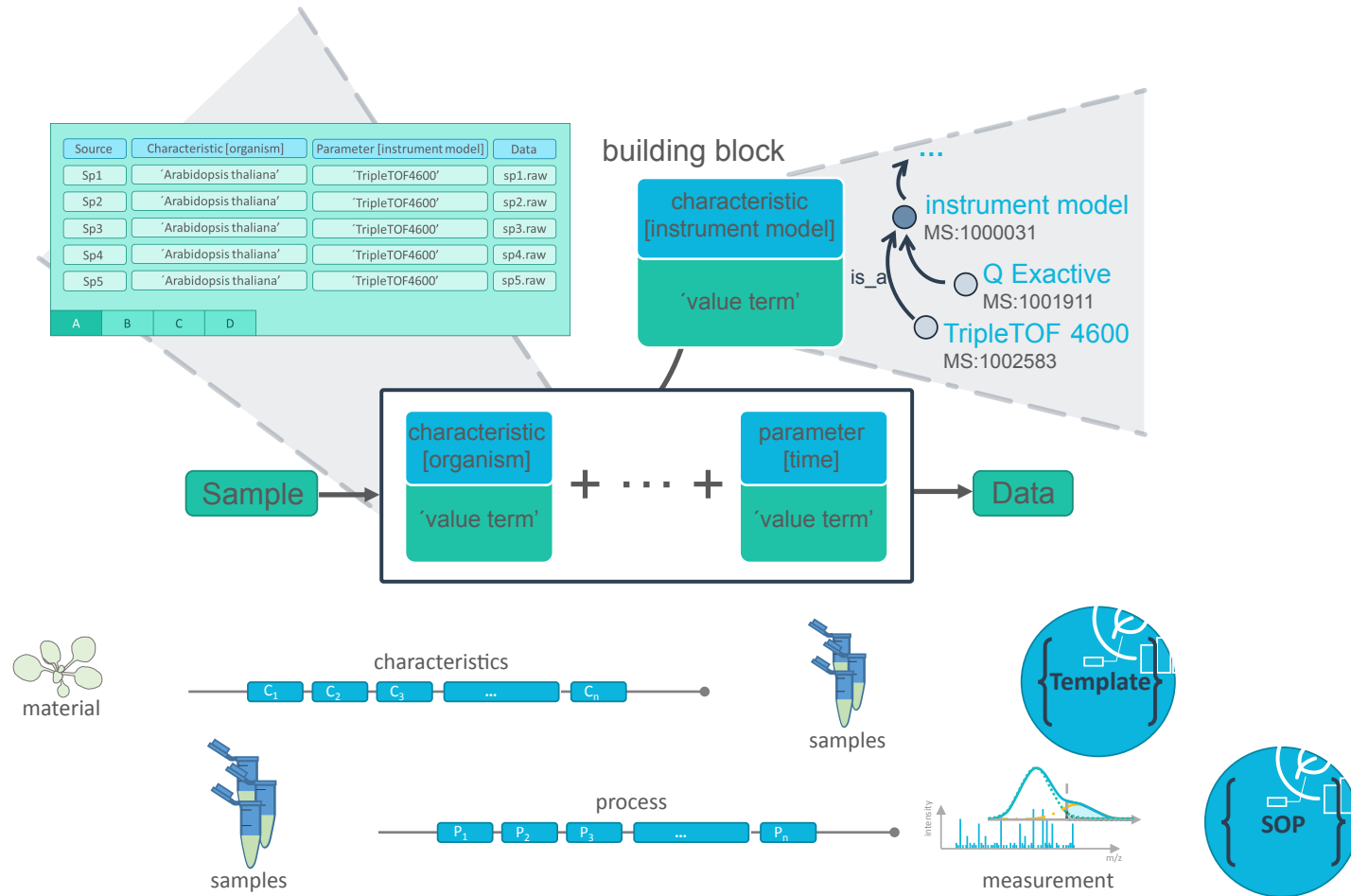
Realization of lab-specific metadata with templates



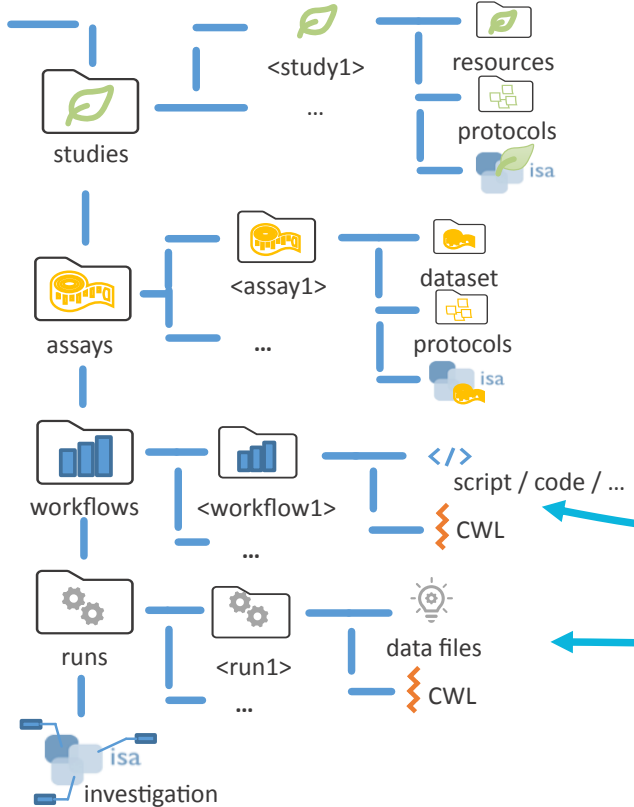
Applying standard procedures to sample record



Save time with templates and SOPs



Data analysis: CWL workflows and runs



```
python
Copy code

import pandas as pd
import plotly.express as px

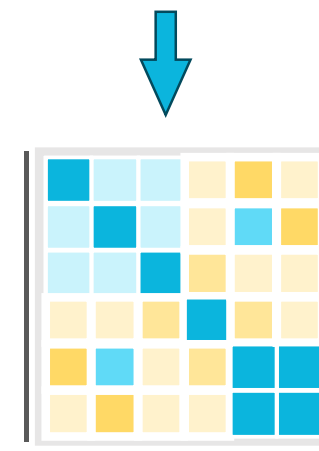
# Read the CSV file
data = pd.read_csv('result.csv')

# If the CSV doesn't have a clear index or row/column names, you may need to manually set
# For example, if the first column should be the index, you can use:
# data = pd.read_csv('result.csv', index_col=0)

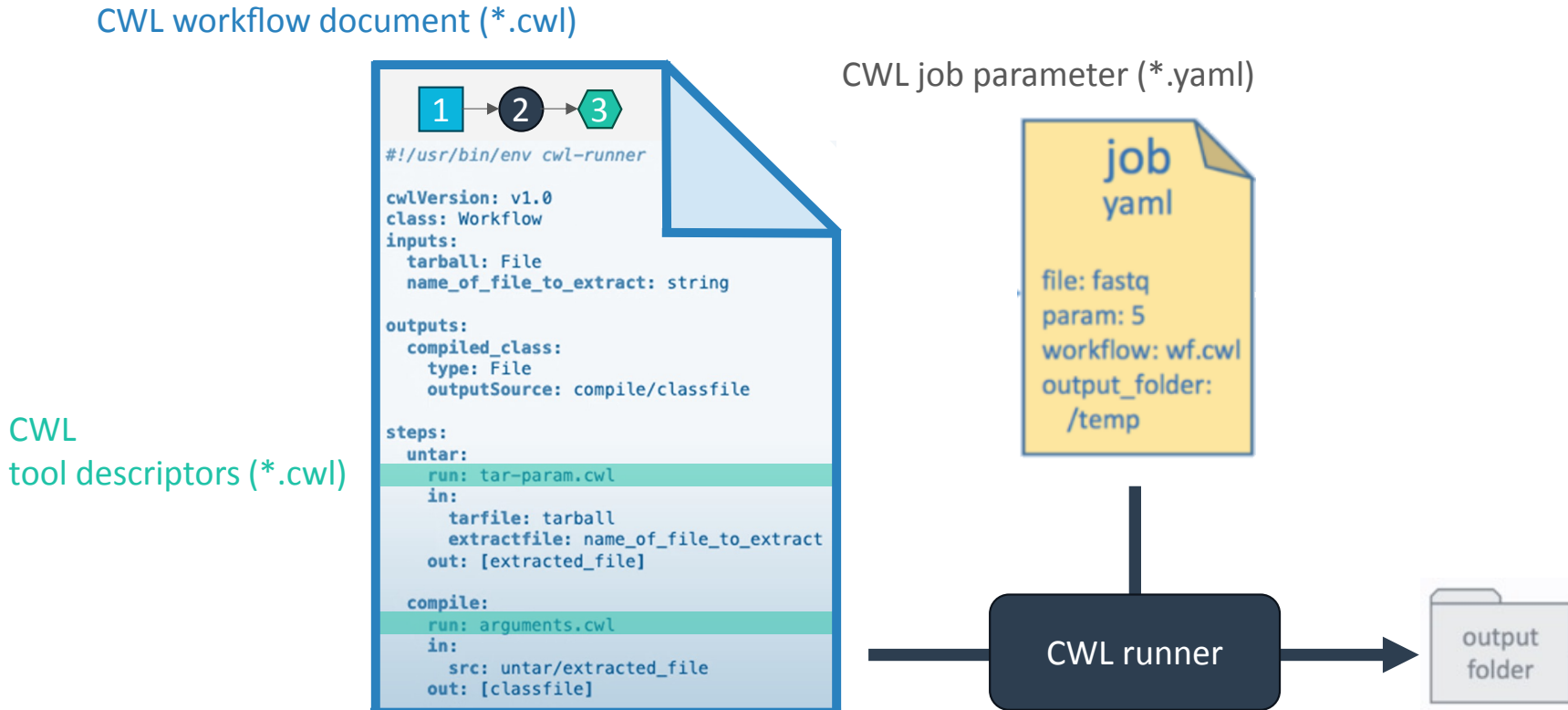
# Create a heatmap using Plotly
fig = px.imshow(data,
                 labels=dict(x="Columns", y="Rows", color="Value"),
                 x=data.columns,
                 y=data.index)

# Show the heatmap
fig.show()
```

