# openBIS tutorial

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## Table of Contents

Overview	2
Registration of samples and protocols in the Inventory	3
Registration of new materials and samples in the Inventory	3
Registration of a media sample	3
Batch registration of chemical samples	3
Batch modification of chemical samples	4
Batch registration of storage positions for chemical samples	4
Registration of a plasmid sample with storage info	5
Registration of yeast samples with parents	6
Batch registration of buffer samples with parents	7
Deletion of duplicate Objects	7
Visualization of storage positions with the Storage Manager	8
Registration of a new protocol in the Inventory	8
Registration of a flow cytometry protocol	8
Laboratory Notebook	10
Registration of a Project	11
Registration of an Experiment	11
Registration of a first Experimental Step	12
Data upload to an Experimental step	13
Data visualization	14
Registration of a second Experimental Step	14
Data and metadata exports	15
Access rights assignment to a personal notebook or to a specific Project	15
Searching the ELN	16
Generic text search	16
Saving and re-using searches	16
Data Set Files search	17
Freezing entities	17

## Overview

In this tutorial we will learn how to use the openBIS inventory and lab notebook.

The tutorial covers the registration of samples, protocols and experiments:

- 1. Registration of media, chemicals, buffers, plasmid, yeasts
- 2. Registration of flow cytometry and western blotting protocols
- 3. Description of experiments in the lab notebook
- 4. Data upload
- 5. Searching the ELN
- 6. Freezing entities

We will see different ways of registering and updating samples and we will see how to keep track of connections to entities in the system.

Additional documentation and video tutorials can be found at: <a href="https://labnotebook.ch/">https://labnotebook.ch/</a>.

## Registration of samples and protocols in the Inventory

This part of the tutorial covers the registration of new samples and protocols in the lab inventory.

#### Registration of new materials and samples in the Inventory

We will start by registering a few samples in the Materials inventory: *media, chemicals, plasmid, yeasts, buffers*. We will see:

- 1. How to register single samples
- 2. How to batch register samples
- 3. How to batch update samples
- 4. How to assign storage positions
- 5. How to set relationships with other samples

## Registration of a media sample

We will now register one single media sample.

- 1. Select the Media Collection folder in the Samples folder
- 2. Click the + button in the main page
- 3. Enter Liquid S media in the Name field
- 4. Enter To grow cells in well controlled nutritional conditions in the For what field
- 5. Select Saccharomyces cerevisiae in the Organism field
- 6. Select room temperature from the Storage conditions list
- 7. Save

Click on the **Media Collection** folder. From the **Columns** drop down in the table you can choose which fields to visualize. Select **Name, For what, Organism, Storage Conditions**; deselect **Identifies, Code, Type**. This information is stored per user, so the selection only has to be made once.

## Batch registration of chemical samples

Now we want to register 4 chemicals. We will do this by Batch Registration.

- 1. Select the Chemicals Collection folder in the Samples folder
- 2. Select Batch Register Objects from the Operations drop down menu
- 3. Select Chemical from the Object Type dropdown menu
- 4. Download the template file and open it with Excel
- 5. Remove the **identifier** column. This is done to use identifiers automatically generated by openBIS. Keep this column if you want to manually enter identifiers (e.g. /USERNAME\_MATERIALS/SAMPLES/CHE1).
- 6. Fill in the Name, Art. Number and Supplier as below

NAME	ARTICLE_NUMBER	SUPPLIER
Dimethyl sulfoxide (DMSO)	D5879	Sigma-Aldrich
Ethanol (EtOH)	02860-2.5L	Fluka
Beta-estradiol	E8875	Sigma
Cycloheximide	C7698	Sigma

- 7. Go back to the ELN interface
- 8. Choose to upload the file you just saved
- 9. Accept
- 10. Select Name, Art. Number, Supplier and Supplier from the Columns drop down in the table.

