

MolGears Docs

version 1

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The MolGears's documentation!

An overview of the MolGears

What is it?

This is a opensource framework/database/toolkit based on a [TurboGears](#) web framework and [RDKit](#) open source toolkit for cheminformatics:

- programed in Python2.7 language
- BSD license
- source code on Github (<https://github.com/admed/molgears>)

Software Dependencies/Third-party software component

- RDKit (<http://www.rdkit.org>)
- TurboGears (<http://turbogears.org/>)
- Postgresql Database (<http://www.postgresql.org/>)
- RDKit database cartridge for postgresql (<http://www.rdkit.org/docs/Cartridge.html>)
- Genshi (<http://genshi.edgewall.org/>)
- JSME editor (<http://peter-ertl.com/jsme/>)
- **Python based libraries:**
 - SQLAlchemy (<http://www.sqlalchemy.org/>)
 - razi (<http://razi.readthedocs.org/en/latest/index.html>)
 - Numpy (<http://www.numpy.org/>)
 - Scipy (<http://www.scipy.org/>)
 - Matplotlib (<http://matplotlib.org/>)
 - Xlwt & Xlrd (<http://www.python-excel.org/>)
 - xhtml2pdf (<http://www.xhtml2pdf.com/>)
 - pillow (<https://github.com/python-pillow/Pillow>)

Design goal



- Project management tool
- Efficient data storage
- Sorting, analysis, aggregation and reporting of data
- Data visualization
- Improved data access
- Automation of procedures
- Facilitate communication

Features

- Multi-projects
- Adding molecules by drawing or pasting SMILES code
- Reading molecules from file (csv, smi, sdf, mol, txt)
- Data presenting in sortable columns
- Exporting data to file (file formats: xls, pdf, csv, sdf, txt, png)
- PAINS (Pan Assay Interference Compounds) filtering ([DOI: 10.1021/jm901137j](https://doi.org/10.1021/jm901137j))
- History of changes
- Compound filtering (by similarity, structure, identity, compound name, creator, adding date etc.)
- Stars rating
- IC50 determination based on least squares method,
- Automatic data processing
- Graphs generation
- Access managing
- Tags

License

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Project Workflow Overview

Multiproject

- MolGears allows you divide your work into projects. The solution is designed to handle project management for small compounds.

- Molgears allows you to manage access and grant authorized users the right to use the service and access permission for each project independly.
- **In addition each project has basic workflow divided into tasks:**
 - designing process,
 - ordering synthesis,
 - tracking sythesis process,
 - storage compound in library,
 - result tool for compounds activity.

Project basic workflow

- A repeatable process that brings similar stages each time
- A clear division of responsibility between different people
- A better basis to estimate task length
- A simple method to communicate process and data to all or selected teem members/employees/colabollators/
- Each task has history records so you can see who added, approved and edited each record.

Basic Workflow Schema

For each project in menu bar you have access to 5 main tables connected to the project workflow.

- Compounds - root table. Storage for molecules structures; both: ideas and existing compounds. Unique records for each structure.
- Requests - put ideas into real things. Table for synthesis requests.
- Synthesis - table for tracking synthesis progress, priority managing, and analytical data storage.
- Library - table only for existing compounds. The library of compounds with ability for tracking location and the amount of compound.
- Results - compound activity data storage connected to library instance. It's allowed to add for one library record many results.

Workflow schema for the tables:

I. Add structures to Compounds table.

II. You can choose two path:

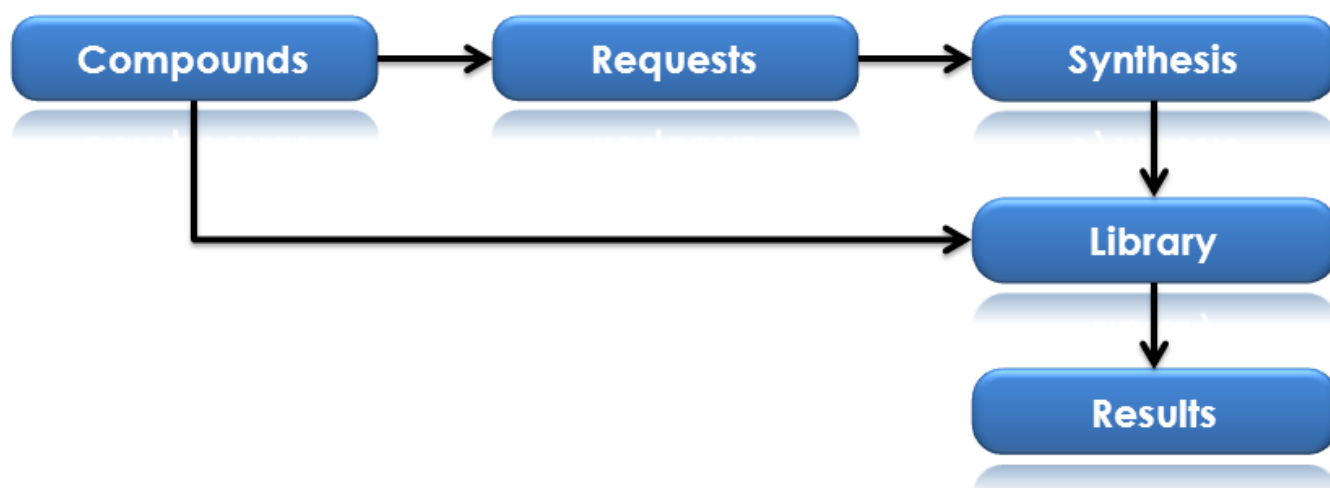
1. Internal synthesis:

- a. Create synthesis request
- b. Accept synthesis by chemist and proceed synthesis phases
- c. Add finished compound from synthesis to library

2. External synthesis (e.g. you're buying ready product):

- a. Accept compound directly to library

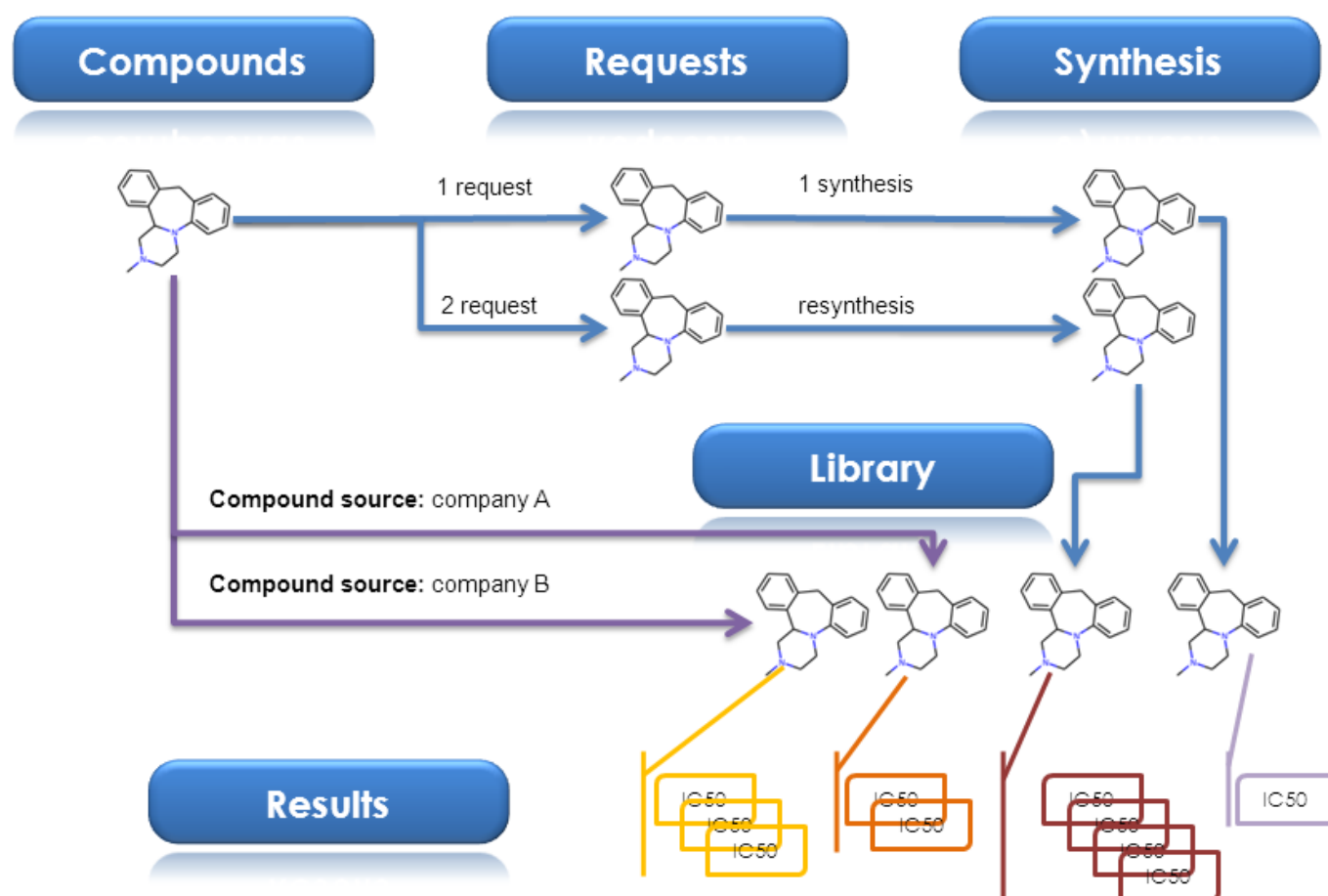
III. Add one or more activity results to library instance.



