DIMSpec Quick Guide - Shiny Web Applications

Jared M. Ragland*

2023-11-29

The Shiny platform and DIMSpec

A driving goal of the Database Infrastructure for Mass Spectrometry (DIMSpec) project is to provide a database that can be easily retasked to support individual projects to manage data coherently and accelerate analyte identification, screening, and annotation processes for non-targeted analysis projects. A DIMSpec mass spectral database incorporates empirical mass spectral data from analytical standards and complex mixtures with relevant analytical method metadata and mass spectral annotation.

For many users, the majority of their interaction with the DIMSpec project will be through graphical user interfaces providing access to the underlying data and functionality. This simplifies their experience and increases the value of the project tremendously. Web applications in the DIMSpec project are powered by the **shiny** R package. Provided applications communicate with the database through the application programming interface (API) provided by the **plumber** R package, though others can be readily developed that connect directly to the database if desired.

This quick guide introduces the three web applications that ship with the project, and the application template designed to accelerate development of new applications. Refer to the DIMSpec User Guide for installation and other details of each application introduced here; information contained here is a selected subset of the DIMSpec User Guide and does not provide an introduction to the shiny platform.

DIMSpec Web Applications

The following subsections provide an introduction to web applications provided in the DIMSpec project.

- 1. Table Explorer
- 2. DIMSpec Quality Control (DIMSpec-QC)
- 3. Mass Spectral Match (MSMatch)
- 4. Application Template

Every effort has been made to make launch of web applications as straight forward as possible. Toward that end, all applications that ship with the project can be launched directly from the command line and will establish all necessary aspects automatically. From an R console opened at the project directory, issue the command shiny::runApp(file.path("inst", "apps", "X"))¹ where "X" is the name of the directory containing an application. This is even simpler if the compliance file has been sourced in the current session; use the helper function start_app("X") to launch the app in directory "X"².

^{*}NIST | MML | CSD | jared.ragland@nist.gov

¹Apps are also launchable on your local network by using shiny::runApp(file.path("inst", "apps", "X"), host = "0.0.0.0", port = Y) where Y is an integer corresponding to an open port on the hosting machine. This does require that other computers on your network can connect to the host machine.

 $^{^{2}}$ A list of recognized shiny applications is available in the session environment as the named vector SHINY_APPS after the compliance file has been sourced.

Table Explorer

Often, one of the friction points in using a database is understanding its schema. For this reason, the DIMSpec project ships with the "Table Explorer" app to facilitate visual exploration of the DIMSpec database schema. It also served as proof-of-concept for the database/API/shiny approach and was used as the basic skeleton of the template app that ships with the project.

Table Explorer is a simple entity viewer for the attached database. Combining the comment decorations in DIMSpec and reading of entity definitions from the database (see Inspecting Database Properties in the DIMSpec User Guide for details) allows for R to expose a wealth of information about the underlying schema and quickly change which entity is being viewed. See above for details of how to launch this app, but the easiest method is after the compliance.R script has been executed, use start_app("table_explorer") to launch it in your preferred browser.

There is only one page for interactive content, named "Table Viewer" (Figure 1). A navigation bar on the left controls the current page being viewed; collapse the bar using the "hamburger" icon at the top next to the NIST logo. Click the drop down box to change the database table or view being displayed. This will update the definition narrative immediately below the selection box and display the contents of that table to the right.

A graphic of the entity relationship diagram is provided on the second page. Click it in the navigation panel to the left. This can be viewed in full resolution by right clicking the diagram and opening it in a new browser tab (language varies by browser).

