SyRF User Guide

CAMARADES

23 Apr 2021

# Contents

1	Get	ting Started	5
	1.1	Glossary	5
<b>2</b>	Cno	eate an Account	9
4	Cre	ate an Account	9
3	Joii	a a Project	11
4 Create a New Project			
	4.1	My project will have multiple screening stages	13
5	Upl	loading a Systematic Search	15
	5.1	Deduplicating your systematic search	15
	5.2	Uploading files	15
	5.3	Uploading full-text PDFs	17
	5.4	View project studies	17
	5.5	Deleting systematic searches	17
6	$\mathbf{Pro}$	ject Stages	19
	6.1	What are stages?	19
	6.2	Screening stages	19
	6.3	Annotation stages	19
	6.4	Screening and annotating within the same stage	20
7	Scr	eening	21
	7.1	Screening using SyRF	21
	7 2	Number of screeners	22

8	Ann	notation Questions	<b>23</b>	
	8.1	Creating annotation questions	23	
	8.2	Question categories	23	
	8.3	Creating a new question	24	
	8.4	Adding questions to stages	25	
	8.5	Question categories	25	
	8.6	Nesting questions	27	
	8.7	Question editing	27	
0	DA C		29	
9	$9  ext{ FAQ}$			

# Getting Started

Welcome to the SyRF User Guide. This guide is designed to help you use SyRF. If you require general guidance on systematic review or meta-analysis, or have never conducted a systematic review before, please read the Systematic Review Wiki created by CAMARADES Berlin.

### 1.1 Glossary

### 1.1.1 Project

Each project is specific to your systematic review and meta-analysis. Upload your deduplicated studies and add stages to screen and annotate your data.

### 1.1.1.1 Public project

A project that can be seen by anyone with a SyRF account. Other users can request to join a public project with your approval.

### 1.1.1.2 Private project

A project that is not visible to other SyRF users unless they have requested and have been granted permission to join the project.

### 1.1.2 Protocol

A structured description of what you set out to do in your systematic review and meta-analysis, which should be finalised before you start your systematic review.

We recommend that this is published or shared publicly with PROSPERO.

### 1.1.3 Screening

Screening refers to making a decision whether to include or exclude a study retrieved in your systematic search based on the inclusion/exclusion criteria defined in your protocol. Screening is described in more detail in

SECTION 7.

### 1.1.4 Annotation

You may want to annotate your studies by labelling or extracting relevant information from them. This part of a systematic review project is fairly flexible and therefore you can define your own annotation questions. Annotation questions should address all questions you want to ask as specified in your protocol. You can choose at which stage of the project you want to answer specific annotation questions. This is described in more detail in

SECTION 8.

### 1.1.5 Data extraction

Where you extract data from graphs or tables in the form of means/medians and corresponding error.

### 1.1.6 Experiment

An experiment refers to any grouping of cohorts where an experiment is carried out at the same time and any of them can be compared with each other.

### 1.1.7 Cohort

A cohort refers to a group of animals - same species, strain, source, comorbidities (if applicable) - which all receive the same procedure and treatments and can be compared to other cohorts. For example, an experiment may involve the following cohorts:

Example Experiment 1:

• Cohort 1: Treatment

• Cohort 2: Sham

• Cohort 3: Control

1.1. GLOSSARY 7

### Example Experiment 2:

• Cohort 1: Control

• Cohort 2: Treatment 1

• Cohort 3: Treatment 2

Cohorts are created in SyRF projects by combining disease models and treatment details. This is described further in

SECTION 8.

<< ADD FLOWCHART DIAGRAM >>

## Create an Account

You will need to create a free account to create or participate in projects.

<< Add link to new SyRF to create an account >>

Creating an account allows us to keep your data secure and allows the administrator of each project to control who has access to the project data. Read our Data Management and Sharing Policy here.

<< Link to FAQs >> I don't see an email in my inbox from SyRF

Once you have created an account and logged in you can access public projects or create a new project via the Projects tab in SyRF.

# Join a Project

The Projects tab in SyRF will show all the projects you are a member of, as well as all public projects. To join a project, you will need to click 'Request to join' on the project's homepage which will send a message to the project's administrator. The project administrator will need to apprive your request to join before you can access the project.

<< Link to FAQs >> My collaborators can't see my project

# Create a New Project

You can create new projects via the Projects tab. Enter your project details in the pop-up form that appears.

As part of project creation, you will be asked to specify the inclusion/exclusion criteria for your project. These should be pre-specified in your protocol.

Once you have created your project, you can keep track of your project progress through the Project Details Page.