Example for the various sources of sample

Compounds table is containing unique structures of molecules. This means you can't add the same structure twice to Compounds Table. Only exception for this rule is adding compound as isomer. **[references]**. This rule also don't apply to other tables. You can create many request for one compound (structure) and then accept them to synthesis. One of the example of using this mechanism is performing synthesis of small amount of some compound. Later when we need more we can order the resynthesis by putting the same compound second time into request and follow the workflow once again.

Separate case is when you don't want to carry out the synthesis procedure and want to add a compound directly to the library e.g. when you're buying ready product from *company A*. Then follow the second path in workflow (see [Project basic workflow](#)). But you can also add this like that many times. If you add this twice and earlier when you performed 2 synthesis procedure as a result you will have 4 instances in library table referenced for the same structure. Than you can add one or more activity results independently for each of library instance of your compound. The shema illustrating this is presented below:



Database model

Simple Model

Database model was built based on Project Workflow (see: [Project Workflow Overview](#)).

Compounds and **Projects** tables are connected by many to many relationship with allowing to adding one compound to many projects and many compounds to one project.

Compounds Table is containing basic information about molecule structure like:

- SMILES and InChi code,
- name
- fingerprints
- molecular weight
- logP
- number of atoms with and without Hydrogens
- number of rings
- Hbond acceptors
- Hbond donors
- Teoretical PSA

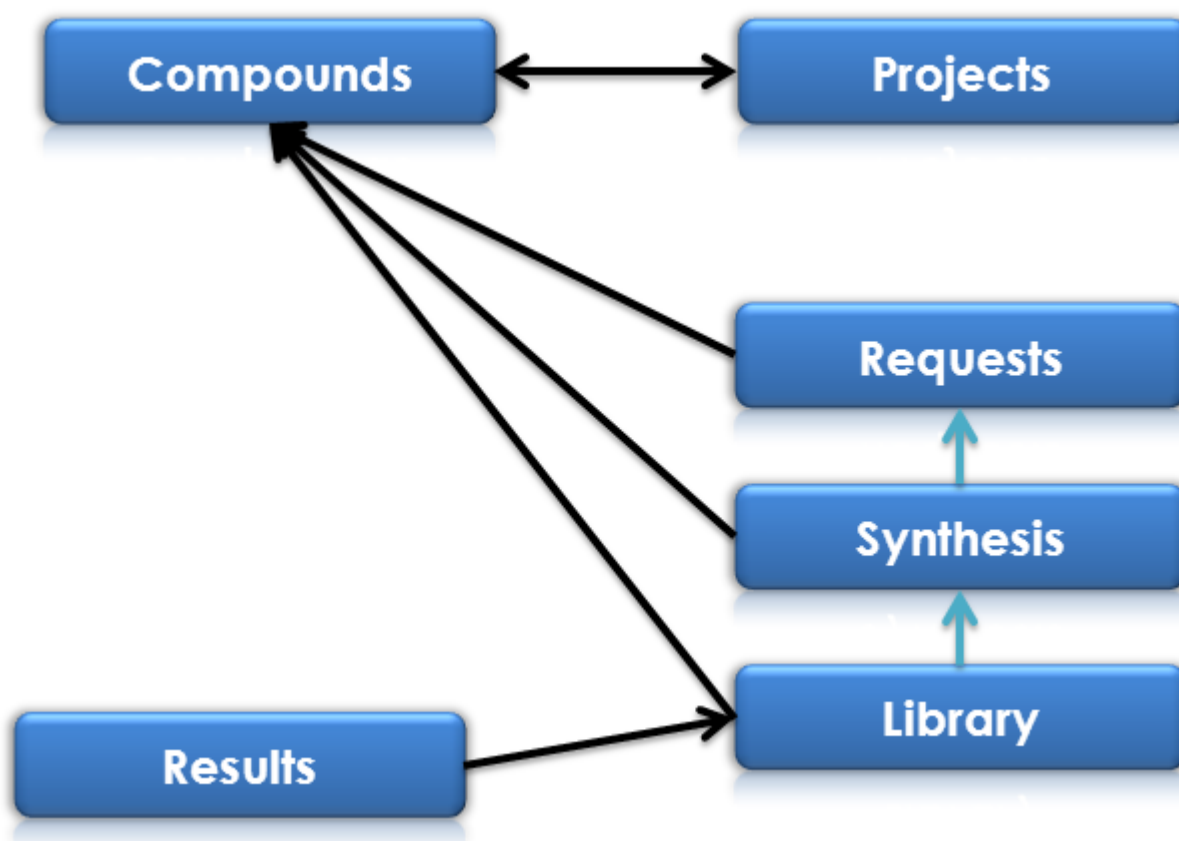
Requests, **Synthesis** and **Library** tables are referring to this data by relationship to the **Compounds**. This mean that editing and changing structure for Compound will make changes for all branches in all connected tables.