## Batch modification of chemical samples

When we registered the chemicals before, we forgot to enter the information about the storage conditions. To correct this in all samples, we can batch update the samples:

- 1. Select the Chemicals Collection folder in the Samples folder
- 2. Select only **Identifier** and **Storage Conditions** from the **Columns** dropdown
- 3. Select Export visible columns with visible rows from the Options dropdown list
- 4. Edit this file with Excel:
  - a. enter RT under the Storage column in the first 3 rows
  - b. Enter 4 under the Storage column in the last row
  - c. Save the file
- 5. Go back to the ELN
- 6. Select Batch Update Objects from the Operations list
- 7. Select Chemical from the Object Type list
- 8. Select the previously modified file from the Documents
- 9. Accept

The storage info has now been added to the chemicals.

### Batch registration of storage positions for chemical samples

Now we want to register the storage positions for the 4 chemicals previously registered. Please follow these steps to do this in batch mode:

- 1. Select the Chemicals Collection folder in the Samples folder
- 2. Select Batch Register Objects from the Operations drop down menu
- 3. Select Storage Position from the Object Type dropdown menu
- 4. Download the template file and open it with Excel
- 5. Remove the identifier column. This is done to use identifiers automatically generated by openBIS.
- 6. Fill in the **parents** column: this field requires the identifiers of the Chemicals we registered before (e.g. /USERNAME\_MATERIALS/SAMPLES/CHEx)
- 7. Fill in the remaining information about the storage as follows:

\$STORAGE_POSITION.STORAGE_CODE	\$STORAGE_POSITION.STORAGE_RACK_ROW	\$STORAGE_POSITION.STORAGE_RACK_COLUMN	\$STORAGE_POSITION.STORAGE_BOX_NAME	\$STORAGE_POSITION.STORAGE_BOX_SIZE	\$STORAGE_POSITION.STORAGE_BOX_POSITION	\$STORAGE_POSITION.STORAGE_USER	\$XMLCOMMENTS	\$ANNOTATIONS_STATE
BENCH	1	1						
BENCH	1	1						
BENCH	1	1						
FRIDGE1_ROOM_A1.1	1	1	Box_your_usernam e	4x4	A1,A2			

- 11. Go back to the ELN interface
- 12. Choose to upload the file you just saved, in Batch Register Objects -> Storage Position
- 13. Accept
- 14. Select Name and Storage from the Columns drop down in the table.

To visualize the samples in the respective storages:

- 1. Go to Storage Manager under Utilities
- 2. Select **Bench** or **Fridge 1 Room A1.1** from the **Storage** dropdown

## Registration of a plasmid sample with storage info

- 1. Go to the **Plasmid Collection** folder in the main menu
- 2. Click the + button
- 3. Enter insul-(lexA-box)4-PminCYC1-CitrineA206K-TCYC1 in the Name field
- 4. Select pRS30y from the Backbone list
- 5. Select bla from the Bacterial antibiotic resistance list
- 6. Select URA 3 from the Marker list
- 7. Enter Notl, Kpnl in the Flanking Restriction Enzymes field
- 8. Define a storage position:
  - a. Click the + button under the storage table
  - b. Select the -80°C Room A1.2 freezer
  - c. Select the 1,1 rack
  - d. Enter Username\_Plasmid\_box in the Box Name field

- e. Select the 10 x 10 size
- f. Select position A1
- g. Accept
- 9. Save

## Registration of yeast samples with parents

First we will register one yeast:

- 1. Go to the Yeasts Collection folder in the main menu
- 2. Enter LexA-DBD-long-HBD-B42 in the Name field
- 3. Select BY4741 from the Genetic Background list
- 4. Select a from the Mating Type list
- 5. Select met15- from the Background-specific markers list
- 6. Select ura3- leu2- from the Common markers list
- 7. Select cir+ from the Endogenous 2micron plasmid in yeast list
- 8. Select transformation from the Origin list
- 9. Select PCR from the Strain Check list
- 10. Define a storage position:
  - a. Click the + button under the storage table
  - b. Select the -80°C Room A1.2 freezer
  - c. Select the 2,1 rack
  - d. Enter Username\_Yeast\_box in the Box Name field
  - e. Select the 10 x 10 size
  - f. Select position A1
  - g. Accept