heatmap.py

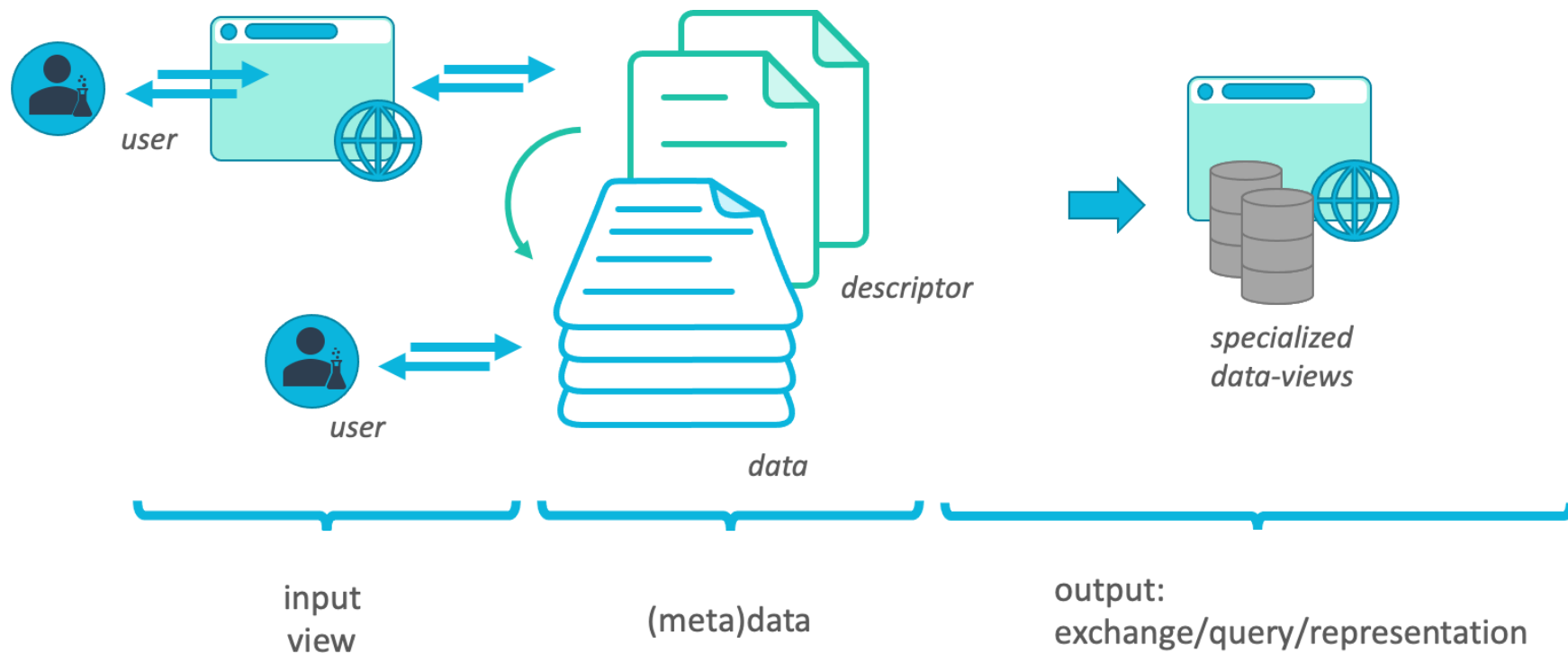


Metadata annotation – from data to result

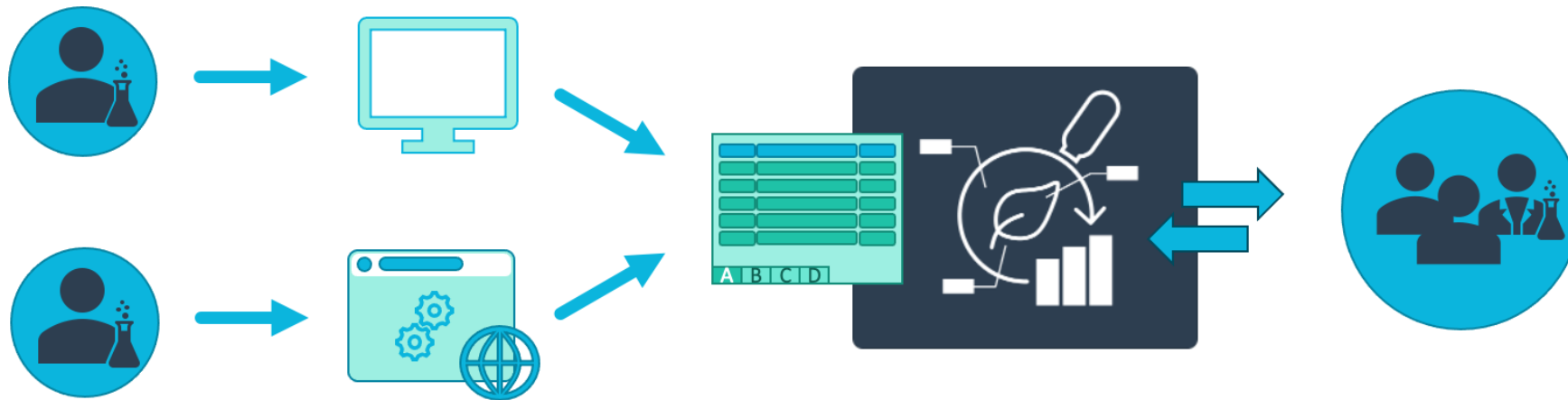


Everything is a file

The ARC is a data-centric approach to RDM

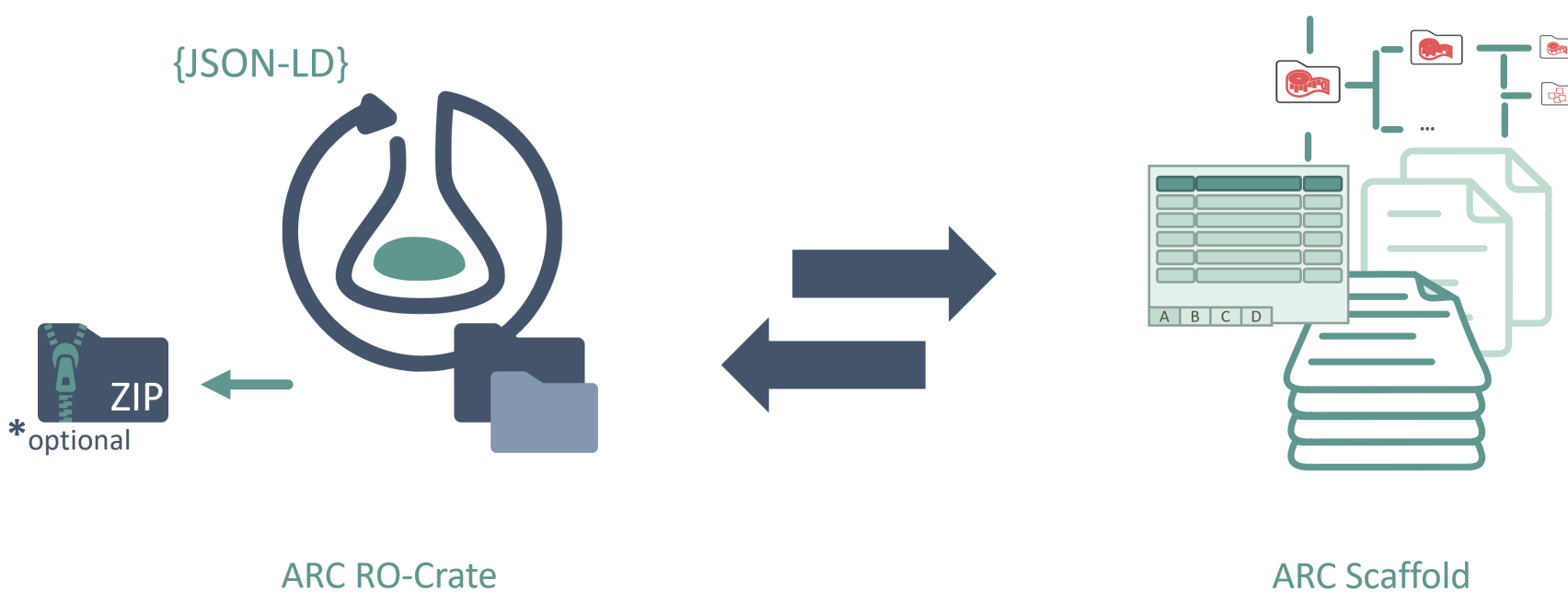


No technical lock-in



(Meta)data transparency with tool assistance but **no technical lock-in**

Two representations of the ARC



Two sides of the same coin

"Developer View": RO-Crate

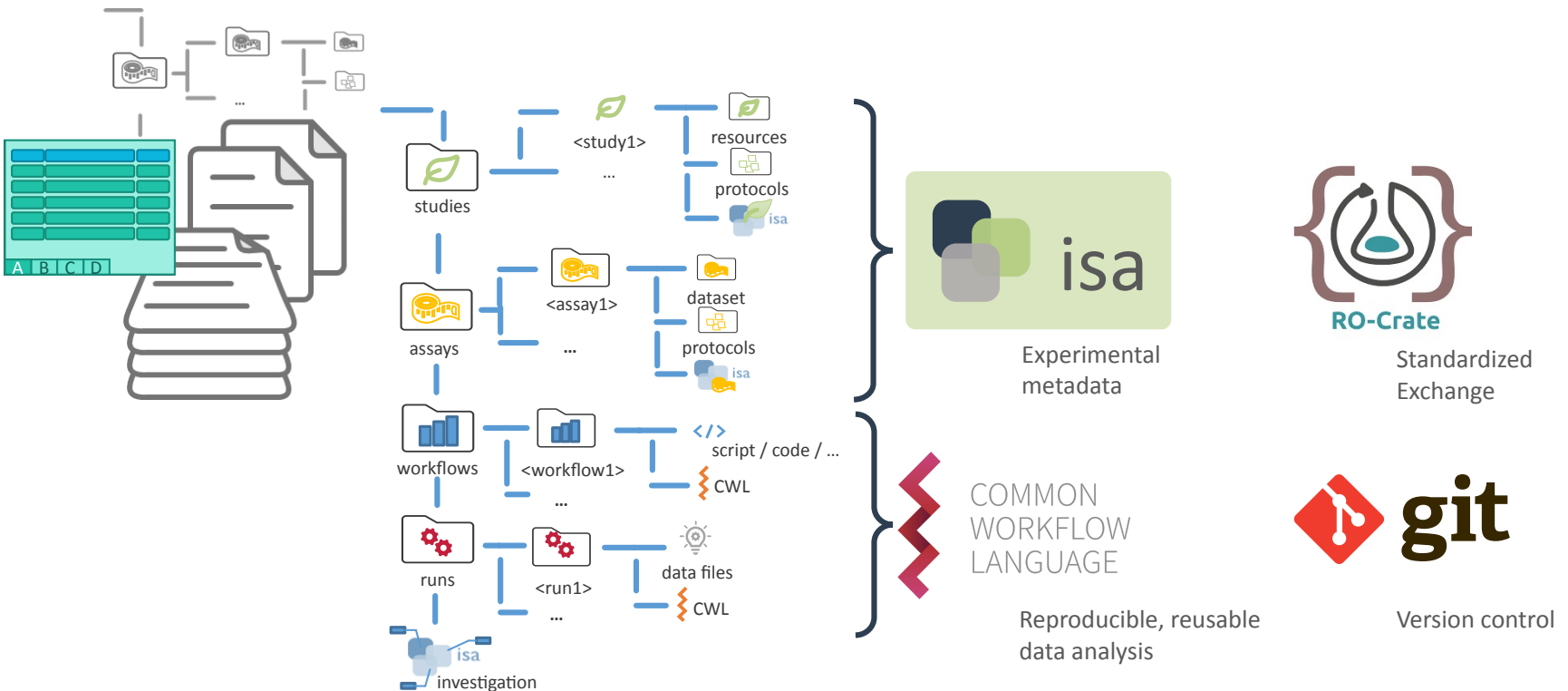
```
{  
  "Identifier": "Proteomics_MS",  
  "MeasurementType": {  
    "annotationValue": "Proteomics_MS",  
    "termSource": "MS",  
    "termAccession": "https://purl.obolibrary.org/obo/FMS_1003348"  
  },  
  "TechnologyType": ...  
  ...  
  "Tables": [  
    {  
      "name": "ProtDigest",  
      "header": [  
        {  
          "headertype": "Parameter",  
          "values": [  
            {  
              "annotationValue": "sample mass",  
              "termSource": "MS",  
              "termAccession": "https://purl.obolibrary.org/obo/FMS_1003348"  
            }  
          ]  
        ]  
      ]  
    }  
  ]  
}
```

"User View": ARC Scaffold and metadata tables

The screenshot shows the ARCIct application interface. On the left, the ARCIct sidebar displays a file structure for a project named 'AthalianaColdStressSugar'. The 'assays' folder contains sub-folders like 'Proteomics_DataAnalysis' and 'SugarMeasurement', each with its own README.md file. On the right, a detailed view of the 'Assay Metadata' for the 'Proteomics_MS' assay is shown. This view includes fields for Identifier (Proteomics_MS), Measurement Type (Proteomics_MS), Technology Type (Mass Spectrometry), Technology Platform (timsTOF Pro 2), and Performers (PeptideMS_Bruker, ProtDigest). Below this, a table lists parameters: 'Parameter [sample mass]' (10 microgram), 'Parameter [Protein Precipitation]' (acetone), 'Parameter [alkylating agent]' (Chloroacetamide), and 'Parameter [red' (TCEP). The ARCIct interface is version v0.0.55.

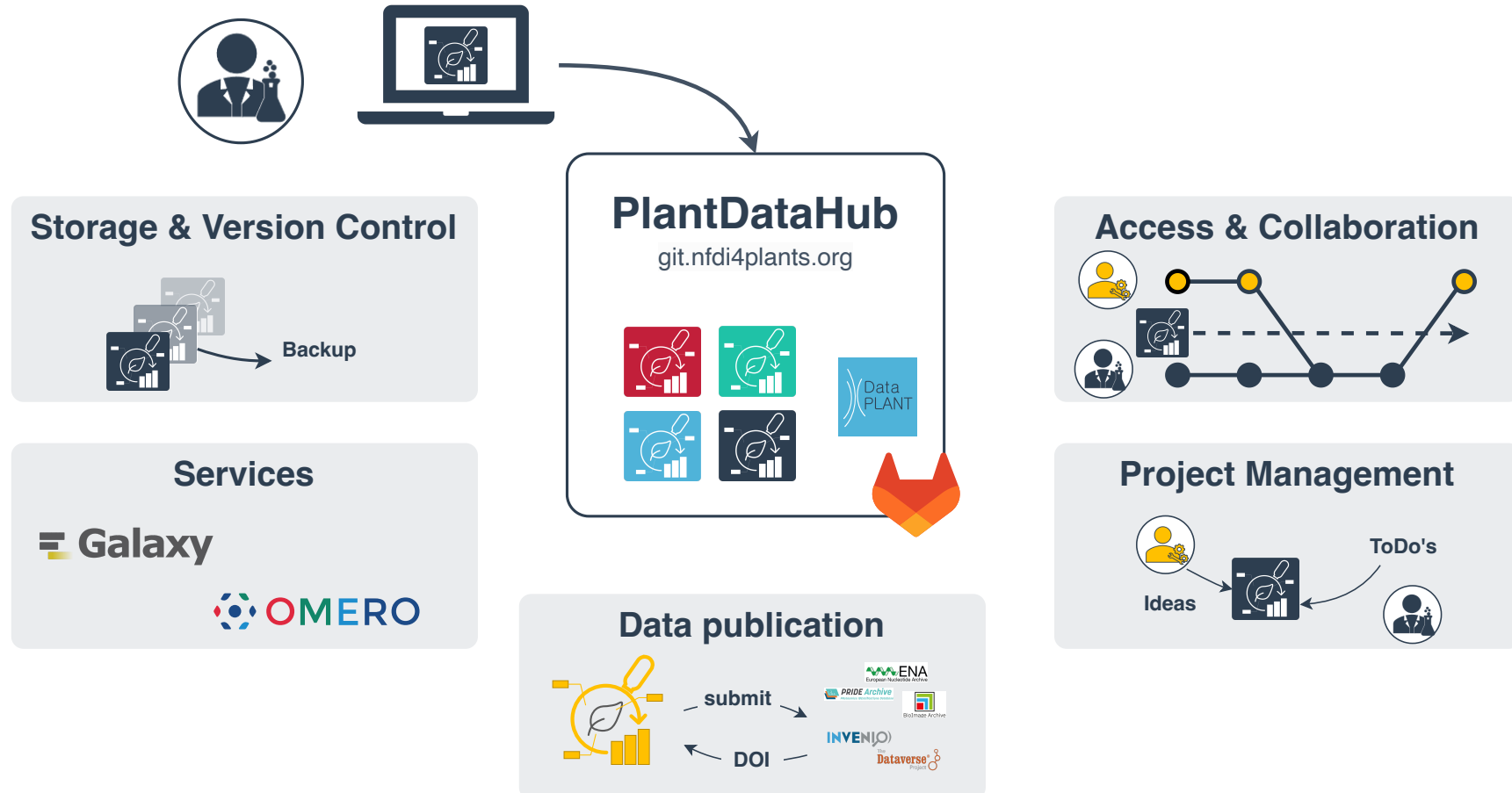
Parameter [sample mass] >>	Parameter [Protein Precipitation] >>	Parameter [alkylating agent] >>	Parameter [red]
10 microgram	acetone	Chloroacetamide	TCEP

ARC builds on standards

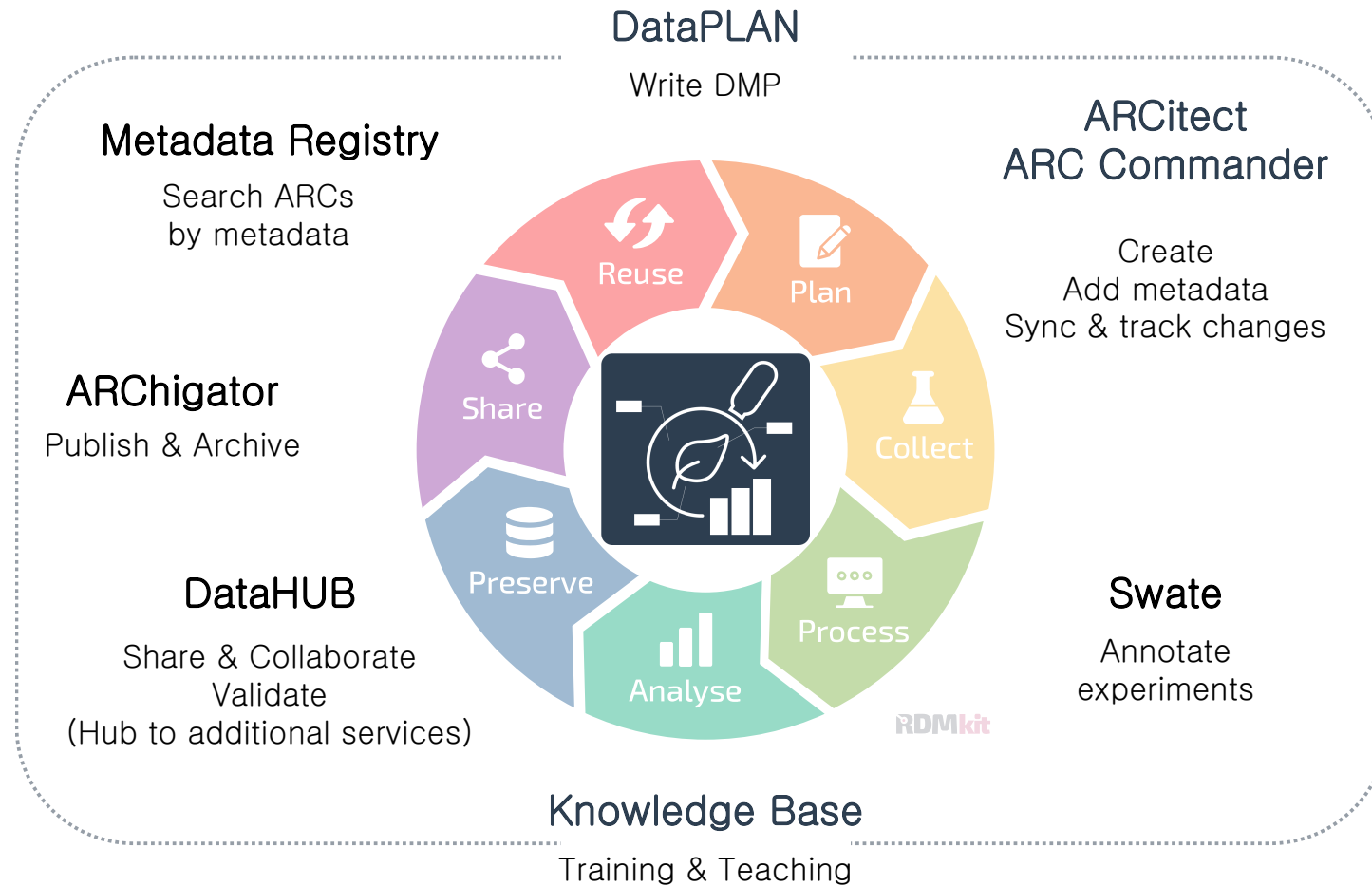


<https://isa-tools.org> | <https://www.commonwl.org> | <https://www.researchobject.org/ro-crate> | <https://git-scm.com>

ARC and DataHUB as entry point



The ARC ecosystem



Acknowledgements



Team Kaiserslautern

- Timo Mühlhaus
- Lukas Weil
- Kevin Frey
- Kevin Schneider
- Jonas Lukasczyk

Team Freiburg

- Dirk von Suchodeletz
- Jonathan Bauer
- Marcel Tschöpe
- Julian Weidhase

Team Jülich

- Stella Eggels
- Angela Kranz



- Björn Usadel
- Vittorio Tracanna
- Yaser Alashloo



- Sabrina Zander

Goals for today

- Get an idea of the ARC concept
- Create and upload a simple ARC
- Explore and discuss ARC features
- ...