# 4.1 My project will have multiple screening stages

Currently, you can only have one set of inclusion/exclusion critiera per SyRF project. If you wish to have multiple screening stages in your systematic review, for instance title and abstract screening followed by full-text screening, you can do this by exporting your included studies from SyRF following your first screening stage and uploading them to a new SyRF project to complete your second screening stage.

Contact us for a link to a Shiny App which will allow you to export your data.

# Uploading a Systematic Search

### 5.1 Deduplicating your systematic search

If you have searched for studies using multiple databases there will be duplicate studies in your systematic search. Currently SyRF does not support deduplication of studies, and this must be performed before your studies are uploaded to SyRF. You can deduplicate your studies automatically using the CAMARADES deduplication tool.

### 5.2 Uploading files

You can upload your systematic searches as an:

- EndNote XML file
- Comma separated value (CSV) file
- Tab separated value (TSV) file

### 5.2.1 Uploading your studies from an EndNote export

- 1. Highlight (Ctrl+A) all references all the references in your EndNote library that you wish to upload to  ${\rm SyRF}$
- 2. Click File -> Export (NB: if you do not highlight all references only the first reference on your list will be exported)
- 3. Change the file type to XML

4. Name and save your XML file, which is now ready to be uploaded to SyRF

Please note that upload of studies with screening decisions is currently not supported with EndNote file uploads. If you have screening decisions for the studies you wish to upload, please use a CSV or TSV format instead.

### 5.2.1.1 Uploading from a Zotero export

Please note that you cannot use the 'EndNote XML' export option in Zotero to upload an EndNote file to SyRF. If you are using Zotero to manage your study references, please export as a CSV file and follow the CSV upload instructions.

<< Insert link to FAQ >> I am trying to upload an EndNote XML file that was created by importing from a place other than an electronic database and getting an error

### 5.2.2 Uploading your studies as a CSV or TSV

To upload your systematic search studies as a CSV or TSV file, you will have to make sure to format your data with following the column headings in order to make the upload work: \* Title \* Authors \* Publication Name \* Alternate Name \* Abstract \* Url \* Author Address \* Year \* DOI \* Keywords \* Reference Type \* PDF Relative Path

You can download a template with the correct column headings and example data here: << link to example csv >>

Even if you don't have information for all the columns specified, they will need to be in your file in order to make the upload work. SyRF will accept empty fields for any of these variables.

Files must first be saved as either Text - Tab delimited (\*.txt) or CSV - Comma delimited (\*.csv) files. This can be done in excel using the 'Save as type:' dropdown control in the 'Save As' dialog.

### 5.2.2.1 Upload studies with screening decisions

If you would like to upload studies along with screening decisions already made outside of SyRF, you should add separate columns for each user and SyRF's wizard will allow you to select which column headers in your file correspond to project members.

Within screening columns, decisions should be represented with the value 1 for inclusion and 0 for exclusion.

Your file should only contain the columns above and columns specified with screening decisions. If any columns are missing or additional columns are added (not specified for screening) the upload wizard will fail.

### 5.3 Uploading full-text PDFs

If you require full-text PDFs for each of your studies at any stage of your SyRF project, it is important that you have already retrieved these full-text PDFs before uploading your search file.

<< Link to EndNote guidance for PDF retrieval>>

In the systematic search file that you upload (csv/tsv spreadsheet or XML from Endnote) make sure the column "PDF Relative Path" contains relative path links (i.e. relative to the root of the folder you send to us) to your PDFs for each record.

You will then need to contact us with the name of your project, a folder containing your PDFs (sent via Google Drive or similar) and a CSV file containing the file path to each PDF and the title or SyRF study ID of each study, so we can match PDFs with your studies in SyRF.

We will upload these PDFs to the SyRF database and these can be opened from the screening form.

### 5.3.1 PDF file names

Please avoid using invalid characters (e.g. , < > : " \ / | ? \*) in file names as it may cause issues. By default, software like EndNote uses Author and Title information to name files, which can cause invalid characters to be added to your PDF file names. You can change the default to name PDFs using another column such as RecordID. Whichever columns you chose to name your PDFs with, the data should be unique.

<< Insert Link to FAQ >> I am performing a two-stage screening process and need to add PDFs only for my included studies for full-text screening

### 5.4 View project studies

You can now view project studies by clicking on the 'View Project Studies' button. This will show you all the studies you have uploaded to your project.

### 5.5 Deleting systematic searches

If you need to delete your systematic search, you can do so In SyRF. Be aware, however, that if you have used SyRF to screen or annotate these studies, deleting your systematic search will also delete these screening decisions and annotation answers.

# Project Stages

### 6.1 What are stages?

Stages are sections of your SyRF projects that you add to perform tasks such as screening and data annotation. Currently, SyRF allows you to add one screening stage and mutiple annotation stages. You can also have screening and annotation in the same stage.