#### 11. Save

Now we will register a second yeast, which was made using the first yeast and the previously registered plasmid. These can be set as **parents.** 

We can **copy** the yeast we just register and modify only certain fields that are different:

- 1. Go to the Yeasts Collection folder in the main menu
- 2. Click on the existing yeast sample
- 3. Click on the copy button from the menu toolbar
- 4. Leave everything unselected
- 5. Accept
- 6. The new yeast is created. Edit the form.
- 7. Remove LexA-DBD-long-HBD-B42 from the Name field and enter demo.
- 8. Add relationships:
  - a. Add yeast parent:
    - i. In the parents section, click on the + next to Yeast parents
    - ii. Select the yeast previously registered from the table
  - b. Add plasmid parent:
    - i. Click the + next to Plasmid
    - ii. Select the only plasmid in the table
    - iii. Select integration from the Plasmid Relationship dropdown
    - iv. Enter URA3 in the Plasmid annotation field
- 9. Change value to leu2- for the Common markers field

- 10. Define a storage position:
  - a. Click the + button under the storage table
  - b. Select the -80C Room A1.2 freezer
  - c. Select the Username\_Yeast\_box in the 2,1 rack
  - d. Select position A2
- 11. Accept
- 12. **Save**





## Batch registration of buffer samples with parents

We will now register 3 buffers using batch registration. Each of the buffers has 2 chemicals as parents, which we need to set in the file.

The first buffer, cycloheximide, has DMSO and cycloheximide as chemical parents. The second and third buffers, bet-estradiol, have Ethanol and beta-estradiol as chemical parents.

#### Setting parents in file:

- 1. Open the **buffers.tsv** file with Excel to visualize its content
- 2. Go to the Chemicals Collection
- 3. Select Identifier and Name from the table
- 4. In the **buffers.tsv** file, fill in the **parents** column with the **Identifiers** corresponding to the chemicals mentioned above:
  - i. DMSO and cycloheximide for the first buffer
  - ii. EtOH and beta-estradiol for the second and third buffer

#### **Batch upload**

- 5. Go to the **Buffers Collection** folder in **Samples** in the main menu
- 6. Select Batch Register Objects from the Operations drop down menu
- 7. Select Solution Buffer from the Object Type list
- 8. Select the buffers.tsv file just modified
- 9. Accept

Select Name, Parents, Details, For what, Stock Concentration and Storage Conditions from the Columns dropdown.

Click on one of the entities in the table (e.g. cycloheximide) and visualize the connections to the chemicals with the hierarchy tree button

## Deletion of duplicate Objects

In the step above we registered 3 buffers, but two of them are the same. We want to delete one of them:

- 1. Go to the **Solutions Buffers Collection** folder in **Samples** in the main menu
- 2. Select *Object* **beta-estradiol** from the table (select button in first column)
- 3. Select **Delete selected** from the **Options** drop down in the table
- 4. Enter duplicate in the Reason field
- 5. Accept

When *Objects* (and also *Experiments/Collections*) are deleted they are first moved to the trashcan, In order to be completely removed from the database they have to be deleted also from the trashcan. Alternatively, it is also possible to revert deletion from the trashcan, if something was accidentally moved here.

- 1. Go to the Trashcan under Utilities in the main menu
- 2. Select **Delete Permanently** from the **Operations** drop down in the table
- 3. Read the warning message
- 4. Accept

## Visualization of storage positions with the Storage Manager

The openBIS storage manager offers an overview of all storages configured for the lab:

- 1. Select Storage Manager under Utilities
- 2. Select the -80°C room A1.2 storage from the Storage list
- 3. Click on one of the boxes and visualize the content

It is possible to drag & drop boxes to change the position of a box inside a storage (or to a different storage). In the same way the position of a sample inside a box can be changed (this is not supported for multiple positions for the same sample).