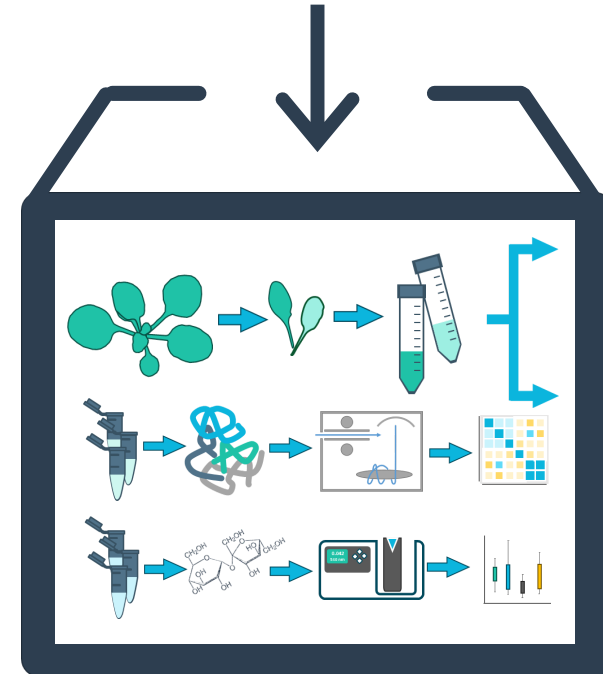
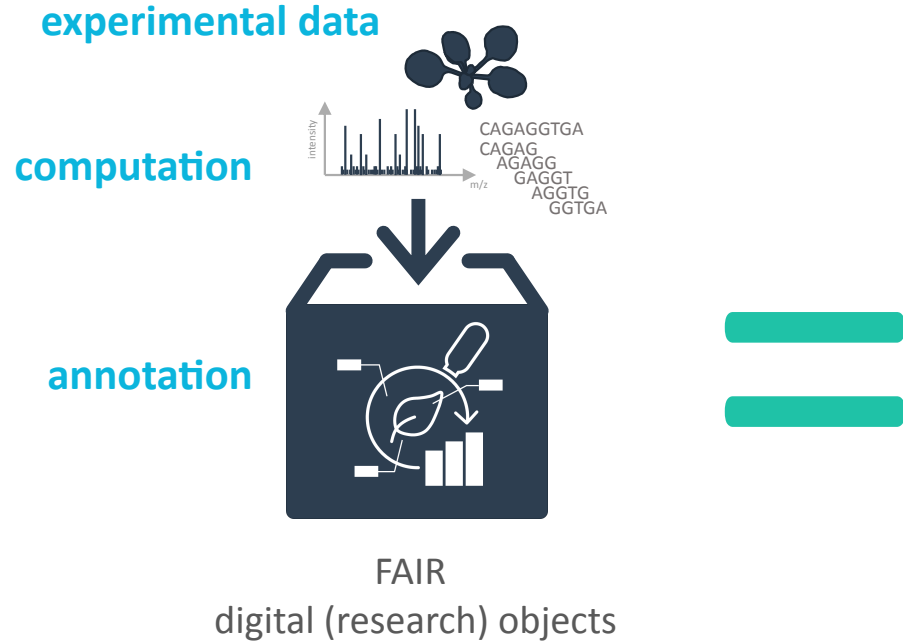
Start an ARC for your investigation with one study



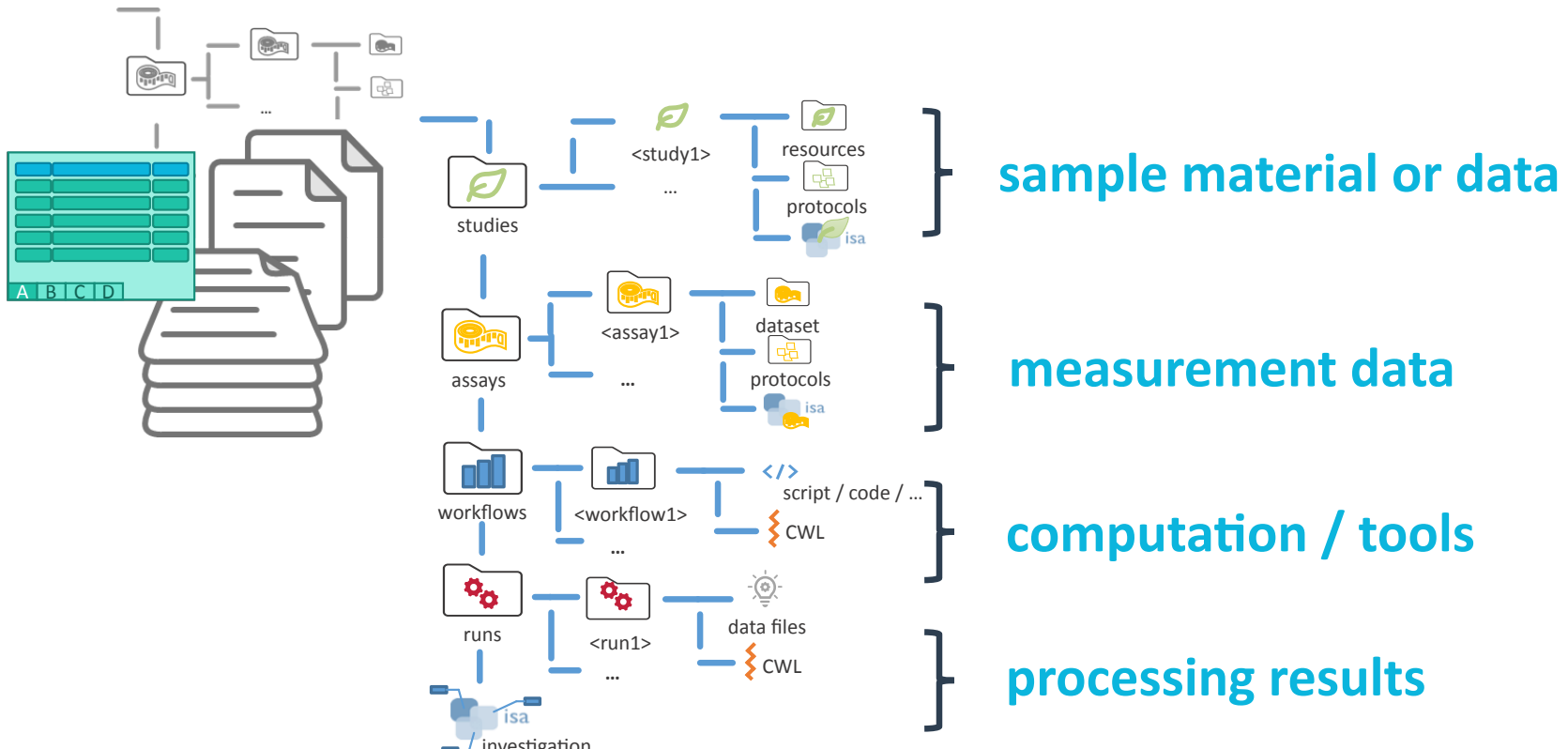
Hands-on

Follow the [Start Here guide](#) until Check point 1

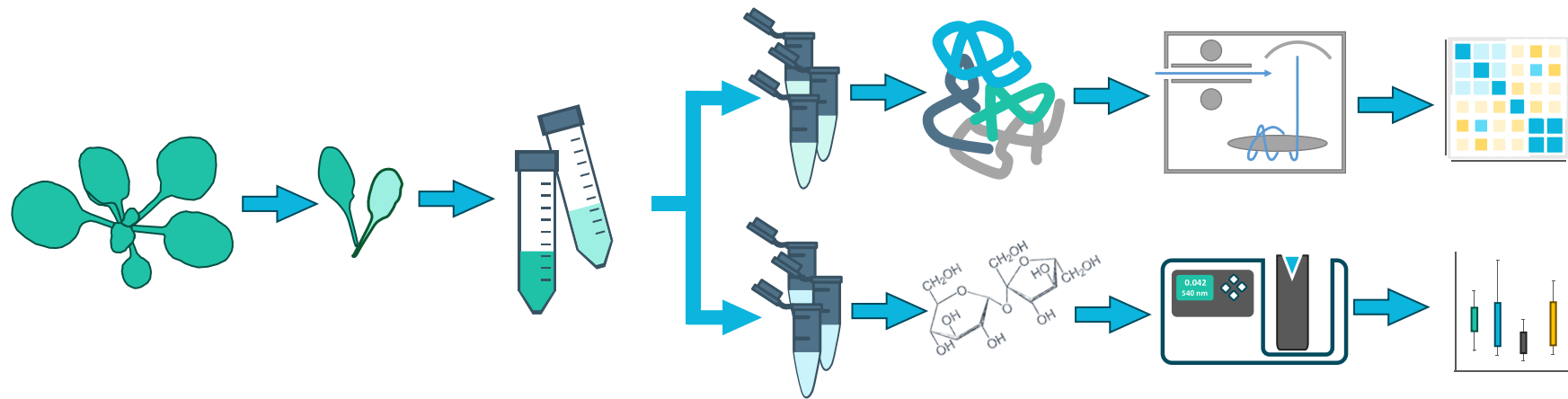
ARC: Annotated research context



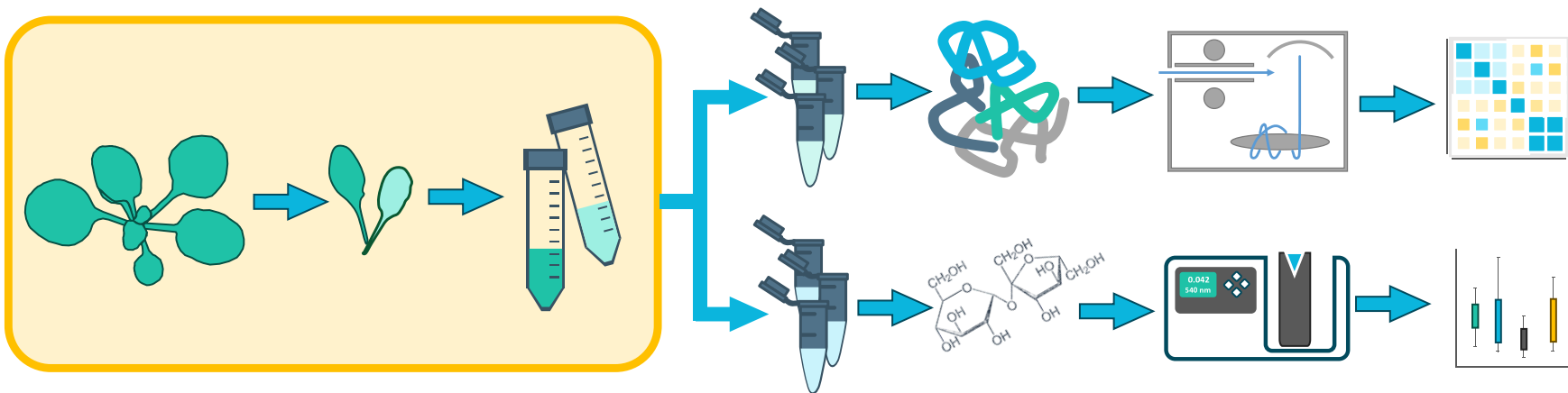
The ARC scaffold structure



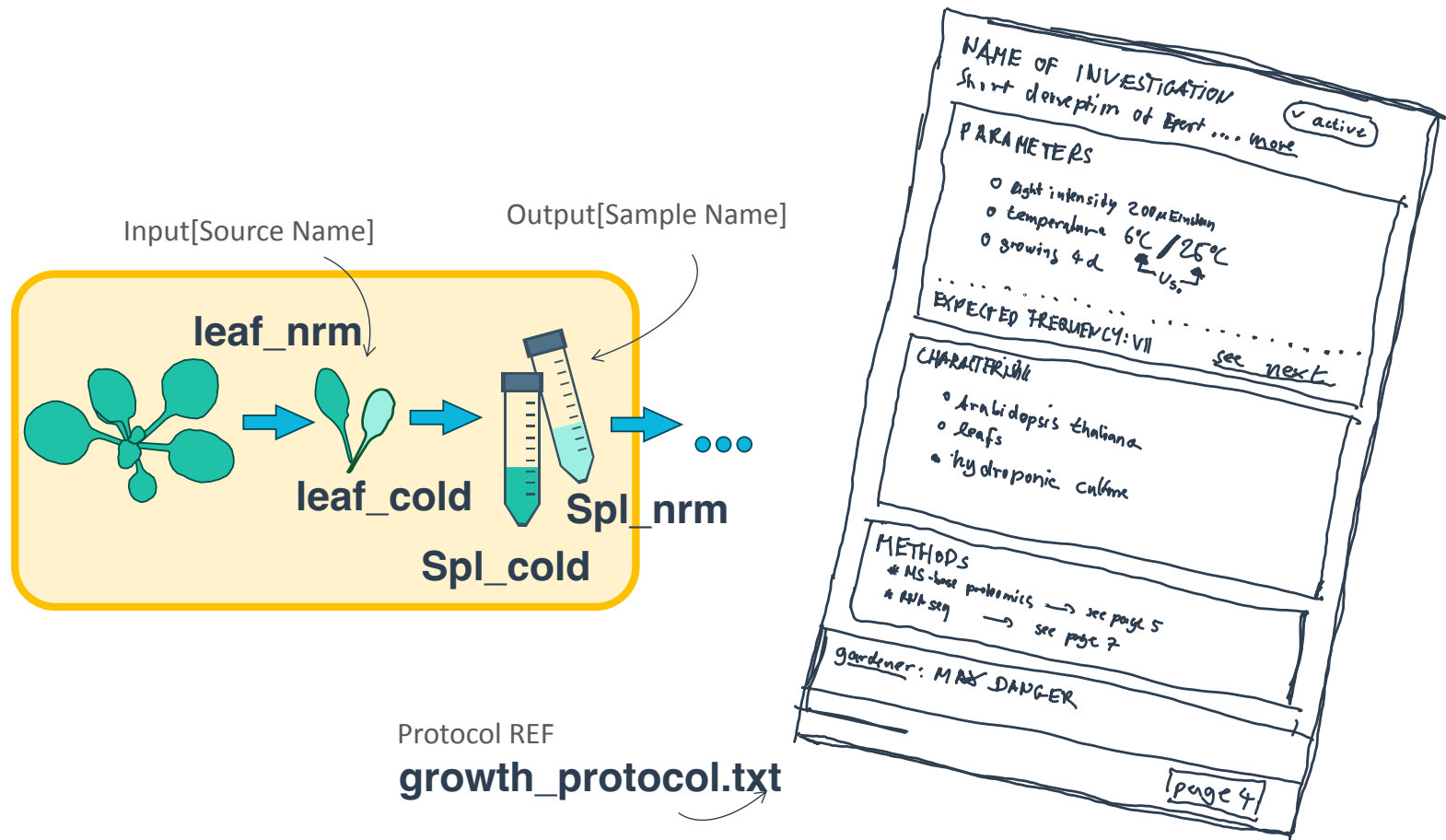
A small prototypic project



Divide and conquer for reproducibility



Identifying the ‘study’ part



A table-based organization schema

Input[Source Name]		Output[Sample Name]
leaf_nrm		spl_nrm
leaf_cold		spl_cold
A	B	C
D		

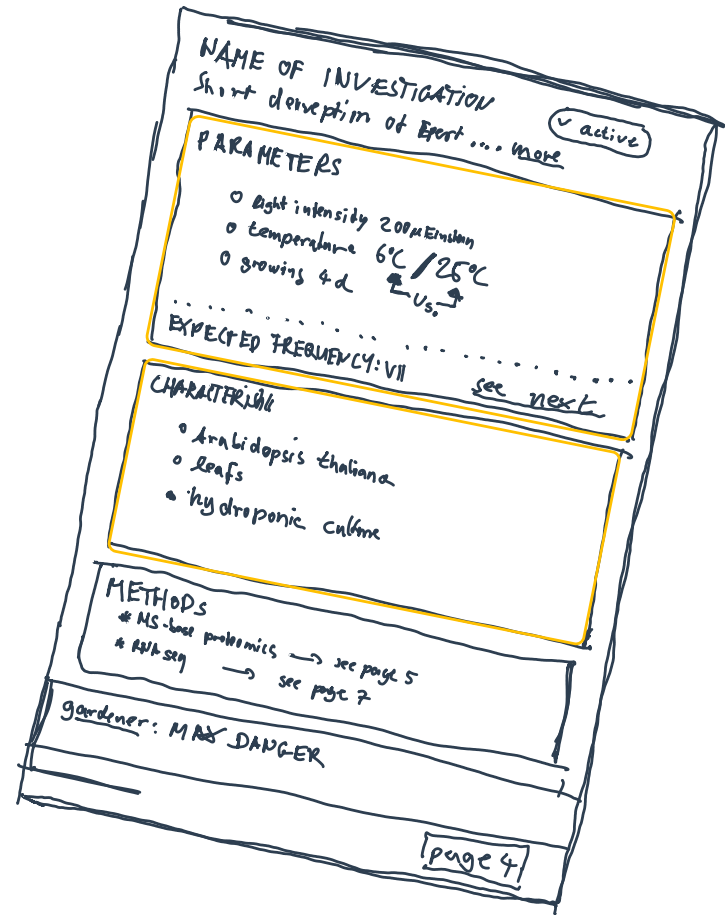
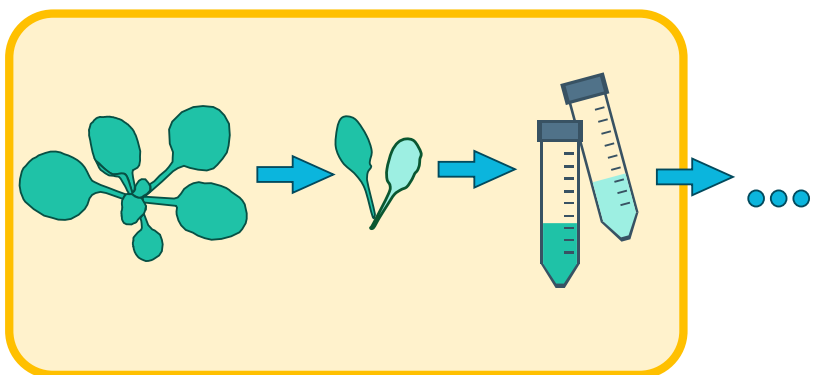
Referencing a protocol

This allows you to reference the free-text, human-readable protocol.

- 💡 It is recommended that the protocol is in an open format (.md|.txt|.docx|…)
- 💡 But everything is possible also an URI to an electronic lab notebook

Input[Source Name]	Protocol REF	Output[Sample Name]
leaf_nrm	growth_protocol.txt	spl_nrm
leaf_cold	growth_protocol.txt	spl_cold
A	B	C
D		

Parameterizing the ‘study’



Finding the right metadata vocabulary

Parameters []

- Light intensity 200 µEinstein
- Temperature 6°C / 25°C
- Growing 4d

Characteristics []

- *Arabidopsis thaliana*
- Leaf
- Hydroponic culture
- Columbia

OLS: Finding the right metadata vocabulary

Temperature Dependence

Temperature:Dependence_Annotation



http://purl.uniprot.org/core/Temperature_Dependence_Annotation

Indicates the optimum temperature for enzyme activity and/or the variation of enzyme activity with temperature variation; the thermostability/thermolability of the enzyme is also mentioned when it is known.