	≣		
Currently connected to DIMSpec for PFAS	Select a table to view additive, aliases	Table data Show 10 v entries	Search:
• Ø Table Viewer	additive_anases	Show to + entries	search:
	List of common aliases for each entry in	additive_id 🍦	alias
📥 Entity Relationship Diagram	norm_additives This table references table norm_additives and is used in	All	All
	view_additive_aliases .		
	additive_Id is an INTEGER column that is required. Foreign key to norm_carriers. It references column id in table norm_additives. alias is a TEXT column that is required and must be unique. Human meaningful name(s) associated with an additive.	1	acetic acid ammonium salt
		1	acetic acid, ammonium salt
		1	Ammonium Acetate
		1	InChI=1S/C2H4O2.H3N/c1-2(3)4;/h1H3,(H,3,4);1H3
		1	USFZMSVCRYTOJT-UHFFFA0YSA-N
		1	631-61-8
		1	C2H4O2H3N
		1	C2H7NO2
		2	Ammonium Formate
		2	CH5NO2
		Showing 1 to 10 of 46 entries	Previous 1 2 3 4 5 Next

Figure 1: The Table Explorer main page.

DIMSpec Quality Control (DIMSpec-QC)

Quality control and assessment is key for trust and ensuring data integrity. Algorithms were developed for the DIMSpec project to validate the quality of new experimental data and build files suitable for database import. The DIMSpec Quality Control (DIMSpec-QC) application was developed to accelerate this quality review process and installs alongside DIMSpec.

Two preprocessing steps are required to use DIMSpec-QC. First, raw data files produced by a mass spectrometer must be converted into mzML format using Proteowizard's MSConvert software (Adusumilli, Ravali and Mallick, Parag 2017) There are specific parameters that must be used during conversion. A more detailed user guide for converting the files is provided as another quick guide.

Filter: Threshold peak filter Threshold type: absolute Orientation: most intense Value: 1 Filter: Peak picking Algorithm: vendor MS levels: 1-2

2. The second step prior to using DIMSpec-QC is to use a macro-enabled Microsoft Excel workbook, called the Non-Targeted Analysis Method Reporting Tool (NTA-MRT) for the systematic collection of sample, method, and compound information related to chemicals identified in a sample. The most up-to-date version of NTA-MRT is publicly available at GitHub. Instructions for completing the NTA-MRT are contained within the tool itself. In order to use the DIMSpec-QC application, a sample. JSON file must be generated using the "Export to JSON file output" button on the first tab of the NTA-MRT.

The file name entered in the NTA-MRT under the Sample tab must exactly match (casesensitive) the paired mzML file name to be used for the DIMSpec-QC.

See above for details of how to launch this app; the easiest method is from the console after the compliance.R script has been executed, use start_app("dimspec-qc") to launch it in your preferred browser. The "About" screen will appear when the DIMSpec-QC application is available (Figure 2). The navigation panel on the left will control which page is currently being viewed; click an entry to navigate to that page. The app itself should guide you through the process in a straightforward manner. Click the "Click Here to get Started" button on the "About" page or click "Data Import" in the navigation panel on the left to get started.

	E					
Mass Spectral Quality Control for PFAS	Quality Control Check Tool for Data Import					
? About						
🗎 Data Import	Click Here to get Started					
🛓 Export Data						
ିକ QC Settings	Welcome to the NIST database for identification of perfluoroalkyl substances (PFAS) by non-targeted analysis (NTA) using high resolution accurate mass spectrometry (HRAMS). The goal of this project is to improve access to and utility of analytical data for the confidence dicovery, identification, and source-attribution of PFAS in environmental matrics.					
	The quality control check tool for the PFAS Mass Spectral database is for the evaluation of mass spectral data for general quality and for the generation of import files that can be directly incorporated into the database.					
	In order to use this application, the following must be done:					
	Raw data files must be converted to mzML (link here for directions) Sample JSON files for each raw data file must be generated, using the Method Reporting Tool					
	More information here					
	This project was funded in part by the Department of Defense Strategic Environmental Research and Development Program grant ER20-1056. The application you are using now was developed by researchers in the Chemical Sciences Division of the Material Nessurement Laboratory at the National Institute of Standards and Technology (NIST). More information about NIST's PFAS program is available at their site. For more information, please contact the NIST PFAS Program at pfas@nist.gov.					