#### Registration of a new protocol in the Inventory

We will now register a flow cytometry protocol in the Methods Inventory.

## Registration of a flow cytometry protocol

- 1. Open the flow-cytometry-protocol.docx file from the training material
- 2. Go to the ELN
- 3. Go to the Flow cytometry protocols folder in the main menu
- 4. Click on the + button in the main page
- 5. Copy/paste the corresponding fields from the doc file in the ELN form
- 6. To add the chemical as parent:
  - a. Click the + button next to Links to Materials and Methods
  - b. Select Chemical from the drop down
  - c. Select Name from the Columns drop down
  - d. Select cycloheximide from the table
- 7. Save

## Laboratory Notebook

In the laboratory notebook, usually each lab member has a personal folder (=Space) to organize Projects and Experiments. Experiments can be further divided in Experimental Steps. An openBIS Experiment is a specific scientific question and it contains Experimental Steps that are individual attempts to answer it.

Experimental Steps can be linked to each other (if needed) and they can also be linked to materials and methods stored in the Inventory.

Some examples:

Lab Experiment	Corresponding openBIS data model				
Monitor decondensation upon transcriptional	Experiment: Monitor decondensation upon				
activation of a given insert in some strains.	transcriptional activation of a given insert in some				
	strains.				
This experiment involves 3 different steps:	Exp Step 1: make plasmids				
Make some plasmids	Exp Step 2: make reporter strains for				
2. Make reporter strains for decondensation	decondensation using plasmids made in Exp step				
using plasmids made before-> PCR	1 (can be a child of Exp Step 1)				
results	Exp Step 3: test strains made in Exp Step 2 using				
3. Test strains made before in different	condition 1 (can be a child of Exp Step 2)				
conditions -> raw data, MATLAB code	Exp Step 4: test strains made in Exp Step 2 using				
	condition 2 (can be a child of Exp Step 2)				
	Exp Step 5: test strains made in Exp Step 2 using				
	condition 3 (can be a child of Exp Step 2)				
	Each Experimental Step can contain the data				
	related to it.				
Induction of a transcription factor in standard	<b>Experiment</b> : Induction of a transcription factor in				
growth conditions with synthetic complete	standard growth conditions with synthetic				
medium containing 2% of glucose.	complete medium containing 2% of glucose.				
	Exp Step 1: flow cytometry				
This experiment involves 3 different steps:	Exp step 2: western blotting 1				
Detection of transcription factor	Exp Step 2: western blotting 2				
induction by flow cytometry					
2. Detection of transcription factor	In this case we are trying different methods to				
induction by Western blotting	detect the transcription factor, but there is no				
3. Detection of transcription factor	connection between them.				
induction by Western blotting					
	Each Experimental Step can contain the data				
	related to it.				
Monitor the expression of a given gene.	<b>Experiment</b> : Monitor the expression of a given				
	gene				
This experiment involves 2 steps:	Exp Step 1: RT-qPCR				
1. RT-qPCR	Exp Step 2: Western blotting				
2. Western blotting	Each Experimental Step can contain the data				
	related to it.				

As an example, we work on a project where we study different inducible transcription factors. In this project we perform several experiments. For example, in one experiment we analyze the abundance of 4 variants of a transcription factor before and after induction. The analysis is done by western blotting.

In another experiment we induce the transcription factor in standard growth conditions and we detect it by flow cytometry and 2 types of western blotting.

In openBIS we would need to register:

- 1. 1 Project
- 2. 2 Experiments
- 3. 1 Experimental Step for the first experiment; 3 Experimental Steps for the second experiment.

In this tutorial we will work with **Example 2** described above and we will register 1 *Project*, 1 *Experiment* and 2 *Experimental Steps* (1 flow cytometry and 1 western blotting). In the Experimental Steps, we will create links to samples and protocols stored in the inventory.