Ontology: UNIPROT RDFS

temperature

AFO:/result#AFR_0001584



http://purl.allotrope.org/ontologies/result#AFR_0001584

A temperature (datum) is a quantity facet that quantifies some temperature. [Allotrope]

Ontology: AFO

temperature

FBcv:0000466



http://purl.obolibrary.org/obo/FBcv_0000466

Mutation caused by exposure to a temperature that is higher or lower than 25 degrees Celsius.

Ontology: FBCV

temperature

PATO:0000146



http://purl.obolibrary.org/obo/PATO_0000146

A physical quality of the thermal energy of a system.

Ontology: PATO

Also appears in:

NGBO

HTN

CAO

ZP

AGRO

OMIABIS

OBIB

MONDO

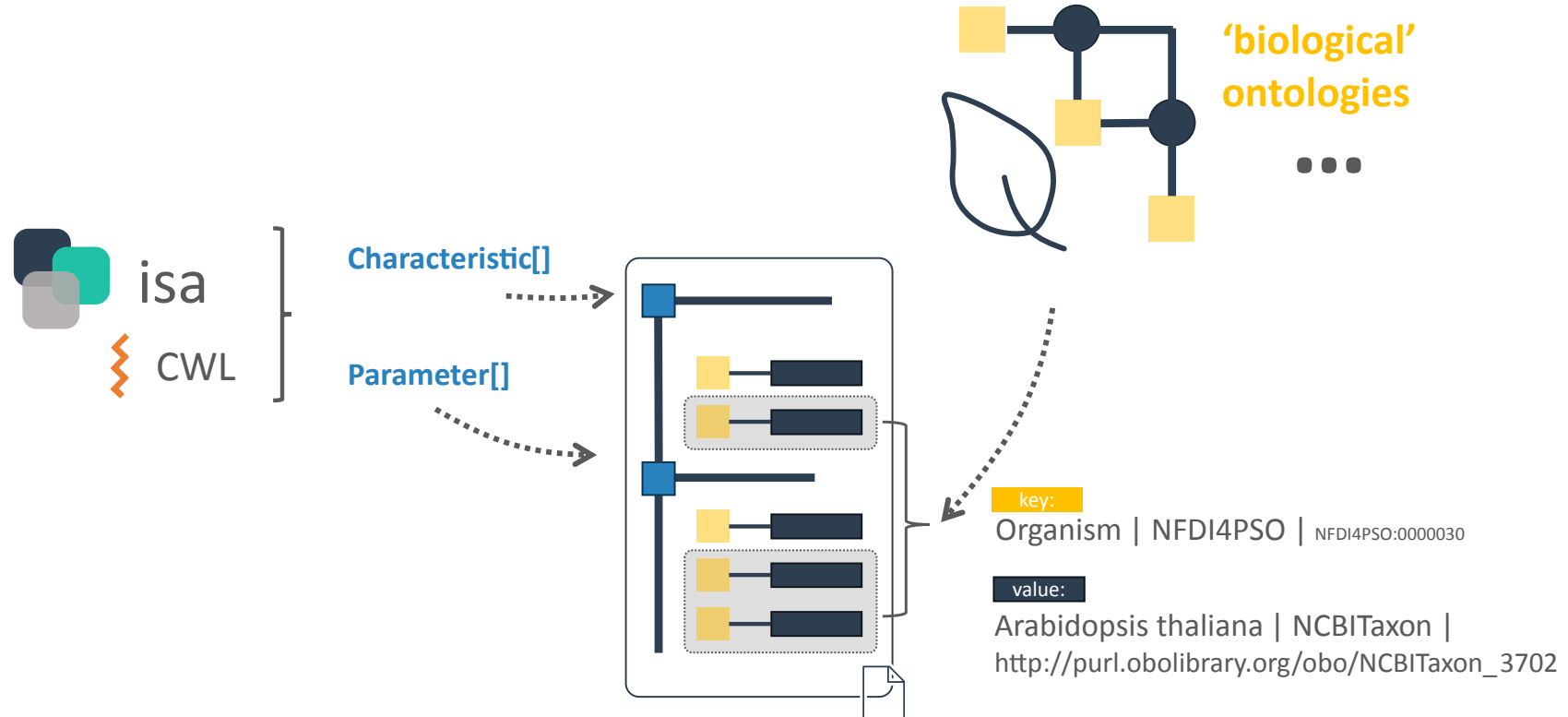
TXPO

MCO

+

Ontology Lookup Service (OLS): <https://www.ebi.ac.uk/ols4/>

Finding the metadata vocabulary and descriptors



Finding the metadata vocabulary and descriptors

Parameters []

- 
 -  Light intensity 200 µEinstein
- 
 -  Temperature 6°C / 25°C
- 
 -  Growing 4d

Characteristics []

- 
 -  Arabidopsis thaliana
- 
 -  Leaf
- 
 -  Hydroponic culture
- 
 -  Columbia

Finding the metadata vocabulary and descriptors

Parameters []

-  Light intensity
 -  200 µEinstein
-  Temperature
 -  6°C / 25°C
-  Growth time
 -  4d

Characteristics []

-  Organism
 -  Arabidopsis thaliana
-  Tissue
 -  Leaf
-  Growth medium
 -  Hydroponic culture
-  Ecotype
 -  Columbia

Check point 1

Task

- ✓ I've created an ARC for my project!
- ✓ I've added metadata about the overall investigation
- ✓ I've added a study for my samples

ARCify your research project

Hands-on ARC introduction @ IRTG 2843

Dominik Brilhaus, [CEPLAS](#)

October 30th, 2025



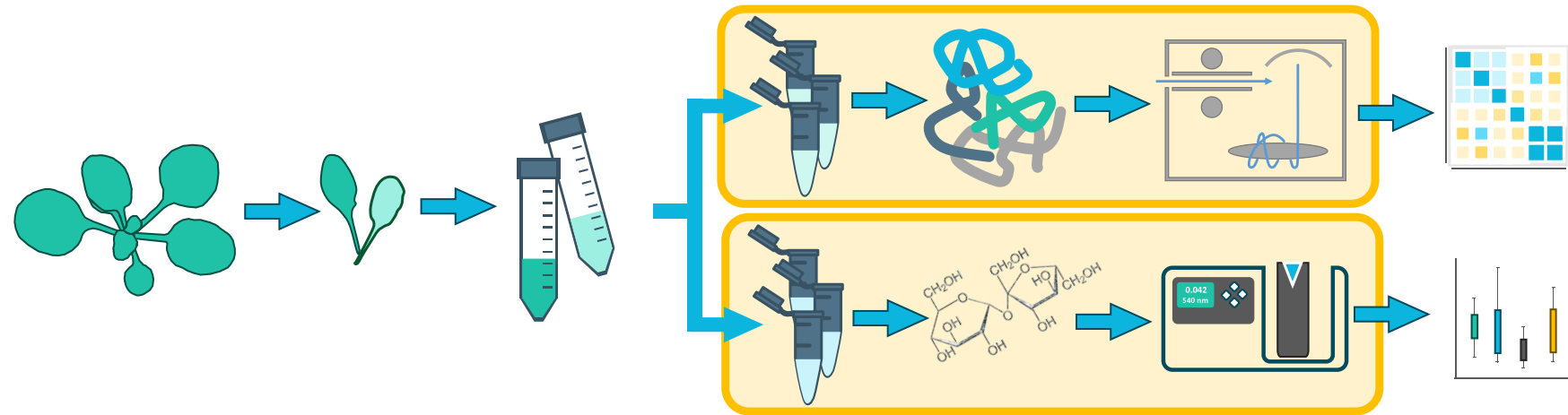
Add an assay and share your ARC



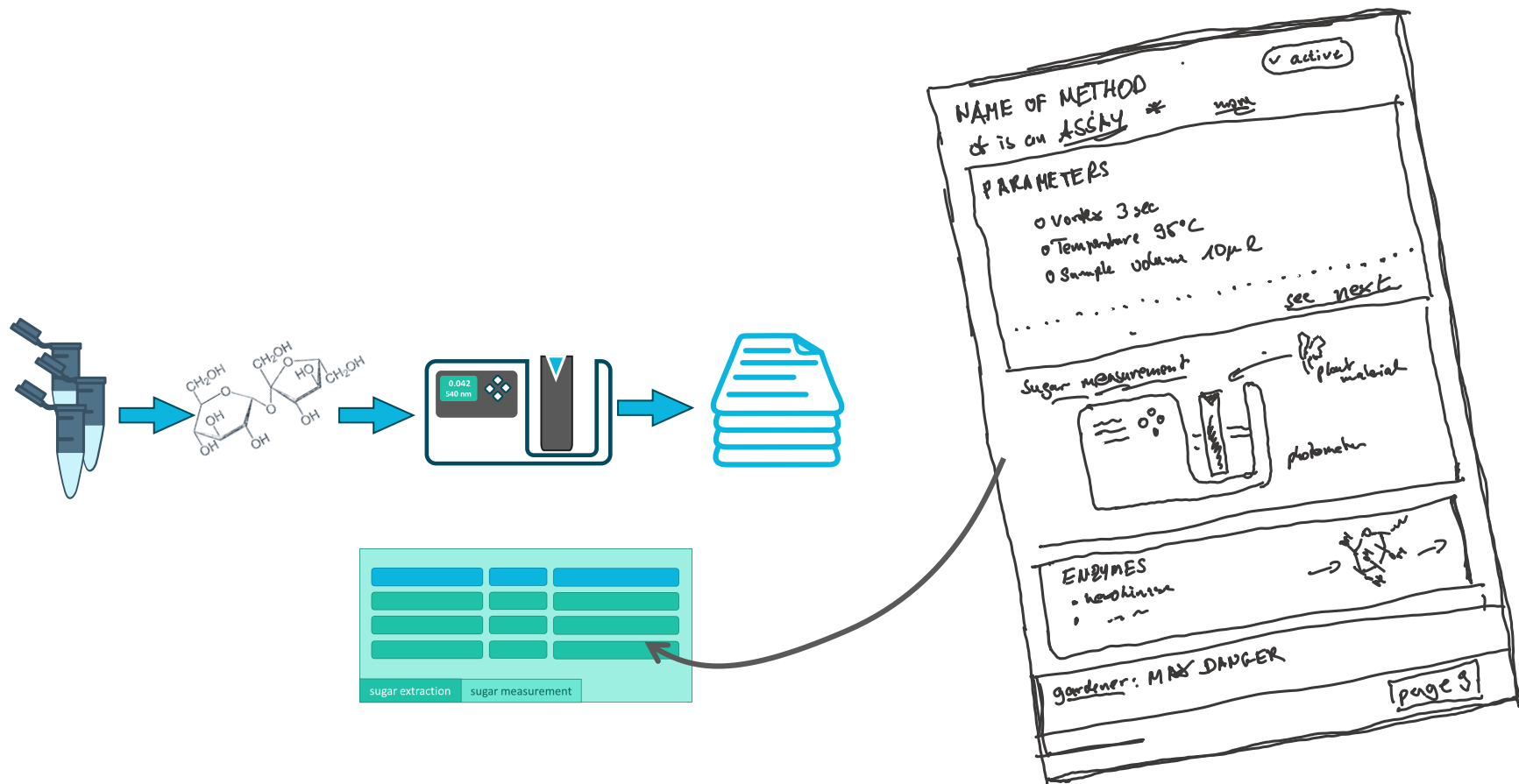
Hands-on

Follow the [Start Here guide](#) until Check point 2

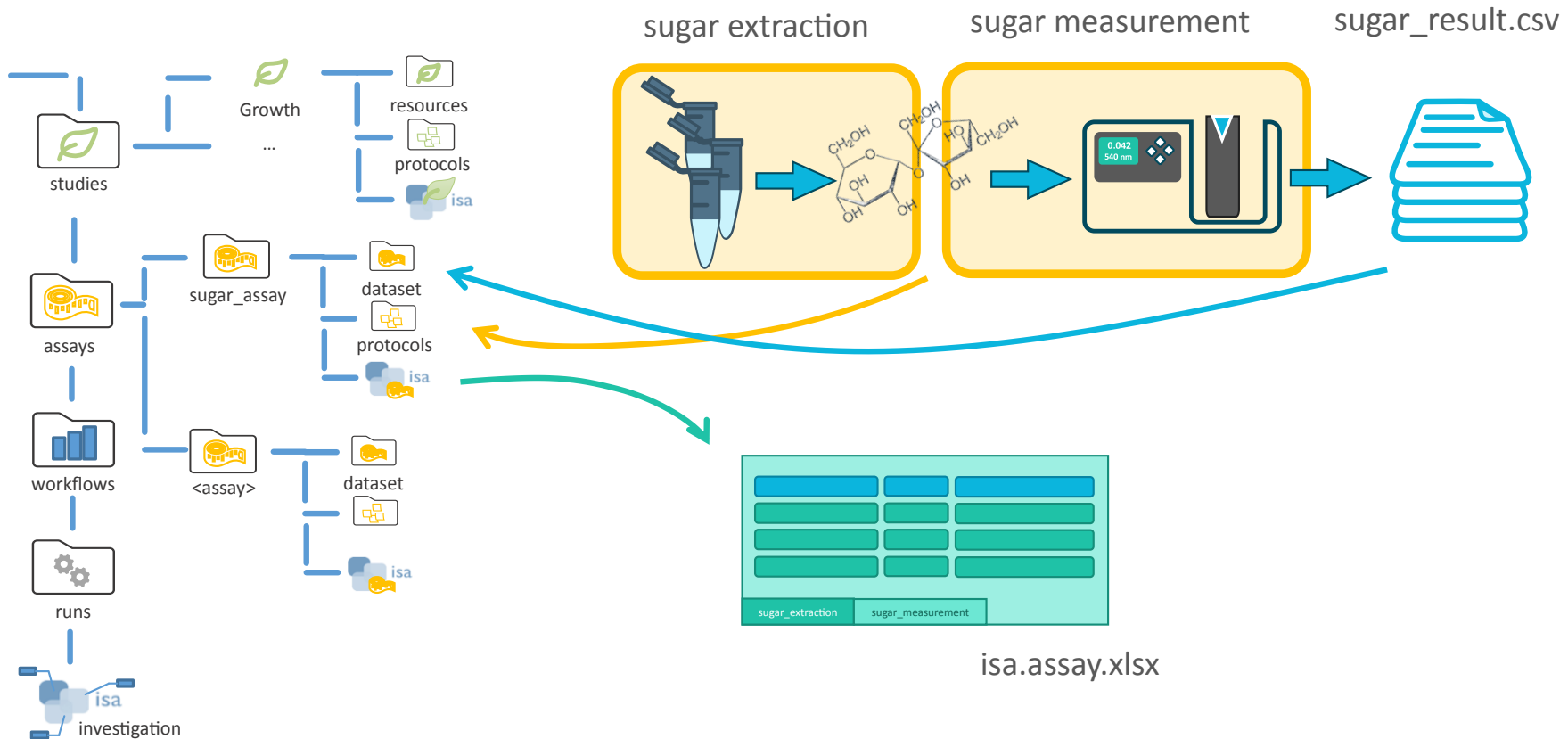
Identifying assays



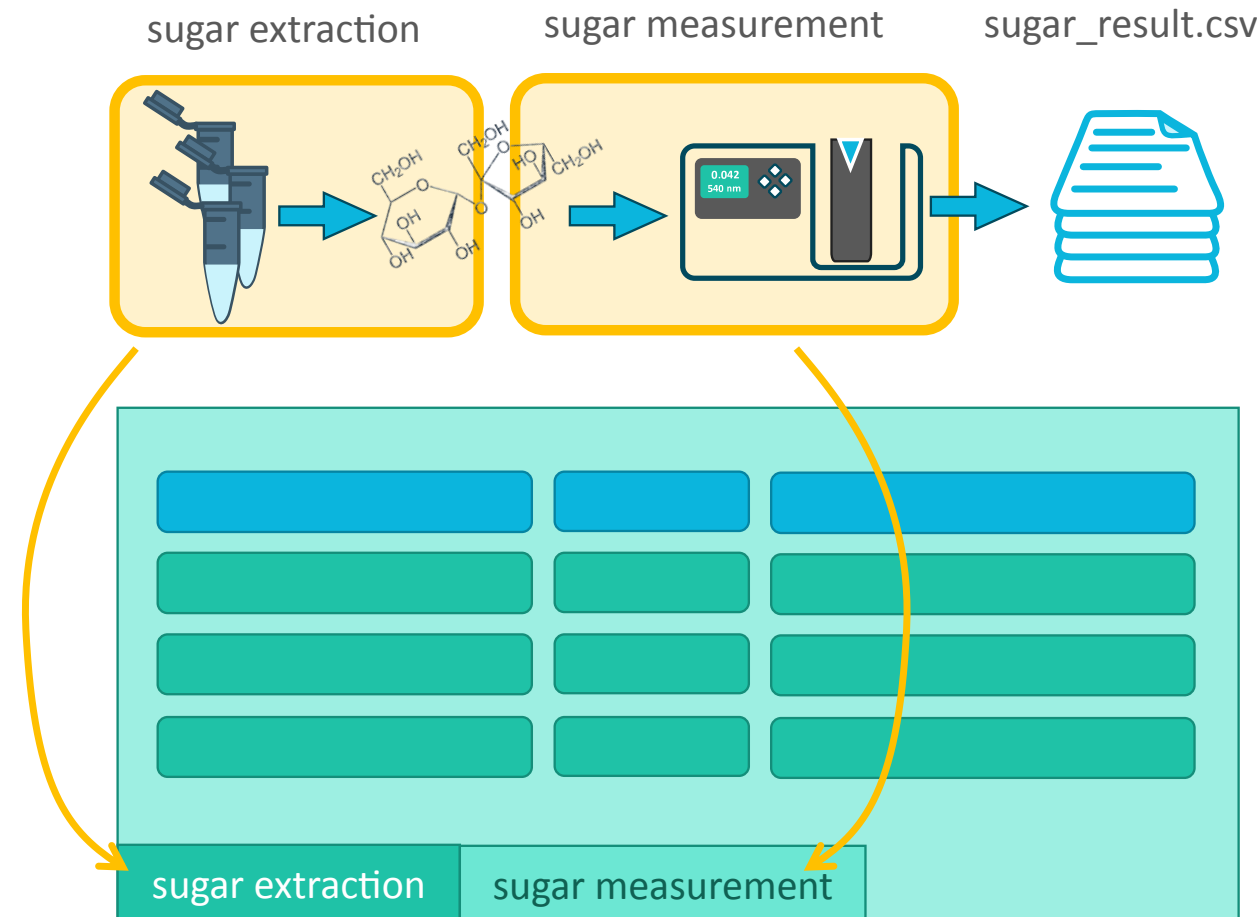
Assay for sugar measurement



Separating different assay elements



Isolating the lab processes in an assay



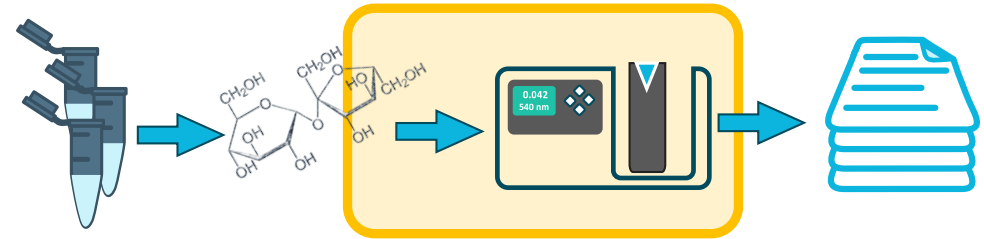
Parameteterization: sugar extraction

- Vortex Mixer
 - 3 seconds
- Temperature
 - 95 degree celsius



Parameteterization: sugar measurement

- ■ technical replicate
 - ■ 1,2,3,...
- ■ sample volume
 - ■ 10 microliter
- ■ buffer volume
 - ■ 190 microliter
- ■ cycle count
 - ■ 5