Figure 2: The home page for the DIMSpec-QC web application contains basic information about the application and can be tailored easily for each use case.

Step 0 (Optional) - Set QC Parameters

Prior to file processing, verify or set parameters to be used for quality control analysis from the "QC Settings" page. The default settings are recommended, but can be modified if needed.

Step 1 - Import Data

From the "Data Import" page, upload paired mzML and sample JSON files (conventionally named "filename_mzml_sample.JSON" by the NTA-MRT macros). Multiple files not exceeding 250 MB may be uploaded; see the shiny documentation for details on changing file size limits if necessary.

- 1. Load the mzML files of interest using the top "Load" button or by dragging the data file(s) to the input widget labeled 1) Load raw mzML file(s).
- 2. Load the paired sample JSON files using the second "Load" button or by dragging the JSON file(s) to the input widget labeled 2) Load Sample JSON file(s). The sample JSON files do not need to be selected in the same order as the mzML files and will be matched by name when processed.
- Once mzML and Sample JSON files are uploaded, the application will automatically check to see if there are valid pairs of mzML and Sample JSON files. A successful upload will update the current screen. If multiple files are loaded, only files with verified matches will be included in the resulting table and available for further processing.

Step 2 - Process Quality Control Metrics

After files have been loaded and matched, a button will appear on the "Data Import" window that is labeled "Process Data". Click the "Process Data" button and it will sequentially process each mzML file. This can take up to 5 minutes per raw file depending on the number of compounds per file, so a large number of files may take a long time to process. Progress indicators are provided. If a raw file does not have a valid Sample JSON, the files can still be processed, but the invalid rows will be excluded.

Step 3 - Evaluate Data Quality

Once processed, the QC results can be reviewed by selecing "Quality Review" in the left menu or clicking the "Quality Review" button that appears below the "Process Data" button. The top table, which is the only visible table when starting a new review, shows the raw files that have been processed and the respective quality control check results ("PassCheck"). If all compounds in the raw file passed all QC checks, the PassCheck result will be true. If any compound in the raw file failed any of the QC checks, the PassCheck result will be false. To review the compounds within a single raw file:

- 1. Click a row in the table at the top left to select an mzML file. This will display a second table of all compounds within the selected raw file. The PassCheck result for each compound is displayed in this table. If all QC checks for each compound in the raw file passed all QC checks, the PassCheck result will be true for that compound. If a compound in the raw file failed any of the QC checks, the PassCheck result will be false for that compound. To review the individual QC checks (described in the DIMSpec User Guide), select a row for the peak to review in the table labeled
- 2. Click a row in the second table on the left to see metrics for that peak. This will display boxes to the right containing all QC checks for that compound. Expand a specific QC check by clicking on the box header to display the results of the QC check as a table. A note below the two tables on the left will indicate whether any QC checks failed.

Step 4 - Export Data

Once data are processed, all data can be exported (regardless of quality review status) by selecting "Export Data" in the left menu. Additional options may be added in the future to refine the export process such as selecting only peaks and files that pass all defined quality checks. Clicking the button labeled "Export all data", will write the peak JSON files and download them in a single .zip file. This file can be unzipped and the peak JSONs can be directly incorporated into the DIMSpec database using the import routine described in the DIMSpec User Guide or the Quick Guide - Importing Data.

Using the DIMSpec-QC application greatly accelerates the process of performing quality control and assurance tasks for non-targeted data and ensures checks are performed consistently across data files. It serves as a key component of using the DIMSpec project in an active research lab setting when adding to the mass spectral library.