## Registration of a Project

We want to register a project called **Inducible transcription factor**:

- 1. Select your folder in the Lab Notebook part of the main menu
- 2. Click the + button in the main page
- Enter Construction and characterization of a beta-estradiol-inducible transcription factor for Saccharomyces cerevisiae in the Description field
- 2. Enter Inducible\_transcription\_factor in the Code field
- 3. Save

Please note that Projects only have Codes, not names. The Code can only have alphanumeric characters and no spaces. It is recommendable to use "\_" to separate words. The ELN will translate this to a space in the main menu.

### Registration of an Experiment

In the Project registered before, we want to register a first Experiment, called **Induction of a transcription** factor in standard growth conditions with synthetic complete medium containing 2% of glucose:

- 1. Select the Project folder
- 2. Click the + button in the main page
- 3. Enter Induction of a transcription factor in standard growth conditions with synthetic complete medium containing 2% of glucose in the Name field
- 4. Check **Show in project overview.** This is a way to mark important experiment, so that they are shown in the Project page. Usually this would be done at the end of the Experiment.
- 5. Enter 10.01.2019 in the start date
- 6. Enter 20.04.2019 in the end date
- 7. Enter Analyze the transcription factor in a concentration series of inducers in the Experimental goals field
- 8. Enter Both variants of transcription factor tested induced in a concentration series of inducer. The variant LexA-ER-B112 is stronger than LexA-ER-B42. in the Experimental results field.
- 9. Save

## Registration of a first Experimental Step

Now we want to register the first *Experimental Step*, which is a flow cytometry experiment. We also want to create links to samples and protocols stored in the inventory.

- 1. Click the + button in the Experiment page
- 2. Enter Detection of LexA-ER-B42 induction by flow cytometry in the Name field
- 3. Click Show in project overview

Now we want to create links to materials and methods stored in the Inventory. We want to add a link to 1 media, 2 buffers, 1 yeast and 1 protocol. These are the samples and materials we used in this experimental step and the protocol we followed.

#### 4. Add media:

- a. Click the + button next to Links to materials and methods
- b. Select Media from the list
- c. Show the Media Name in the table (Columns dropdown)
- d. Select the only media available in the table

#### 5. Add buffers:

- a. Click the + button next to Links to materials and methods
- b. Select **Solutions Buffer** from the list
- c. Show the Names of the buffers in the table (Columns dropdown)
- d. Check both buffers in the table
- e. Select Add selected from the Options dropdown in the table

### 6. Add yeast:

- a. Click the + button next to Links to materials and methods
- b. Select Yeast from the list
- c. Show the Names of the yeasts in the table (Columns dropdown)
- d. Select the **demo** yeast

#### 7. Add protocol:

When we add a protocol we have two options:

- i. Create a link to an existing protocol in the inventory. This is suitable when the protocol is followed "as it is"
- ii. Copy the protocol to our experiment folder in the lab notebook in order to modify it. This is suitable when the main protocol is modified during the experiment.

We will now see the procedure to create a local copy:

- b. Click + next to **Protocols**
- c. Show the **Name** of the protocols in the table (Columns dropdown)
- d. Select the flow cytometry protocol
- e. Choose Use as template from the Operation dropdown in the table
- f. Enter a code in the Code field
- g. Accept

- h. Right click on the Protocol code and open in new tab. This is a copy of the original protocol, created in your personal folder (check the identifier). You can see that this protocol is linked to the original protocol in the inventory. In this way you can modify your local copy and leave the template untouched. If you do not need to make any modification to the protocol, you do not need to use the **Use as template** option.
- 8. Enter Machine used: LSRII Fortessa in the Experimental description field
- 9. Save

## Data upload to an Experimental step

Data can be uploaded to *Experiments* or *Experimental Steps* using the same procedure.

In the same way, files can also be added to *Objects* in the Inventory (i.e. to samples or protocols).

In this case we want to upload: 1. the raw data that was obtained from the flow cytometer; 2. the R script that was used to analyze the data; 3. the final result pictures.