DataHUB terminology and data sharing

Choosing the proper role

When inviting new members to an ARC or group, you can choose between different levels.

Permissions & Roles

Roles are assigned when adding a user to an ARC or to a group. This is a very simplified summary.

Guest – Can only see the ARC's wiki

Reporter – Can read, but not add or edit data

Developer – Reporter permissions + can read, add, and edit data

Maintainer – Developer permissions + can add new members

Owner – Maintainer + can delete ARC, manage memberships and permissions

💡 By default you are **Owner** of an ARC you create or upload to the DataHUB.

Projects and Groups are not the same

- "Project" = ARC
- "Groups" = Group of users

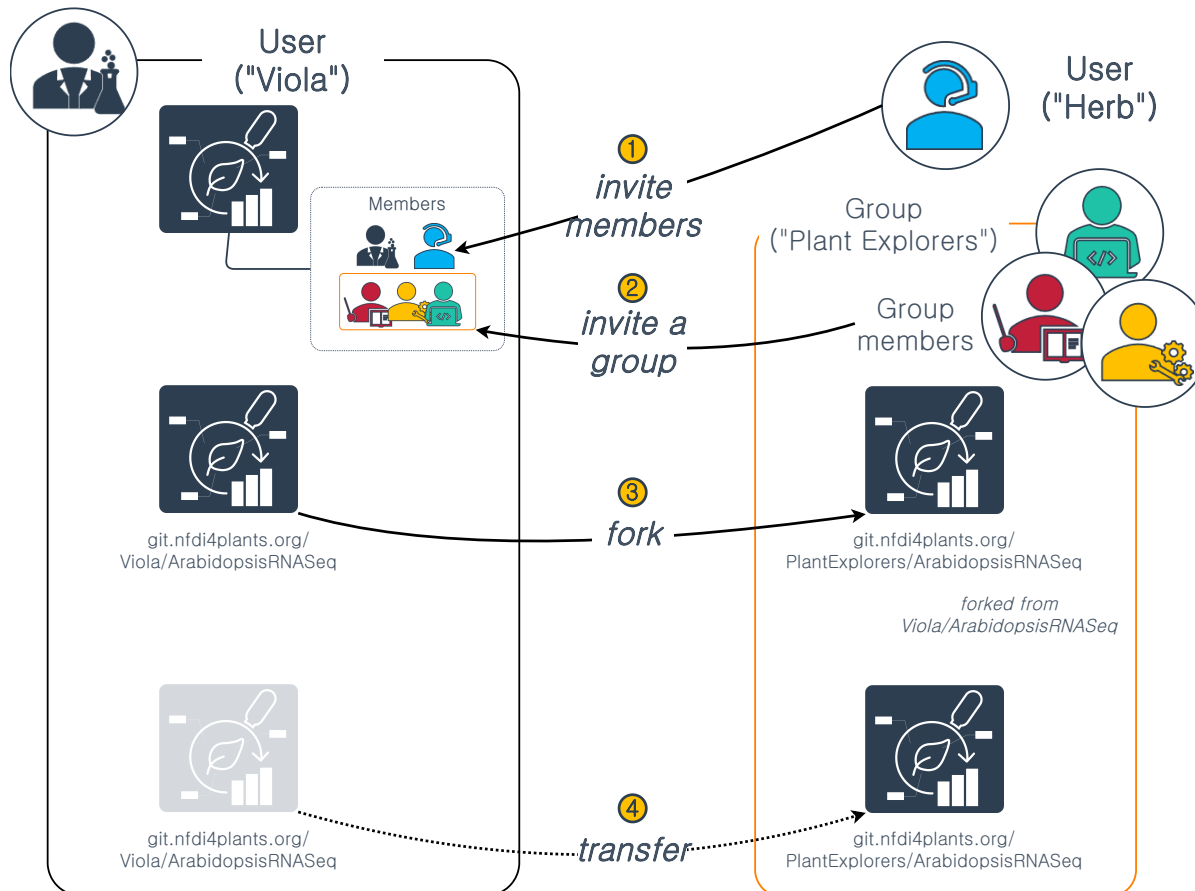
Project = ARC

- In the DataHUB, ARCs are called "projects"; they are the same.
- An ARC can be shared with individual users (invited as "members") or a group.

DataHUB Groups = Group of members (e.g. lab)

- A "Group" is a group of users with specific [permissions](#)
- A group can share ARCs
- A group can be invited to an ARC
- Groups can have subgroups

Options to share an ARC via the DataHUB



Namespaces

- Every user has a personal namespace, where they can upload or create new ARCs
- Every group and subgroup has its own namespace

Type	URL	Namespace	Name
A personal ARC	https://git.nfdi4plants.org/brilator/Facultative-CAM-in-Talinum	brilator	Dominik Brilhaus
An group-shared ARC	https://git.nfdi4plants.org/hhu-plant-biochemistry/Samuilov-2018-BOU-PSP	hhu-plant-biochemistry	HHU Plant Biochemistry

Visibility

The visibility of ARCs and groups can be managed individually for each ARC or group

Visibility

The visibility of each ARC can be managed in the settings of the ARC

Private – ARC access must be granted explicitly to each user or group.

Internal – ARC can be accessed by any logged in user.

Public – ARC can be accessed without authentication.

💡 By default every ARC and every group is set to **private**.

ARC DataHUB members // ARC Investigation contacts

The screenshot shows the 'Project members' section of the DataHUB interface. It includes a search bar, navigation links for 'Import from a project', 'Invite a group', and 'Invite members'. Below this, there are tabs for 'Members' (18) and 'Groups' (1). A filter bar allows users to 'Filter members'. The main table lists four members:

Account	Source	Max role	Expiration	Activity
Adriano Nunes-Nesi @nunesnesi	HHU Plant Biochemistry by Sebastian Triesch	Maintainer	Expiration date	User created: Jul 05, 2023 Access granted: Jul 10, 2023 Last activity: Aug 09, 2023
Andreas Weber @andreas.weber	HHU Plant Biochemistry by Sebastian Triesch	Maintainer	Expiration date	User created: Mar 10, 2023 Access granted: Jul 31, 2023 Last activity: Sep 11, 2023
Dominik Brilhaus It's you @brilator	Direct member by Dominik Brilhaus	Owner	Expiration date	User created: Feb 21, 2022 Access granted: Dec 06, 2023 Last activity: Mar 26, 2024
Franziska Fichtner @franziska.fichtner	HHU Plant Biochemistry by Sebastian Triesch	Maintainer	Expiration date	User created: Aug 11, 2023 Access granted: Aug 11, 2023 Last activity: Aug 11, 2023

DataHUB: ARC members

https://git.nfdi4plants.org/hhu-plant-biochemistry/Samuilov-2018-BOU-PSP/-/project_members

The screenshot shows the 'ARCIct' interface for the 'Samuilov-2018-BOU-PSP' project. The left sidebar includes 'Login', 'New ARC', 'Open ARC', 'Download ARC', 'Save ARC', 'Explorer', 'Commit', 'DataHUB Sync', and 'History'. The main area displays the project structure and contact information.

Identifier: Samuilov-2018-BOU-PSP

Description: (empty)

Contacts:

Contact	Score
Sladjana Samuilov <orcid>	4/10
Nadine Rademacher <orcid>	3/10
Samantha Flachbart <orcid>	3/10
Leila Arab <orcid>	3/10
Saleh Alfarraj <orcid>	3/10
Franziska Kuhnert <orcid>	3/10
Stanislav Kopriwa <orcid>	3/10
Andreas P. M. Weber <orcid>	4/10
Tabea Mettler-Altmann <orcid>	3/10

ARCIct: Investigation Contacts

- 💡 Investigation contacts are not automatically invited as members to the ARC.

Version control

Check out the **commit history** of your ARC via Repository (2) or directly via commits (7)

The screenshot shows the DataPLANT interface with the following numbered callouts:

- Manage
- Plan
- Code
- Build
- Secure
- Deploy
- Operate
- Monitor
- Analyze
- Settings
- Help

Key UI elements include:

- D Demo_ARC**: Project name in the header.
- Code**: Selected tab in the top navigation bar.
- History**: Tab in the top navigation bar.
- Find file**: Tab in the top navigation bar.
- Edit**: Tab in the top navigation bar.
- Project information**: Pipeline status (passed), Star count (0), Fork count (0), and a three-dot menu.
- 1 Commit**: Summary of project activity.
- 2 Branches**: Summary of branches.
- 0 Tags**: Summary of tags.
- 7 KIB Project Storage**: Summary of storage usage.
- Auto DevOps enabled**: Project status.
- Add README**, **Add LICENSE**, **Add CHANGELOG**, **Add CONTRIBUTING**, **Add Kubernetes cluster**, **Add Wiki**, **Configure Integrations**: Action buttons.
- Created on**: Date of project creation (July 13, 2024).

The commit history table lists the following files and their last commit details:

Name	Last commit	Last update
assays	arc init	4 minutes ago
runs	arc init	4 minutes ago
studies	arc init	4 minutes ago
workflows	arc init	4 minutes ago
.gitignore	arc init	4 minutes ago
isa.investigation.xlsx	arc init	4 minutes ago

Check point 2

Task

- ✓ I've added measurement data as an assay
- ✓ I've uploaded my ARC to the DataHUB to have a version-controlled save copy
- ✓ I've shared the ARC with colleagues, so they can explore my findings and contribute

Goals for today

- Continue preparing Viola's ARC
- Start to create an ARC for your own project
- More parts of the ARC ecosystem
 - ARC Commander
 - ARCtrl
 - Templates
 - Workflows / Runs

Use SOPs and annotate your data analysis



Hands-on

Follow the [Start Here guide](#) until Check point 3

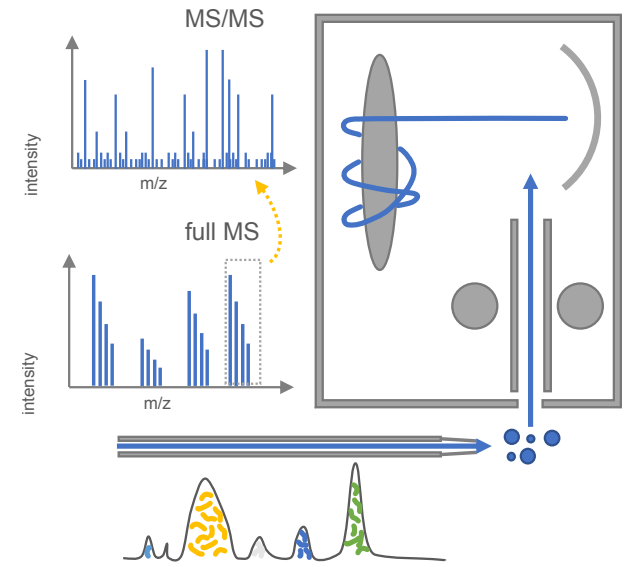
Save time using standard methods and SOPs

Parameter []

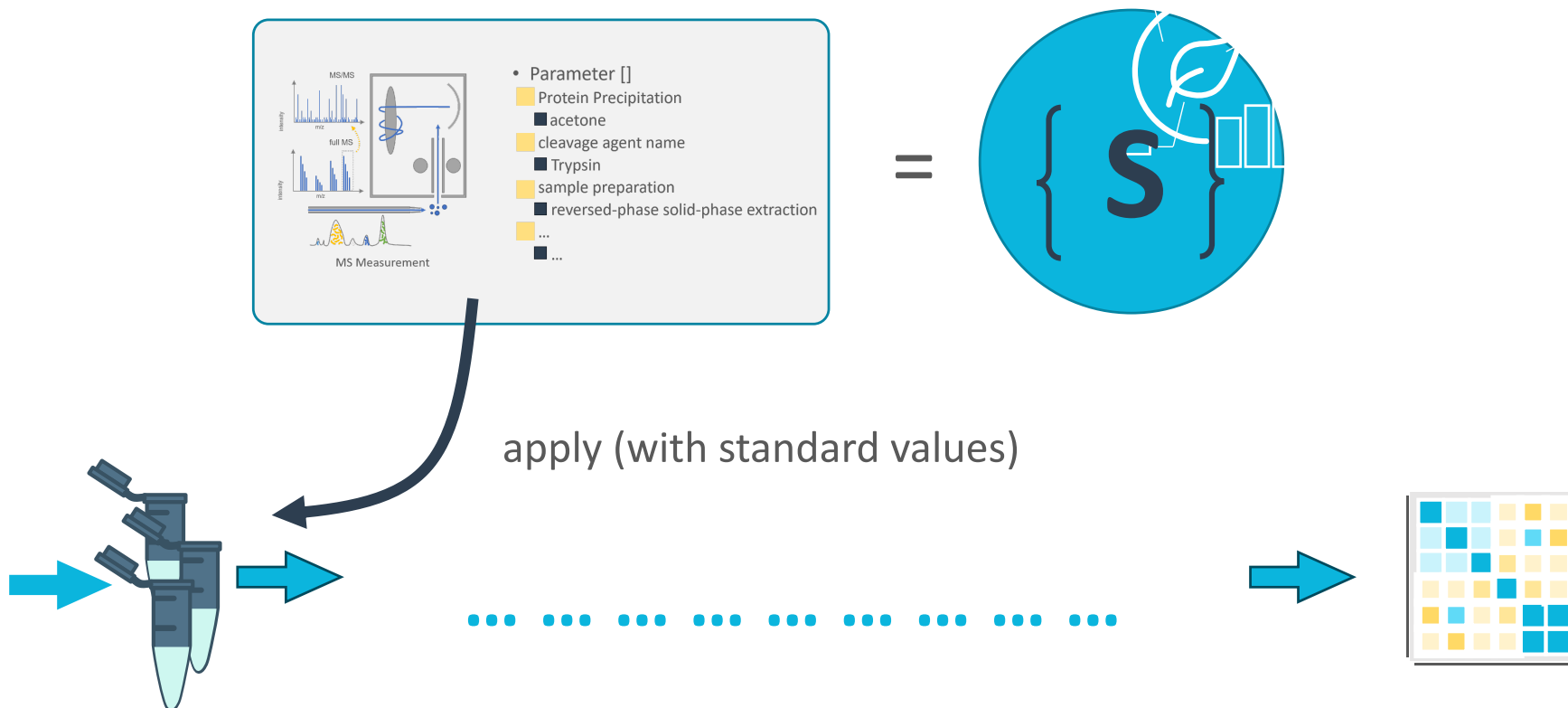
- ■ Protein Precipitation
 - ■ acetone
- ■ cleavage agent name
 - ■ Trypsin
- ■ sample preparation
 - ■ reversed-phase solid-phase extraction
- ...