Mass Spectral Match for Non-Targeted (MSMatch)

A common user need addressed by DIMSpec is identification of analytes in a sample measured by high resolution mass spectrometry. Often the components are unknown but may be suspected. Ability to identify not only an analyte from its mass spectra, but to identify known fragments for unknown analytes is a key need; when the analyte itself cannot be identified, non-targeted analysis workflows make use of fragmentation patterns to determine or reconstruct its identity. The Mass Spectral Match for Non-Targeted Analysis (MSMatch) application was built to accelerate non-targeted analysis projects by searching experiment result data in mzML format for matches against a curated mass spectral library of both compounds and annotated fragments.

As with the DIMSpec-QC app application, raw data files produced by a mass spectrometer must be converted into mzML format using Proteowizard's msConvert software. There are specific parameters that must be used during conversion.

Filter: Threshold peak filter Threshold type: absolute Orientation: most intense Value: 1 Filter: Peak picking Algorithm: vendor MS levels: 1-2

A more detailed user guide for converting the files is provided as another quick guide. This data format conversion step is the same as for the DIMSpec-QC application.

See above for details of how to launch this app, but the easiest method is after the compliance.R script has been executed, use start_app("msmatch") to launch it in your preferred browser. The "Home" screen will appear when the MSMatch application is available (Figure 3, next page). The navigation panel on the left will control which page is currently being viewed; click an entry to navigate to that page. The app itself should guide you through the process in a straightforward manner. Click the "Click Here to get Started" button on the "Home" page or click "Data Input" in the navigation panel on the left to get started.

Every effort has been made to make using MSMatch as intuitive as possible. Hints in the form of tooltips are provided throughout; hover over question mark icons or controls to see them. These can be toggled on and off at any time using the "Show Tooltips" toggle button at the bottom left of the application window (see Figure 3 inset at bottom right. If enabled, advanced search settings can be similarly toggled on and off for the session (see the DIMSpec User Guide for instructions on how to set default accessibility and settings for tooltips and advanced settings). The "hamburger" icon at the top left of the screen will collapse the

left-hand navigation panel to provide more horizontal room on smaller screens, though the application will rearrange itself when screens are smaller than a minimum width.

Click the "Click Here to Get Started" button to begin (Figure 3, top). This will activate the "Data Input" page (Figure 4, next page). Example data files are provided in the project directory ("/example/PFAC30PAR_PFCA2.mzML" and "/example/example_peaklist.csv").

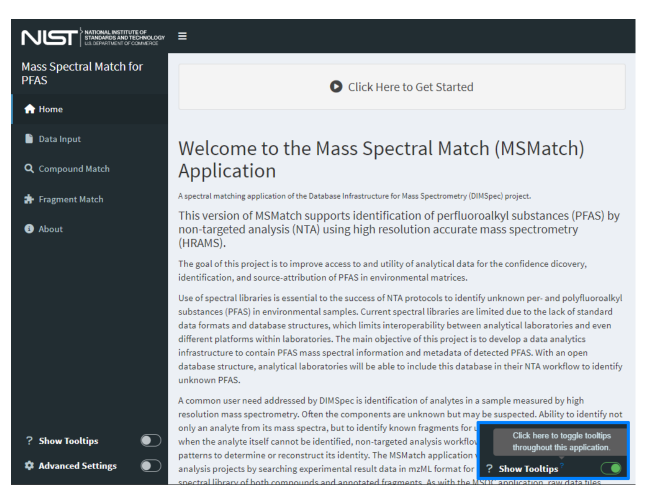


Figure 3: The home page for the MSMatch web application contains basic information about the application and can be tailored easily for each use case for reuse.

Step 1 - Load an mzML Data File

MSMatch only accepts files in the mzML format (see the note on preprocessing above for details). Either click the "Load" button to select a file or click and drag one from your file system to that widget. Set instrument parameters to match those used during data collection using the controls provided. Files must be less than 250 MB; see the shiny documentation for details on changing file size limits with shiny.maxRequestSize if necessary.