#### **Upload** raw data

- 1. Click the Upload Dataset icon form the menu toolbar
- 2. Select Raw data from the Data Set type dropdown
- 3. Enter Flow cytometry data in the Name field
- 4. Select the FC\_LEXA-ER-B42-raw.zip file in the Documents folder to upload
- 5. Check Uncompress before import
- 6. Save
- 7. Open the Raw data folder in the Experimental step to see the content

#### **Upload the R script**

- 1. Click the Upload dataset icon form the menu toolbar
- 2. Select Code from the Data Set type dropdown
- 3. Type flow in the Parent field and select the dataset that comes up in the list. This step is not necessary, but it allows you to establish a relationship between datasets. In this case we establish a connection to the raw data that was analyzed with the R script we are uploading.
- 4. Enter R script in the Name field
- 5. Select the FC LEXA-ER-B42-script.R file in the Documents folder to upload
- Save
- 7. Open the **Code** folder in the Experimental step to see the content

#### Upload the analyzed data

- 1. Click the Upload dataset icon form the menu toolbar
- 2. Select Analysed data from the Data Set type dropdown
- 3. Type script in the Parent field and select the dataset that comes up in the list.
- 4. Enter Analysis results in the Name field
- 5. Select the FC\_LEXA-analyzed\_data.zip file in the Documents folder to upload
- 6. Check Uncompress before import
- 7. Save
- 8. Open the Analyzed data folder in the Experimental step to see the content

### Data visualization

To open data files stored in openBIS we recommend to use software that allows to mount openBIS as a drive on your computer. Examples are:

- 1. Mountain Duck for MacOS (https://mountainduck.io)
- 2. NetDrive for Windows (https://www.nsoftware.com/sftp/netdrive/).

Files can be opened with the desired application in read-only mode.

Any other FTP solution can also be used (e.g. Cyberduck, Filezilla, etc).

Please note that data files stored in openBIS are read-only!

## Registration of a second Experimental Step

In our main experiment, called **Induction of a transcription factor in standard growth conditions with synthetic complete medium containing 2% of glucose,** we performed a flow cytometry experiment and two western blotting experiments. We now want to register one western blotting experiment as *Experimental Step* in openBIS.

- 1. Select the Experiment Induction of a transcription factor in standard growth conditions with synthetic complete medium containing 2% of glucose from the Lab Notebook main menu
- 2. Click the + button in the main page
- 3. Enter Detection of LexA-ER-B42 induction by western blotting in the Name field
- 4. Select Show in project overview

Now we want to create links to some materials stored in the Inventory. We want to add a link to 1 media, 2 buffers, 2 yeasts. These are the samples and materials we used in this experimental step.

- 5. Add media:
  - a. Click the + button next to Links to materials and methods
  - b. Select **Media** from the list
  - c. Show the Media Name in the table (Columns dropdown)
  - d. Select the only media available in the table
- 6. Add buffer:
  - a. Click the + button next to Links to materials and methods
  - b. Select Solution Buffer from the list
  - c. Select the **beta-estradiol** buffer from the table
- 7. Add yeast:
  - a. Click the + button next to Links to materials and methods
  - b. Select Yeast from the list
  - c. Select the **demo** yeast from the table
- 8. Enter Analyze the full induction of LexA-ER-B42 on western blot by doing a dilution series in the Experimental goals field.
- 9. Enter LexA-ER-B42 full fold induction is between 128 and 256 in the Experimental results field
- 10. Add a gel picture to the Experimental results field:
  - a. Select the image icon in the Experimental results text editor
  - b. Go to the **Upload** tab

- c. Choose the WB\_LEXA-ER-B42-actine.png file you received from us
- d. Click Send it to server
- e. Resize the width to 500
- f. Press OK

#### 11. Save

## Data and metadata exports

It is possible to export a complete lab notebook or only parts of it (*Project, Experiment, Experimental Step, Datasets*).