Component []

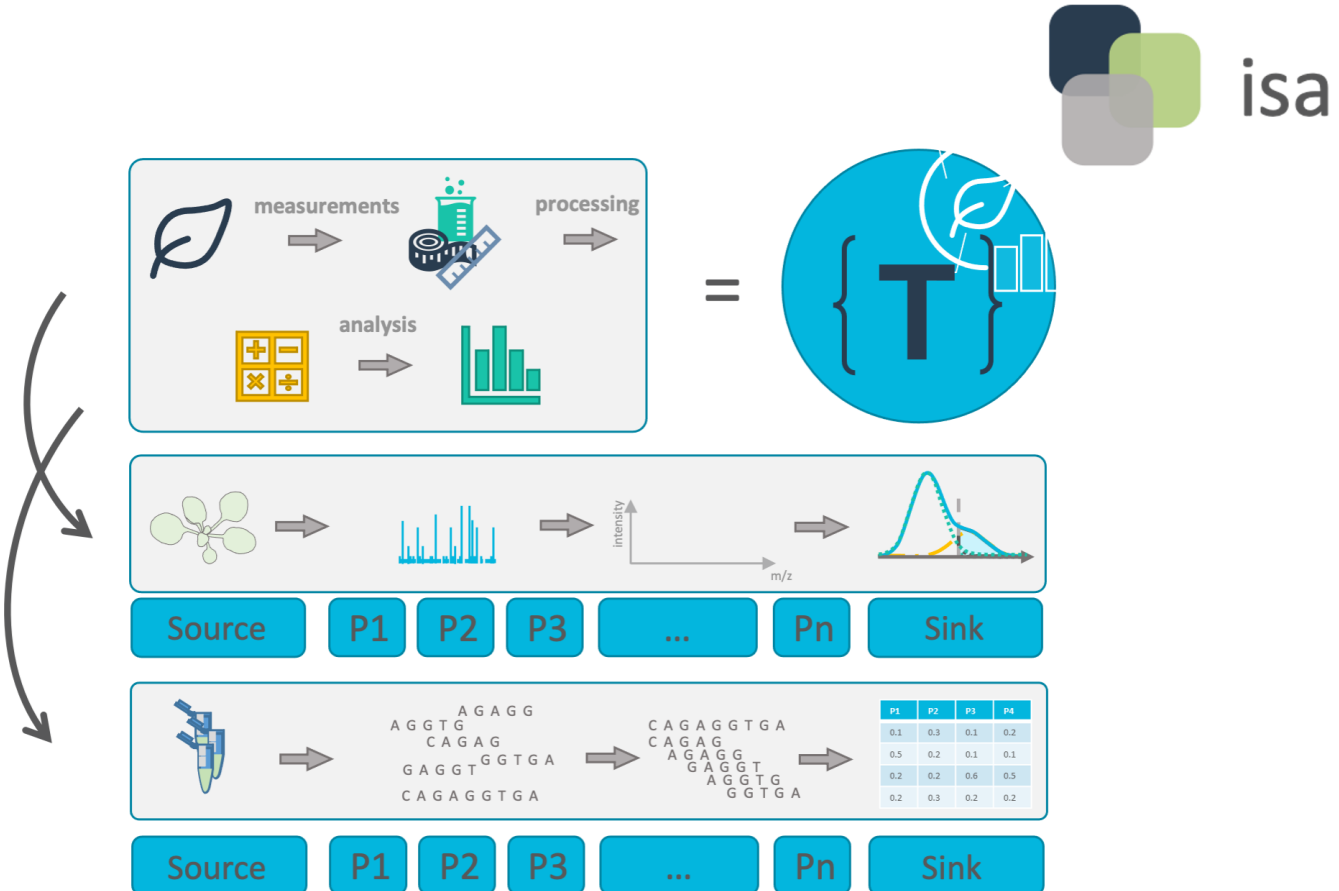
- ■ chromatography instrument model
 - ■ nanoElute2
- ■ chromatography column model
 - ■ PepSep C18 1.9u, 25cm x 75um ...



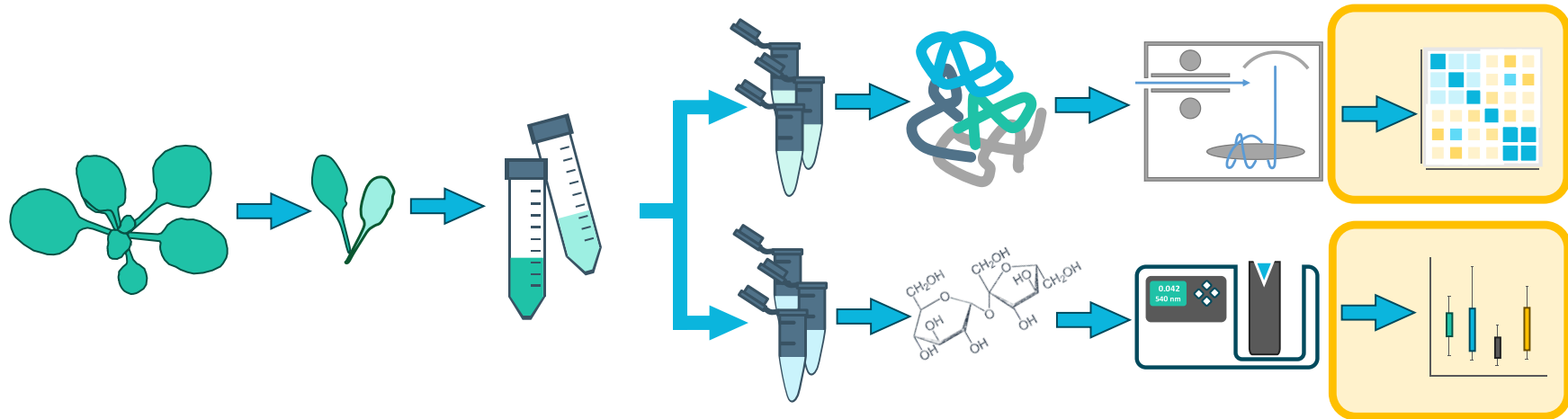
Applying standard procedures to sample record



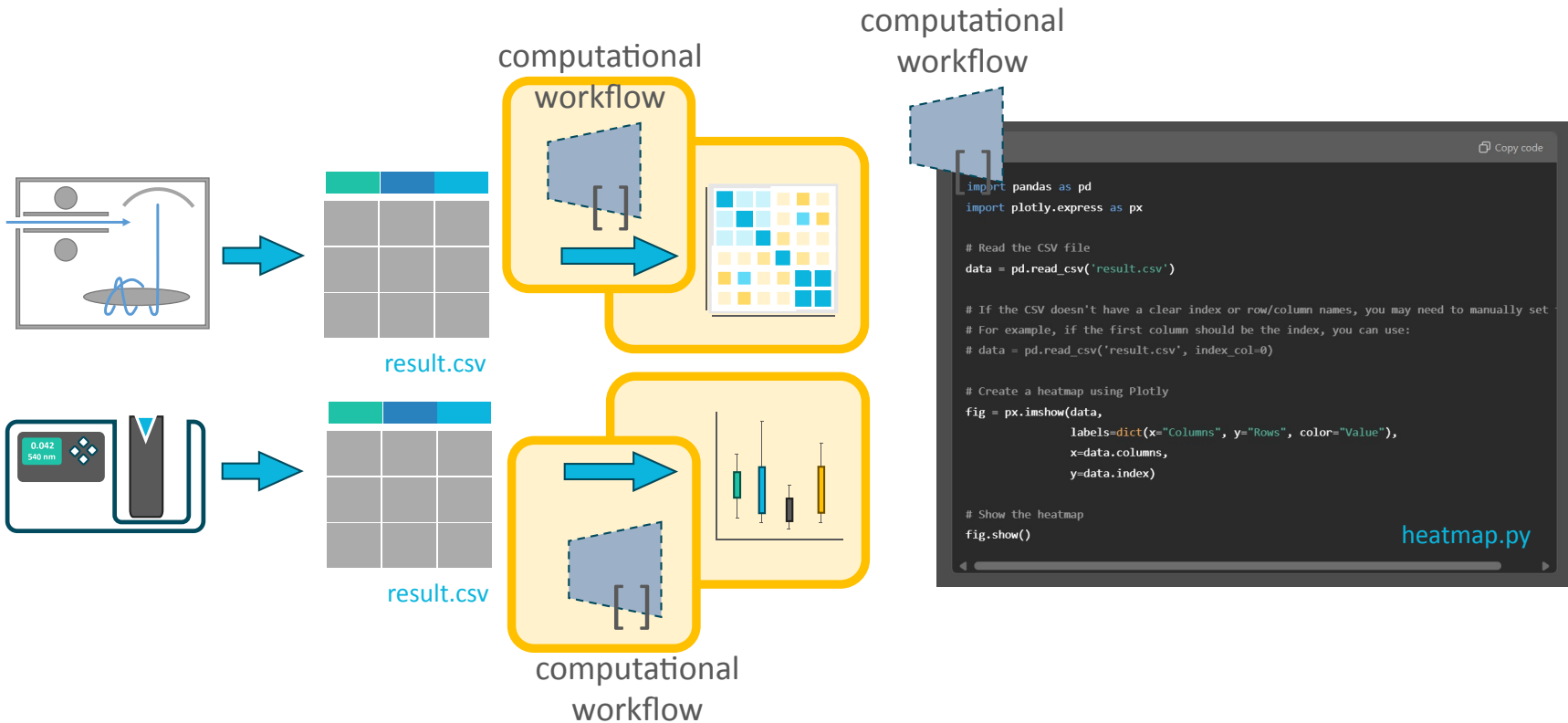
Realization of lab-specific metadata with templates



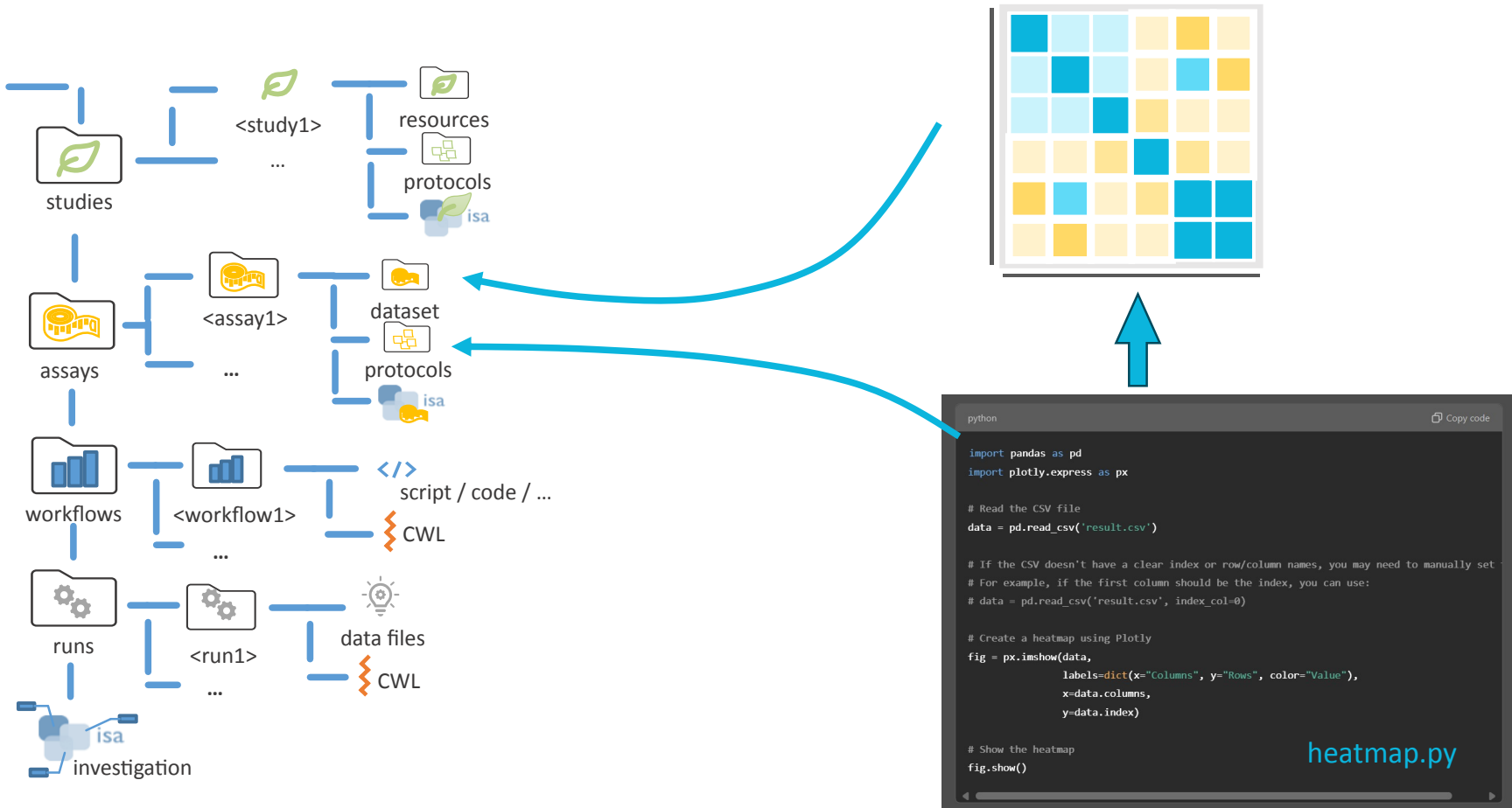
Options to annotate the data analysis



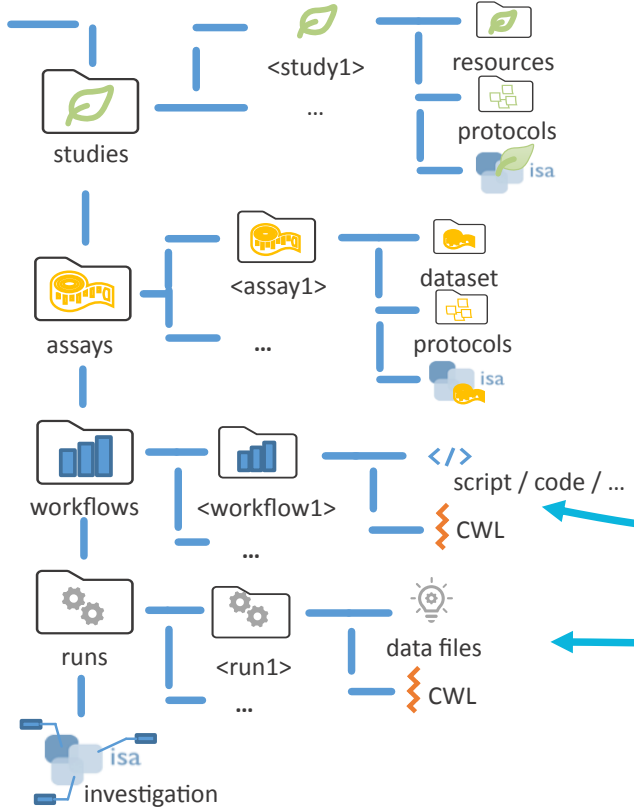
A computational workflow is like a protocol