### 6.2 Screening stages

A screening stage allows you to screen the studies you have uploaded to your SyRF project using the inclusion/exclusion criteria you entered when creating your project. Currently, as inclusion/exclusion criteria are defined at the project level, you can only have one screening stage per project.

If you wish to have multiple screening stages in your systematic review, for instance title and abstract screening followed by full-text screening, you can do this by exporting your included studies from SyRF following your first screening stage and uploading them to a new SyRF project to complete your second screening stage.

Contact us for a link to a Shiny App which will allow you to export your data.

### 6.3 Annotation stages

An annotation stage allows you to annotate data from your study, according to pre-defined questions in your systematic review protocol. SyRF has a question builder tool to allow you to design questions for your project.

Once you have designed all annotation questions, you should specify the stage at which you want to answer each question. You can do this before you start screening or after you have finished screening.

To add questions to your stage of interest go to the 'Stages' section of your project homepage and click 'Enter Stage'. You will then need to click on 'Stage Design' to start editing the stage.

### 6.3.1 Data extraction

To extract data from graphs in your systematic review studies, you will have to turn on data extraction in addition to annotation within your stage.

# 6.4 Screening and annotating within the same stage

If you want to carry out screening and annotation at the same time, you will also need to have these functionalities turned on (e.g. even if you have screened at a separate stage, you might want to have the functionality of being able to exclude a study at a later time point when you have read the full-text).

You will then be able to select the questions that you want to be included in this stage by checking the box next to the relevant questions.

# Screening

### 7.1 Screening using SyRF

You can screen the studies in your SyRF project against the inclusion/exclusion criteria you defined in your systematic review protocol and that you specified when you created your project by creating a screening stage in SyRF.

By default, two screeners have to agree on a study for it to be marked as insufficently screened. This is the recommended methodology for systematic reviews. If you are doing a student project and require only one screener, please contact us so that we can edit this for you.

### << UNLESS THIS IS AN OPTION IN THE NEW RELEASE >>

When you start reviewing in a screening stage, you will be shown the title and abstract of a random study from your systematic search uploads. If you have also uploaded PDFs, there will be a button to allow you to view the PDF.

Use the include and exclude buttons to record whether or not your study meets the inclusion/exclusion critiera. If you are unsure and want to come back to the study, click 'Next' to skip it and move on to another study.

Once you have decided to include or exclude a study, SyRF will record it as completed by you and you will not be shown it again. If you think you have made a mistake, it is possible to click the back button on your browser to go back to your previous study and re-screen it.

#### 7.1.1 Studies unavailable to screen

You will not be presented with studies that have been sufficently screened by other reviewers on the project. Instead, you can see how many studies have been sufficiently screened by other reviewers on your progress bar within the screening stage, marked as 'Unavailable'. Information on how to configure the number of reviewers required to sufficently screen each study can be found in the 'Number of Screeners' section below.

### 7.2 Number of screeners

By default, SyRF expects each study to be screened by two indepentent reviewers, with disagreements reconciled by a third reviewer, meaning you need at least three people to screen on your project. SyRF will check which studies have to be reconciled and they will become automatically available to a third person on the project.

If you are doing a student project and don't have other screeners, you can configure your stage to allow single screening in the Stage Settings. This is only recommended for student projects, and not for systematic reviews which you plan on publishing in a scientific journal.

If you wish to manually reconcile your screening disagreements, you can also configure this in the Stage Settings. Please note that this option is not recommended and you will have to contact our Help Desk to help you access your data for manual reconciliation.

<< Insert Link to FAQ >> My project only has two screeners, how can I see screening decisions?

# **Annotation Questions**

### 8.1 Creating annotation questions

In your systematic review protocol, you will have specified certain information you want to extract from each of your studies, such as 'Were animals randomised to experimental groups?' or 'What concentration of drug treatment was used?'. In SyRF you can annotate your studies with this information using annotation questions.

To design annotation questions for a stage of your project, go to 'Question Design' under 'Project Settingss' on the left-hand navigation bar of your project.

<< INSERT IMAGE >>

### 8.2 Question categories

Annotation questions are entered into the following categories depending on what information they ask about:

- Study level questions questions that apply to the entire study
- **Disease model induction questions** questions that apply to disease models
- Treatment questions questions that apply to treatment/s administered
- Outcome assessment questions questions that apply to the methods used to assess outcome
- Cohort questions questions that apply to experimental groups

### 8.3 Creating a new question

To add a question, simply click the '+' button next to the category you want to design a question in.