Load Data	Feat	ure Identification	
Choose a data file (.mzML) Sele	ect par	rameters identifying peaks to examine.	
Load Select a file to begin		+ Add	
Ir	mport	Select a file to import parameters	
Set Instrument Parameters			
MS Experiment Type			
DDA (data-dependent acquisition) 🔻			
Relative Error (ppm)			
5			
Minimum Error (Da)			
0.002			
Isolation Width (Da)			
0.7			

Figure 4: The Data Input page for the MSMatch web application.

Step 2 - Identify Features of Interest

Add features of interest from the Data Input page (Figure 4) by providing the mass-to-charge ratio and retention time start/centroid/end values for each feature. Two methods (Figure 5) are supported to identify features of interest by mass-to-charge ratio and retention time properties. Either use case is fully supported. Users may:

- 1. Import a file (Figure 5, left; either .csv, .xls, or .xlsx, though workbooks should have relevant data in the first worksheet) and identify which columns contain the correct information.
 - Click "Import" and select a file of interest from your local computer or drag and drop a file to this input.
 - Use this method if you have a file containing features of interest from other procedures or software outputs to quickly import many feature properties.
 - Select a column that corresponds to each property.
 - To append to the current list, keep the checkbox checked. To overwrite, uncheck this box.
 - Click "Load Parameters" to validate and add parameters or "Cancel" to abort this operation.
 - Repeat until all files are imported. OR

- 2. Click the "Add" button and enter search parameters one at a time (Figure 5, right, next page); repeat this process to add more.
 - Add numeric values for all items.
 - Click "Save Parameters" to validate and add or "Cancel" to abort this operation
- Users receive feedback on the form if values are left blank or if they do not meet expectations (e.g. centroid is after peak start and before peak end).
- Values should all be numeric in nature.
- This list may be edited after import by clicking the "Edit" button Figure 6 (next page).

Import peak search parameters	Peak search parameters
Select one column from your file	Precursor m/z
corresponding to each of the following parameters.	327.4586
	Retention Time (Centroid)
Precursor m/z	13.213
mz	Retention Time (Start)
Retention Time (Centroid)	12.987
rt 💌	Retention Time (End)
Retention Time (Start)	13.687
peak_starttime 💌	Save Parameters × Cancel
Retention Time (End)	
peak_endtime 🔻	
Append to the current parameter list	
Load Parameters X Cancel	

Figure 5: Dialogs to identify features of interest by upload (left) or manually by clicking the Add button (right).

Select para	meters identifyir	ng peaks to examin	ie.			Select para	meters
		+ /	Add				+ Add
Data will be	searched for peak	ks matching these c	haracteristics. Select	a row to edit or remove	it.	Data will be	search
P	recursor m/z	♦ RT	RT Start	RT End	÷	Р	recurso
	312.973	10.8	10.5	11.1			312.9
	327.4586	13.213	12.987	13.687			327.45
	362.9699	12.09	11.9	12.4			362.96
	412.9665	13.05	12.8	13.4			412.96
Import	Select a file to in	nport parameters				Import	Selec

Select parameters identifying peaks to examine

	+ Add	🖋 Edit	🖌 Edit 🗙 Remove			
Data will be searched for peaks matching these characteristics. Select a row to edit or remove i						e it.
	Precursor m/z	♦ RT ♦	RT S	tart 🍦	RT End	$\frac{\mathbb{A}}{\mathbb{V}}$
	312.973	10.8	10	.5	11.1	
	327.4586	13.213	12.9	987	13.687	
	362.9699	12.09	11	.9	12.4	
	412.9665	13.05	12	.8	13.4	
Import	Select a file to i	mport parameters				

Figure 6: Manage the feature identification list (left) interactively by adding, editing, or removing features as needed (right) by selecting a row from the table and clicking the appropriate button.

Data are ready to be processed once features of interest are added. Selecting any row in the resulting table makes two additional functions available (Figure 6, right, next page). With a row selected, click "Remove" to delete it or "Edit" to bring up the same form as above (Figure 5, right), change the values, and click "Save Parameters." All records remaining in the feature of interest list will be available to search widgets on subsequent pages.