Option I: Create a virtual assay



Option II: Create a workflow and run



```
python
Copy code

import pandas as pd
import plotly.express as px

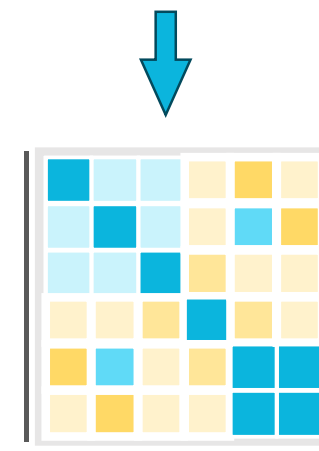
# Read the CSV file
data = pd.read_csv('result.csv')

# If the CSV doesn't have a clear index or row/column names, you may need to manually set
# For example, if the first column should be the index, you can use:
# data = pd.read_csv('result.csv', index_col=0)

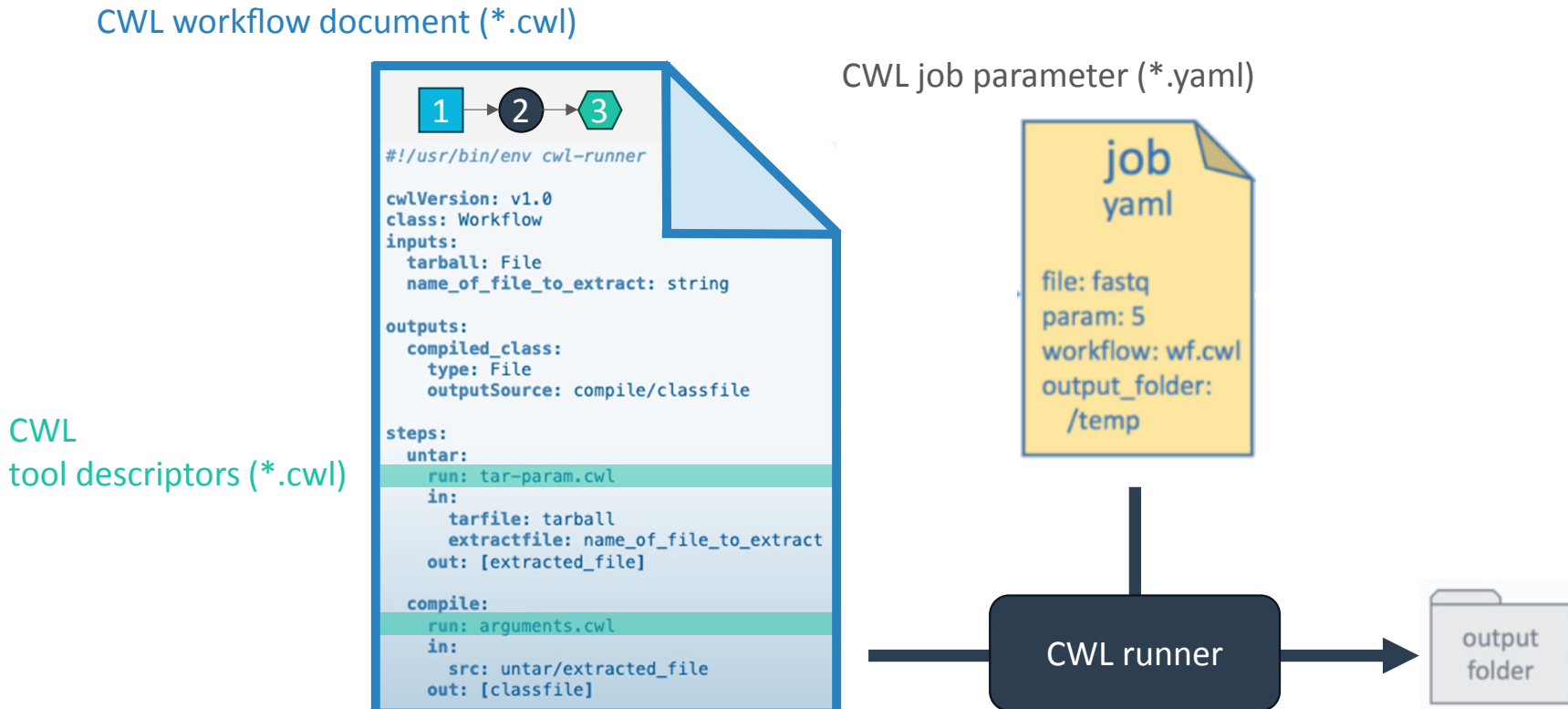
# Create a heatmap using Plotly
fig = px.imshow(data,
                 labels=dict(x="Columns", y="Rows", color="Value"),
                 x=data.columns,
                 y=data.index)

# Show the heatmap
fig.show()
```

heatmap.py



Use CWL to wrap your workflow

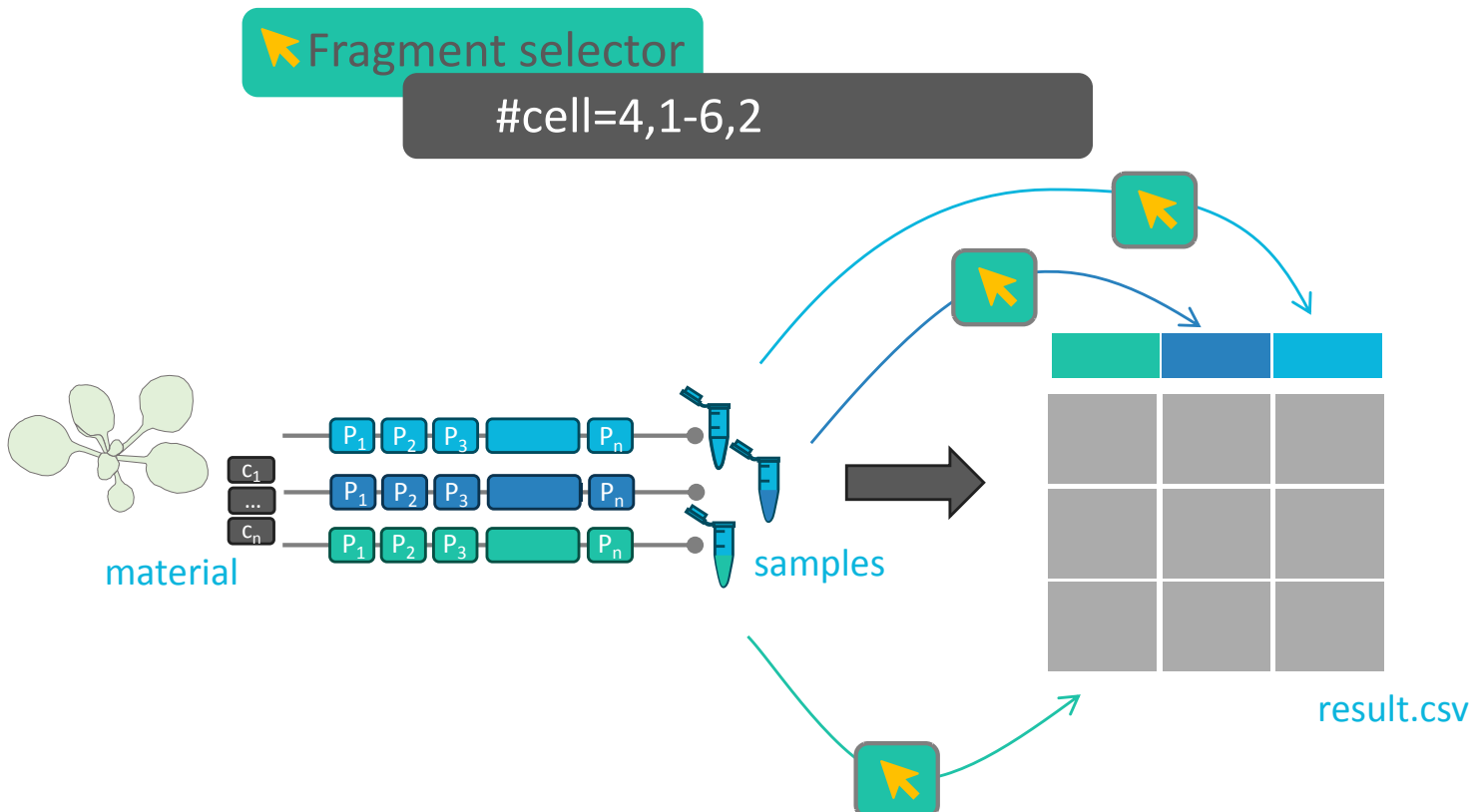


DataMAPs

Mechanisms to annotate the result data

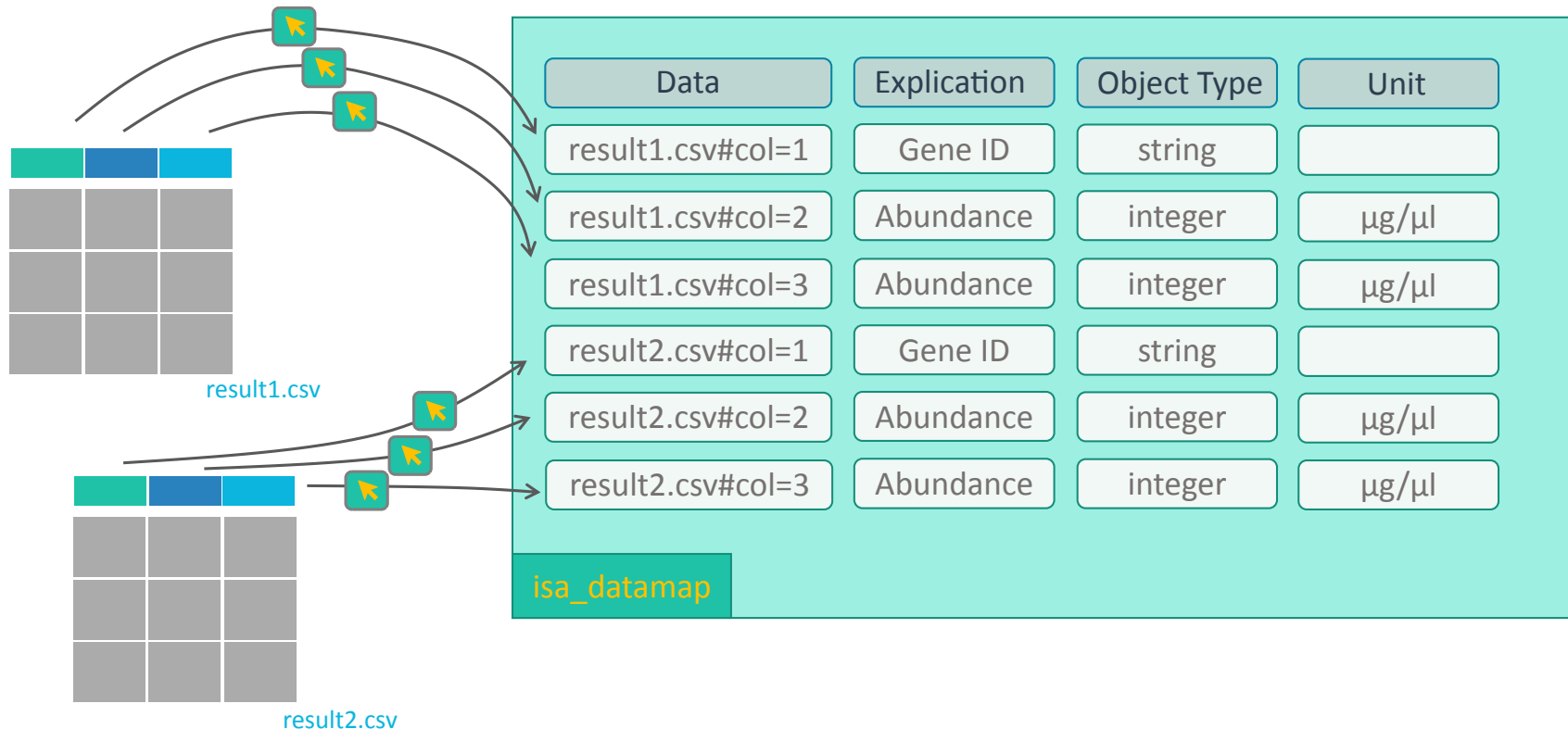
Point into results – Fragment selectors

Connecting samples to data points



DataMAP: Annotation for the fragment selectors

Annotating contents of result files



Check point 3

☒ Task

- ✓ I've added experimental details from SOPs
- ✓ In addition to the raw data, my ARC now contains the annotated data analysis
- ✓ Using a datemap I can point to the content of my result files

Domain-specific data repositories

Repository	Description	Biological data domain
EBI-ENA	European Nucleotide Archive	genome / transcriptome sequences
EBI-ArrayExpress	Archive of Functional Genomics Data	transcriptome
EBI-Metabolights	Database of Metabolomics	metabolome
EBI-PRIDE	PRoteomics IDEntifications Database	proteome
EBI-Biolimage Archive	Stores and distributes biological images	imaging, microscopy
e!DAL-PGP	Plant Genomics & Phenomics Research Data Repository	phenome
NCBI-GEO	Gene Expression Omnibus	transcriptome
NCBI-GenBank	Genetic Sequence Database	genome
NCBI-SRA	Sequence Read Archive	genome / transcriptome sequences

Domain-specific data repositories

Good

- Assign PIDs / DOIs
- Long-term accessible
- Data type specific
- Apply metadata standards
- Usually recommended / required by journals
- Mostly accepted by the community

Intermediate

- User-friendliness
- Different metadata schema
- Complex and versatile submission routines

Generic data repositories

Good

- Allow publication of any kind of data Assign PIDs / DOIs
- Long-term accessible
- Very simple to use



<https://zenodo.org>

Intermediate

- Only generic / high-level metadata schema
- Limited reusability



<https://datadryad.org/>



<https://figshare.com>

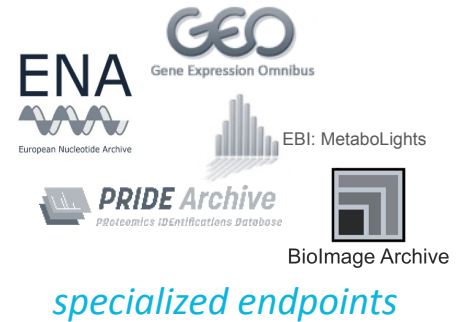
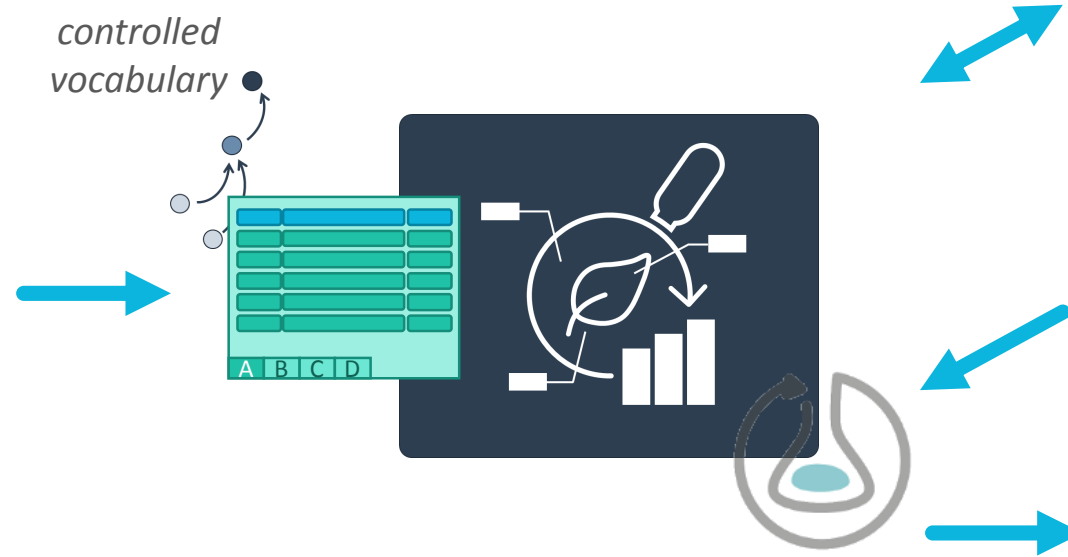
Choosing a data repository

Domain-specific >> Generic >> Institutional

Find repositories at:

- <https://www.re3data.org>
- <https://fairsharing.org>

Building exporters to support versatile publication outlets



Google
OpenAIRE
dataset search

The
Dataverse®
Project
INVENIO

data publication

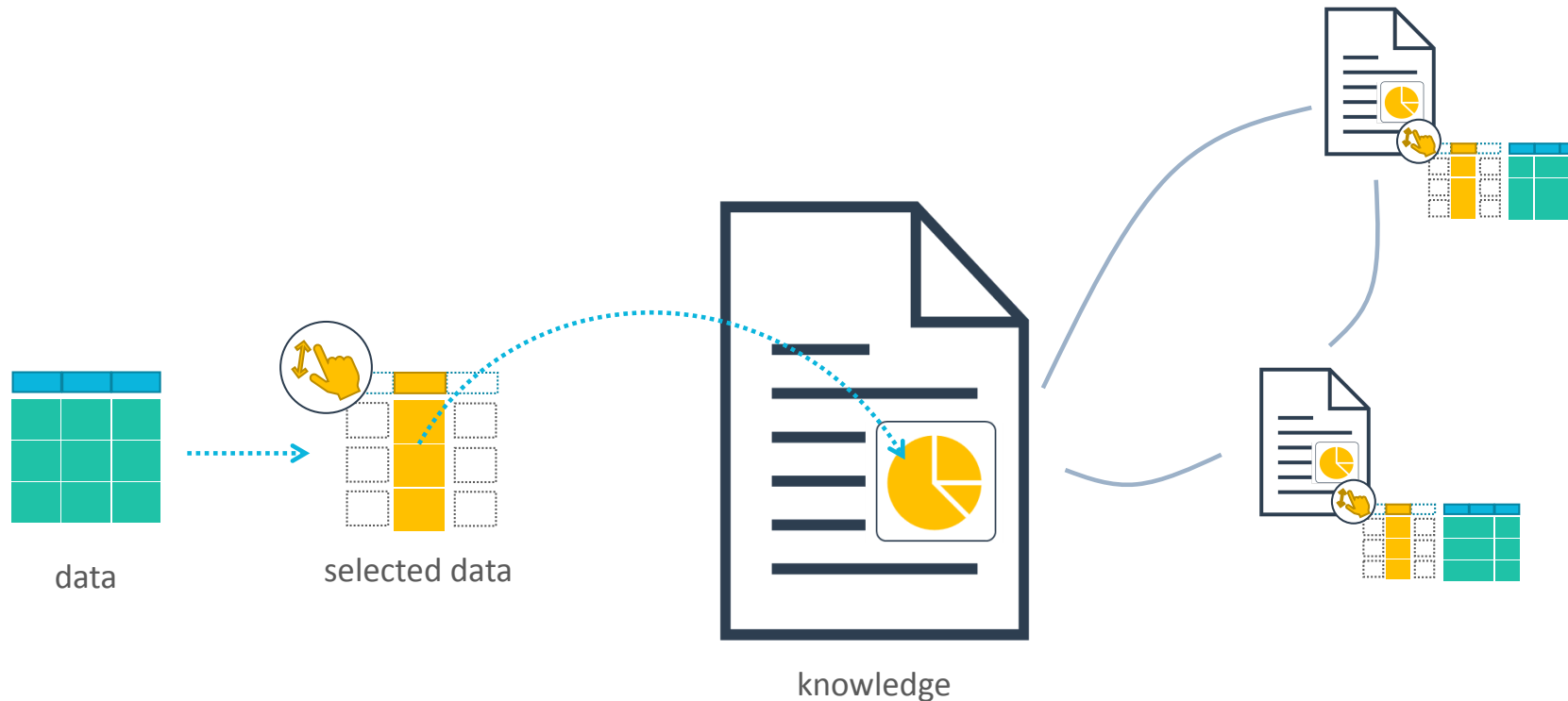
Validate and publish your ARC



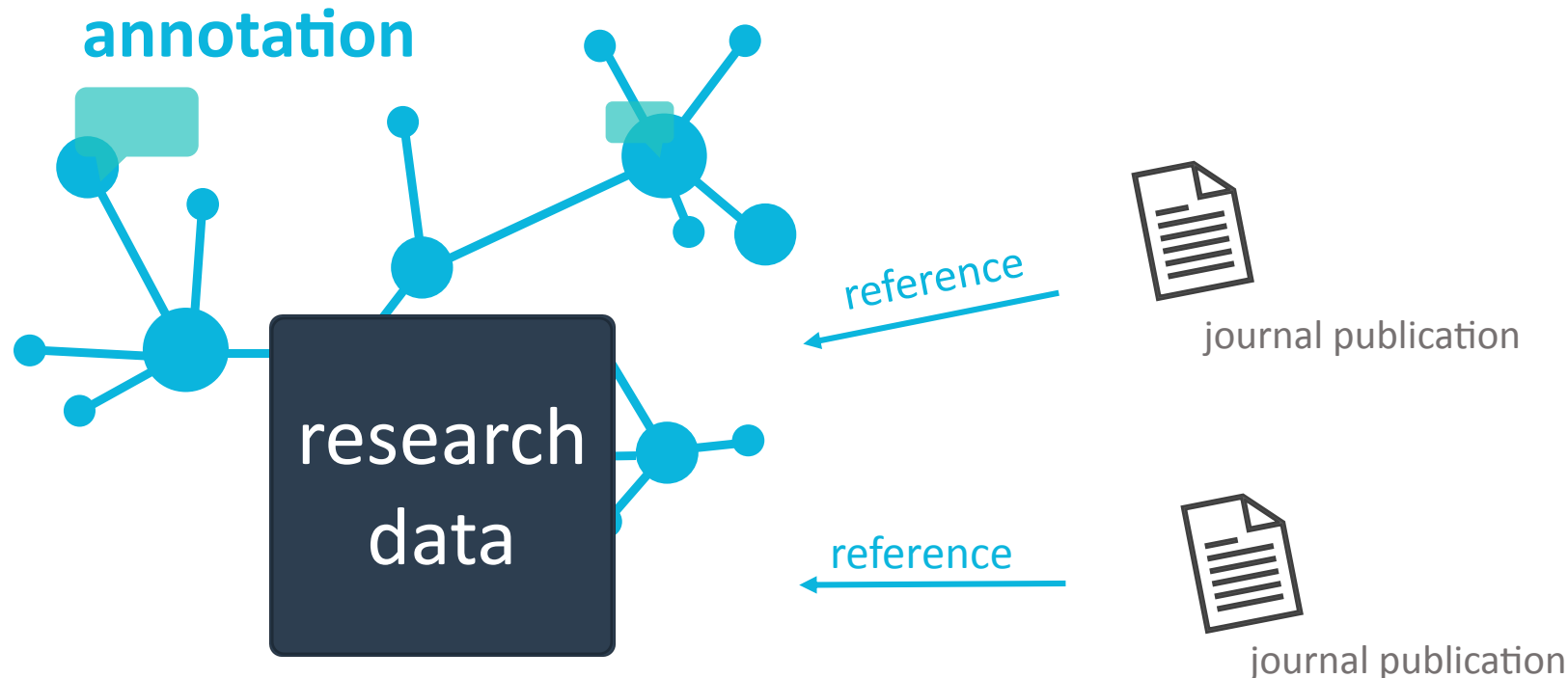
Hands-on

Follow the [Start Here](#) guide until Check point 4

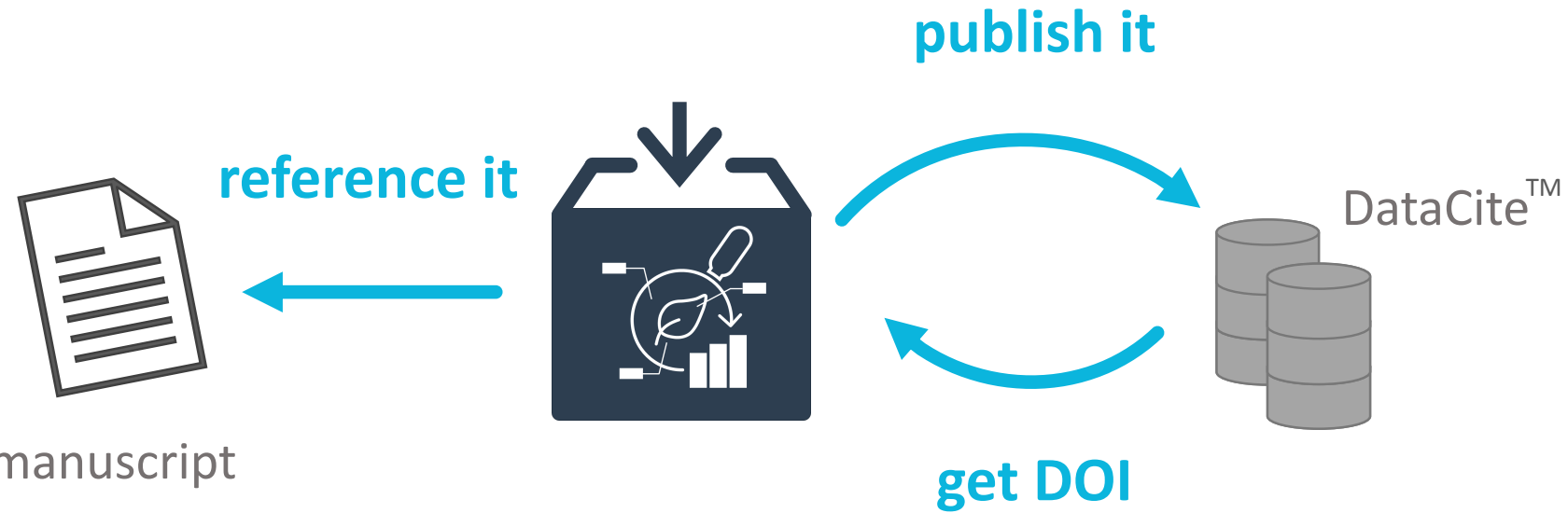
ARC data publication



Moving from paper to data publications



Publish your ARC, get a DOI



Publish your ARC with a few clicks

 **Ru_ChlamyHeatstress** 

Project ID: 122 

 Star 1

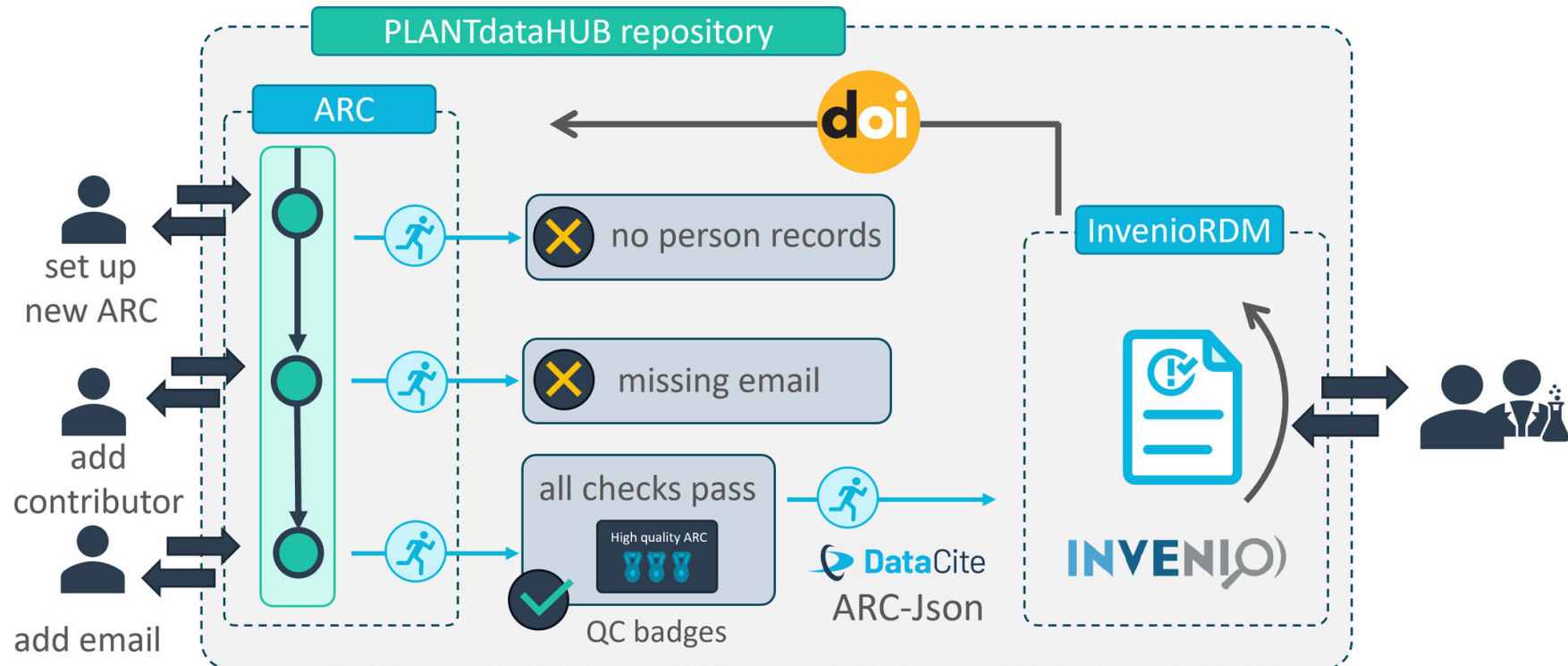
-o 53 Commits  1 Branch  0 Tags  293.9 GB Project Storage

Topics: Chlamydomonas abiotic stress proteomics + 1 more

Algae cultures were grown mixotrophically (TAP). After 24h of 35°C/40°C the cells were shifted back to room temperature for 48h. 'omics samples were taken.

1 pipeline 2 passed 3  Publish ARC arc quality 301/301

Validate & publish



Weil, H.L., Schneider, K., et al. (2023), PLANTdataHUB: a collaborative platform for continuous FAIR data sharing in plant research.
Plant J. <https://doi.org/10.1111/tpj.16474>

Validation towards publication

 **Ru_ChlamyHeatstress** 

 main  Ru_ChlamyHeatstress  Find file  History Find file Code

 add validation_packages.yml  da71d3c9 
Kevin Schneider authored 2 months ago

Name	Last commit	Last update
.arc	add validation_packages.yml	2 months ago
assays	Add missing data files	1 year ago
publication	add publication, add zScores	2 years ago
runs	add tpm data	1 year ago
studies	add publication information	1 year ago
workflows	Update	2 years ago
.gitattributes	rename missing samples; #2	1 year ago
.gitignore	add gitignore	1 year ago
README.md	Add doi badge	1 year ago
isa.investigation.xlsx	add author emails and adapt title	1 year ago

 README.md

Chlamydomonas reinhardtii heat stress time course experiment

DOI [10.60534/9e5jx-75d83](https://doi.org/10.60534/9e5jx-75d83)

 Star 2 

Project information
Algae cultures were grown mixotrophically (TAP). After 24h of 35°C/40°C the cells were shifted back to room temperature for 48h. 'omics samples were taken.

Chlamydomonas abiotic stress
proteomics + 1 more

 pipeline  invenio  10/10

-o- 55 Commits
2 Branches
0 Tags

 README
Auto DevOps enabled

Created on
July 11, 2022

Receive a DOI

Published September 7, 2023 | Version v1

Dataset 

Systems-wide investigation of responses to moderate and acute high temperatures in the green alga *Chlamydomonas reinhardtii*.

Zhang, Ningning¹; Mattoon, Erin¹; McHargue, Will¹ ; Venn, Benedikt² ; Zimmer, David² ; Pecani, Kresti³; Jeong, Jooyeon¹; Anderson, Cheyenne⁴; Chen, Chen⁴; Berry, Jeffrey¹; Xia, Ming¹; Tzeng, Shin-Cheng¹ ; Becker, Eric¹; Pazouki, Leila¹; Evans, Bradley¹; Cross, Fred³; Cheng, Jianlin⁴; Czymmek, Kirk¹ ; Schröder, Michael⁵ ; Mühlhaus, Timo² ; Zhang, Ru¹ 

Show affiliations

Style

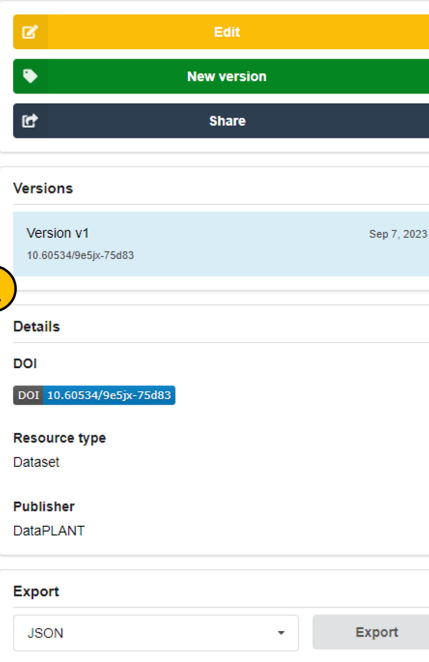
APA

1

Citation

Zhang, N., Mattoon, E., McHargue, W., Venn, B., Zimmer, D., Pecani, K., Jeong, J., Anderson, C., Chen, C., Berry, J., Xia, M., Tzeng, S.-C., Becker, E., Pazouki, L., Evans, B., Cross, F., Cheng, J., Czymmek, K., Schröder, M., ... Zhang, R. (2023). Systems-wide investigation of responses to moderate and acute high temperatures in the green alga *Chlamydomonas reinhardtii*. [Data set]. DataPLANT. <https://doi.org/10.60534/9e5jx-75d83>

2

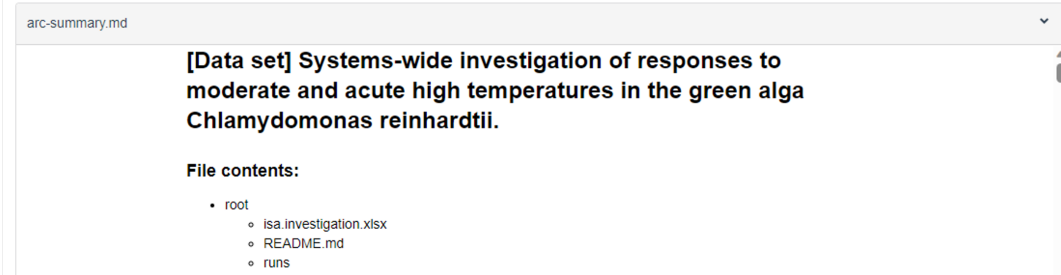


The screenshot shows the right-hand sidebar of a DataPLANT dataset page. At the top are three buttons: 'Edit' (yellow), 'New version' (green), and 'Share' (dark blue). Below that is a 'Versions' section showing 'Version v1' (DOI: 10.60534/9e5jx-75d83, Sep 7, 2023). Further down are sections for 'Details' (DOI: 10.60534/9e5jx-75d83), 'Resource type' (Dataset), 'Publisher' (DataPLANT), and 'Export' (JSON, Export button).

Description

hosted on: <https://git.nfd4plants.org/projects/122>

Files



The 'Files' section displays a single file: 'arc-summary.md'. The content of this file is shown in a scrollable box:

```
[Data set] Systems-wide investigation of responses to moderate and acute high temperatures in the green alga Chlamydomonas reinhardtii.

File contents:
• root
  • isa.investigation.xlsx
  • README.md
  • runs
```

Check point 4

Task

- ✓ My ARC is now validated and submitted for publication!

Resources



Info & materials

- DataPLANT Website: <https://nfdi4plants.org/>
- ARC website: <https://arc-rdm.org>
- Knowledge Base:
<https://nfdi4plants.org/nfdi4plants.knowledgebase/>

Tools and Services

- ARCitect: <https://github.com/nfdi4plants/arcitect>
- DataHUB: <https://git.nfdi4plants.org>

Continuous support

- HelpDesk: <https://helpdesk.nfdi4plants.org>
- Matrix for ad hoc support: <https://matrix.to/#/%23arc-user-support:matrix.org>
- User Support Meeting (2nd Friday of the month | 1 – 2pm):
<https://nfdi4plants.github.io/events/arc-user-support/>
- User Support Mailing List: [Click here to subscribe](#)

Open Source Development

- GitHub: <https://github.com/nfdi4plants>

Acknowledgements



Team Kaiserslautern

- Timo Mühlhaus
- Lukas Weil
- Kevin Frey
- Kevin Schneider
- Jonas Lukasczyk

Team Freiburg

- Dirk von Suchodeletz
- Jonathan Bauer
- Marcel Tschöpe
- Julian Weidhase

Team Jülich

- Stella Eggels
- Angela Kranz



- Björn Usadel
- Vittorio Tracanna
- Yaser Alashloo



- Sabrina Zander