Synthesis table is storing ID of **Request** instance from which it is derived. This making the possibility for tracking the pathway of compound from one to other table and the ability to implement The Project Workflow in a consistent way.

The same connection is between **Synthesis** and **Library** tables.

Results table has many to one relationship to **Library**. In this way it's available to add many results to one library instance.

Basic schema of database model is presented below:



Detailed Model

Entity - Relationship Diagram for MolGears Database model is presented below.

Diagram was generated using [DBSchema](#).

Main 6 Tables are respectively named as:

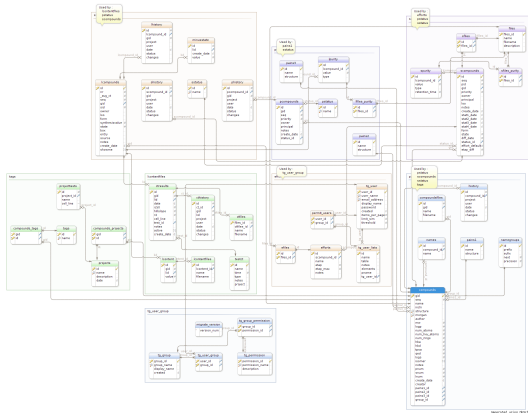
Model Name	SQL name
Projects	projects
Compounds	compounds
Requests	pcompounds
Synthesis	scompounds
Library	lcompounds
Results	ctoxicity

- Each of the main tables (except the Projects Table) has dedicated history tables for storing changes.
- Request and Synthesis tables have dedicated tables for Status.
- Compounds, Synthesis, Library and Results tables have dedicated tables for storing files.
- There are 3 PAINS tables for storing SMARTS codes.

Getting started

- The tables for access managing have "tg_" prefix (tables for Users accounts, Users Lists, Groups and Permissions).
- Other tables are auxiliary and serves i.a. for auto-naming, state and purity monitoring, test description, tags storing etc.

ER Diagram (click the image to enlarge):



Getting started

Installing your doc directory

You may already have sphinx [sphinx](#) installed -- you can check by doing:

```
python -c 'import sphinx'
```

If that fails grab the latest version of and install it with:

```
> sudo easy_install -U Sphinx
```

Now you are ready to build a template for your docs, using sphinx-quickstart:

```
> sphinx-quickstart
```

accepting most of the defaults. I choose "sampledoc" as the name of my project. cd into your new directory and check the contents:

```
home:~/tmp/sampledoc> ls
Makefile      _static      conf.py
_build       _templates   index.rst
```

The index.rst is the master ReST for your project, but before adding anything, let's see if we can build some html:

```
make html
```

If you now point your browser to `_build/html/index.html`, you should see a basic sphinx site.

Fetching the data

Now we will start to customize our docs. Grab a couple of files from the [web site](#) or git. You will need `getting_started.rst` and `_static/basic_screenshot.png`. All of the files live in the "completed" version of this tutorial, but since this is a tutorial, we'll just grab them one at a time, so you can learn what needs to be changed where. Since we have more files to come, I'm going to grab the whole git directory and just copy the files I need over for now. First, I'll cd up back into the directory containing my project, check out the "finished" product from git, and then copy in just the files I need into my sampledoc directory:

```
home:~/tmp/sampledoc> pwd
/Users/jdhunter/tmp/sampledoc
home:~/tmp/sampledoc> cd ..
home:~/tmp> git clone https://github.com/matplotlib/sampledoc.git tutorial
Cloning into 'tutorial'...
```

```
remote: Counting objects: 87, done.
remote: Compressing objects: 100% (43/43), done.
remote: Total 87 (delta 45), reused 83 (delta 41)
Unpacking objects: 100% (87/87), done.
Checking connectivity... done
home:~/tmp> cp tutorial/getting_started.rst sampledoc/
home:~/tmp> cp tutorial/_static/basic_screenshot.png sampledoc/_static/
```

The last step is to modify `index.rst` to include the `getting_started.rst` file (be careful with the indentation, the "g" in "getting_started" should line up with the ':' in `:maxdepth:`):

Contents:

```
.. toctree::
   :maxdepth: 2

   getting_started.rst
```

and then rebuild the docs:

```
cd sampledoc
make html
```

When you reload the page by refreshing your browser pointing to `_build/html/index.html`, you should see a link to the "Getting Started" docs, and in there this page with the screenshot. *Voila!*

Note we used the image directive to include the screenshot above with:

```
test
```

Indices and tables

- *genindex*
- *modindex*
- *search*