When adding a question, you can enter the following details:

- 1. The name of your question
- 2. A description of the question, which will be displayed alongside your question in SyRF
- 3. Whether the question accepts only a single answer or multiple answers
- 4. Whether the question is optional or required
- 5. The type of answer the question accepts (e.g. text, integers, decimals)
- 6. The question type (e.g. dropdown list, checkbox)

<< INSERT IMAGE >>

### 8.3.1 Question types

There are 6 question types in SyRF:

- Dropdown lists
- Autocomplete lists
- Radio buttons
- Checklists
- Check boxes
- Input boxes

We recommend that when using checklists or check boxes, you set the default checkbox status to 'indeterminate'.

### 8.3.2 Allowing multiple answers

If you choose to allow multiple answers, you will be asked if you want these to be split into separate annotations. This refers to how the data will be presented in your output data file.

Choosing to split muultiple answers into separate annotations means that, in your output file, the multiple answers will be separated into different rows. Choosing not to split into separate annotations, means that they will appear in the same row, separated by a semi-colon.

If you have nested questions, it may be best to choose to split your answers into separate annotations so that the nesting displays properly in your output

file. For example, in an 'Outcome Assessment' category question asking 'What behavioural tests are used?', we would allow multiple answers and may choose to split into separate annotations' as this allows us to ask further related questions to the specific behavioural test.

### 8.3.3 Nesting questions and conditional questions

Questions may be nested to allow for hierarchy of conditional information entry (i.e. questions can become active, depending on answers to previous questions).

### 8.4 Adding questions to stages

Once you have designed your questions, to allow them to be presented to reviwers, you need to add them to your annotation stage by going to 'Stage Settings' then 'Annotation' and selecting the questions you want to add to the stage.

### 8.5 Question categories

### 8.5.1 Study level questions

Enter any question that is relevant to the overall study.

e.g. Do the authors provide a study protocol that is available to you? (Yes or No checkbox)

### 8.5.2 Disease Model Induction questions

When adding a question you will be required to select whether the question relates to Control animals, Non-control animals (model animals) or both.

### 8.5.2.1 Control Question

Define questions that are specific to the Model control.

e.g. Do the control animals receive Sham surgery? (Yes or No checkbox)

### 8.5.2.2 Non-Control Question

Define questions that are specific to the Model

e.g. What type of surgery was carried out to induce the model? (Dropdown list with defined options)

### 8.5.2.3 Both

Define questions that are relevant to both Model control and Model animals e.g. What anaesthetic was used for both the model and sham surgery? (Dropdown list with defined options)

### 8.5.3 Treatment questions

When adding a question you will be required to select whether questions related to Control animals, Non-control animals (treatment group animals) or both.

### 8.5.3.1 Control question

Define questions that are specific to the Treatment control

e.g. What is the vehicle given to the control animals? (Dropdown list with defined options)

#### 8.5.3.2 Non-Control question

Define questions that are specific to the Treatment group

e.g. Specify the dose of treatment drug given in mg/kg (Decimal input box)

### 8.5.3.3 Both

Define questions that are relevant to both Treatment control and Treatment animals

e.g. What route of drug or vehicle administration is used in the experiment? (Dropdown list with defined options)

### 8.5.4 Outcome assessment questions

Define questions relevant to each outcome assessment procedure in the study.

e.g. What is the behavioural test used to measure outcome? (Dropdown list with defined options)

### 8.5.5 Cohort level questions

Define questions relevant to each cohort (experimental group) in the study.

e.g. What is the sex of the animals included in the cohort? (Dropdown list with options males, females, both, unknown)

<< Insert FAQ link >> I have cohorts with comorbidities and I'm not clear on how to differentiate between them.

### 8.5.6 Experiment questions

Define questions relevant to each experiment in the study e.g. Was there a habituation period? (Yes or No checkbox)

### 8.6 Nesting questions

As previously, mentioned, for each question you can choose to add related questions, if you want to get answers to additional questions, which are conditional on the answer to the previous question.

e.g. "What type of model is used?" (Drop down list with option of: Pharmacological or Surgical)

We could then add a related question by selecting "Add Related", and in the form "conditionally display based on parent question".

e.g. "What is the drug given?" (Drop down list with options of different drugs)

You could then ask further related questions, by clicking on this question and selecting 'Add Related' and asking for each drug selected: "What is the dose and route of delivery?" If Surgical is selected then we may ask the related questions: "What was the anaesthetic used?" or "What was the site of lesion?".

These questions will nest under the previous question.

### 8.7 Question editing

At present, questions can only be added or deleted. Individual questions cannot be edited or changed order. This is primarility due to our database structure and questions and an editing feature will be implemented in an upcoming release.

# $\mathbf{FAQ}$

If you have a question related to SyRF that is not covered by this user guide, please check our frequently asked questions page on the SyRF website.

<>