Step 3 - Generate the Search Object

Clicking the "Process Data" button. Buttons will appear to navigate to the "Compound Match" and "Fragment Match" pages once processing is complete.

Step 4 - Explore Results

Navigate to the desired search page. Algorithmic matching of provided mass spectral data for features of interest is performed against data stored in the attached database. Matching algorithms are described in detail in the DIMSpec User Guide). Both the "Compound Match" and "Fragment Match" pages are laid out in similar fashion. Select a feature of interest from the drop down box at the top of the page and click the "Search" button. Continue with your investigation of matches for the selected features.

The Compound Match page will search features of interest for known mass spectral pattern matches against the attached database (Figure 7, next page; Figure 8, next page).

Use the Fragment Match page to explore fragmentation patterns in more detail for features of interest with compound matches. This page, however, is of more use when features do not match with a known compound. It will identify and label identifiable fragments, assisting with the NTA process of fragment annotation. It also provides information about the sample and compound(s) from which identified fragments were obtained (Figure 9, next pages; Figure 10, next pages).

Top match is Perfluorohexanoic	Select a row to view the m	atch or send	it to uncerta	inty estimati	on, or click a	button at the botto	m to save this tabl	e.
acid from an analytical standard ✓ This is the most common (n = 9 of 9) compound match. MS1 Score: 0.9989 (rev score 0.9972) MS2 Score: 0.9939 (rev	Compound	MS1 Score	MS1 Score ≑ (Rev)	MS2 Score	MS2 Score 🔶 (Rev)	# Annotated 🝦 Fragments	# Annotated ≑ Structures	# Annotated Citations
score 0.9999) Comparison Mass Spectrum	Perfluorohexanoic acid	0.9989	0.9972	0.9939	0.9999	2	2	2
Your measurement is in black . Comparison spectrum is in red .	 Perfluorohexanoic acid 	0.9883	0.9843	0.7424	0.6647	1	1	1
ACH V ACH	Perfluorohexanoic acid	0.965	0.7849	0.3342	0.4294	1	0	0
i i	Perfluorohexanoic acid	0.9633	0.7331	0.3342	0.3357	0	0	0
0.4-	Perfluorohexanoic acid	0.9717	0.626	0.3342	0.4054	1	0	0
-0.0	Perfluorohexanoic acid	0.9419	0.7254	0.3342	0.2375	1	0	0
Relative	Perfluorohexanoic acid	0.9695	0.7948			1	0	0
-0.4-	Perfluorohexanoic acid	0.9359	0.8033			1	0	0
100 150 200 250	Perfluorohexanoic acid	0.964	0.6962			1	0	0
100 150 200 250 m/z	Copy CSV Excel							

Figure 7: Results of a compound match for the selected feature of interest. Download results using the buttons at the bottom left of the match table.

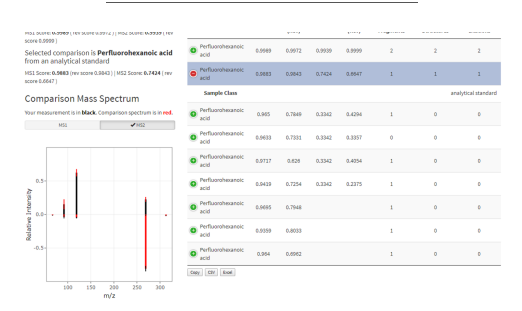


Figure 8: Results change in real time when different rows are selected from the table, updating the narrative, butterfly plot, and method narrative (compare with Figure 7).