In each folder, the menu toolbar on the main form has an option to **Export metadata only** or **Export metadata anly** or **Export metadata anly** or **Export metadata anly** or **Export metadata and anly** or **Export metadata and an analysis** of **Export metadata and an analysis** or **Export metadata analysis** or **Export metadata and an analysis** or **Export metadata analysis** or **Export metadata analysis** 

When you export something, you will receive an email with a link to download your metadata (+data).

The export contains folders with the same structure organization they have in openBIS. Metadata is exported to 4 different file formats: .docx, .html, .txt, .json.

## Access rights assignment to a personal notebook or to a specific Project

It is possible to grant collaborators and colleagues access rights to your complete lab notebook or only to selected *Project*(s).

To grant rights to your folder:

- 1. Select your folder in the Lab Notebook part of the main menu
- 2. Click the **Manage access** icon in the main page
- 3. Select a role from the **Role** dropdown menu. Available roles:
  - a. Observer: has read-only access
  - b. User: can create and modify entities, but cannot delete anything
  - c. Admin: can create, modify and delete entities
- 4. Select **User** from the **grant to:** dropdown
- 5. Enter the username of a registered user
- 6. Grant access

## Searching the ELN

openBIS offers 3 options for searching:

- 1. **Text search** on all fields stored in the database. Searches can be refined using the **Advanced search**.
- 2. **BLAST** search for sequence comparison across sequences stored in the database.
- 3. Data Set Files search. This should be used to find files uploaded as datasets by their name.

#### Create a folder to save queries in your lab notebook

It is now possible to save and reuse searches in the ELN. Searches can only be saved in a *Space* where the user has admin rights. Usually, each user has admin rights in their own personal lab notebook folder.

#### Create a collection folder for searches in your lab notebook folder:

- 1. Go to your **lab notebook** folder
- 2. Click the + button
- 3. Enter OUERIES in the Code field
- 4. Save
- 5. In the Queries form, select Create Experiment from the Operations dropdown
- 6. Choose Collection from the list
- 7. Enter Queries in the Name field
- 8. Save

## Generic text search

We want to find all the flow cytometry Experimental Steps registered by you.

- 1. Enter flow cytometry in the Global search field, on top of the main menu. This search returns a few results. Now we want to narrow down the search.
- 2. Select Object in the Search For dropdown
- 3. Click the + button in the Criteria table
- 4. Select Property from the Field Type dropdown
- 5. Select Object Type from the Field Name dropdown
- 6. Enter \*STEP\* in the Field Value field
- 7. Click again the + button in the Criteria table
- 8. Select Property from the Field Type dropdown
- 9. Select Registrator from the Field Name dropdown
- 10. Enter your username in the Field Value field
- 11. Run the search

In this case, the search returns only one item, but in a similar way, you can run complex searches by combining multiple search criteria.

### Saving and re-using searches

To save a search in your Project folder:

- 1. Select **Save** on top of the page
- 2. Enter demo search in the Name field

- 3. Enter queries in the search entity to store query field
- 4. Save

#### Run a saved search:

- 1. Refresh the page to clear the search
- 2. Select the search from the list of saved searches
- 3. Run the search

### Data Set Files search

We want to find a file that contains "script" in the name.

- 1. Enter script in the Global search field, on top on the main menu
- 2. Select Data Set File from the dropdown next to it

The search returns the dataset that contains the R file script we uploaded to out flow cytometry experiment before.

## Freezing entities

Since the last upgrade (openBIS v19.06.1), openBIS has a new feature that allows to freeze entities that should no longer be modified. When freezing one entity, everything connected to it can be frozen too. The user needs to select what should be frozen and what shouldn't.

Please note that freezing is IRREVERSIBLE!



At all levels of the notebook and inventory, there is a **lock** icon.

When you click the lock, you are presented with a list of things connected to the chosen level that can also be frozen. By default, everything is selected. After the selection, you need to enter your password to be able to freeze entities.