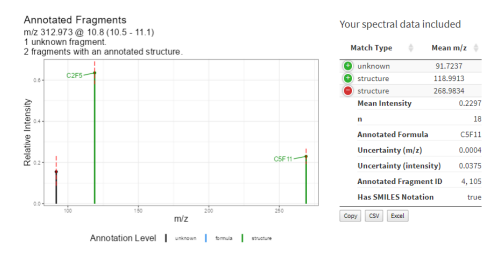
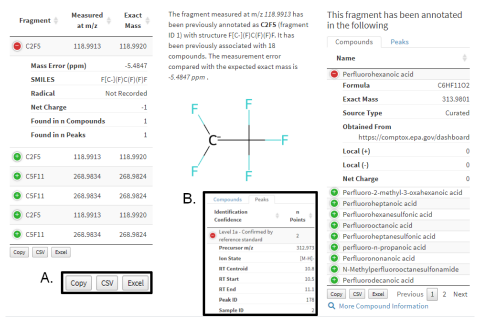


Figure 9: Automatic fragmentation annotations based on fragments listed in the database



Select a row in the left-hand table to view additional information for that annotated fragment.

Figure 10: A variety of additional information is available for known, previously annotated fragments

Application Template

The Table Explorer, DIMSpec-QC, and MSMatch web applications serve to increase utility and usability of the DIMSpec project by making data and code available for casual users. They can be launched locally from the project, or hosted on a network. These are not, however, intended to provide exhaustive functionality for potential users.

The underlying infrastructure enabling these apps has been constructed as to make adding new ones as easy as possible. Toward that end, the Table Explorer app served as a proof-of-concept to develop the application template at /inst/apps/app_template. It is in the "3 file" format and contains subdirectories for data, modal dialogs, and app resources (in the www subdirectory). To start developing a new app within the project, simply copy and rename the entire app_template folder. The new app will benefit from the same environment establishment routines used elsewhere. Simply edit the /global.R file as needed, add any data and image or resource files, set need_files in /global.R to include any .R files you want available when the app is launched, and begin development with an assured connection either directly to a database or by leveraging the plumber API. An isolated /app_functions.R file is also provided for custom functions the new app will require.

A Note on Settings

In order to set up the project to use shiny applications, the environment variable USE_SHINY must be set to TRUE (the default) in /config/env_glob.txt. Shiny apps are then enabled through the environment file at /inst/apps/env_shiny.R and will load necessary packages. These are automatically installed if not present the first time the application is launched on any given system. By default, a logging environment is also enabled to assist with metrics and debugging.

Applications are located in the /inst/apps directory and are self-contained in subdirectories by application name. The three that ship with the project are in the "three file" format of global.R, ui.R, and server.R and make use of the API for database communication; they will launch the API server in a background process if it is not already running. To add a live database connection to a new app, simply add the connection object to global.R for that app and develop as normal.

Pay close attention to the settings in the /config/env_glob.txt file as these will determine whether you are using the project in "local" mode or "network" mode. Once the /R/compliance.R file has been sourced, these settings are available in your environment for reference, and applications are available to be launched locally with start_app("X").

• To launch the project solely for access to the web applications listed here, it is recommended to use the following settings and launch them directly from the command line with shiny::runApp(file.path("inst", "apps", "X")) from the project directory (assuming shiny has been installed previously):

INIT_CONNECT	= FALSE
LOGGING_ON	= FALSE
USE_API	= FALSE
API_LOCALHOST	= TRUE
API_PORT	= 8080
INFORMATICS	= FALSE
USE_SHINY	= FALSE

• To launch the project in development mode, it is recommended to use the following settings:

INIT_CONNECT	= TRUE
LOGGING_ON	= TRUE

USE_API	=	TRUE
API_LOCALHOST	=	TRUE
API_PORT	=	8080
INFORMATICS	=	TRUE
USE_SHINY	=	TRUE

These will launch the project in "local" mode enabling, respectively:

- 1. a minimal footprint session with only what is necessary to run the application requested; and
- 2. a live database connection (by default at con), logging, an API hosted on your local machine at port API_PORT, rdkit integration, and shiny apps.

If you want the application to be available on your network, use the following settings and launch the application with shiny::runApp("X", host = "0.0.0.0") and, for consistency, set the port argument to a predetermined port (otherwise a random open port will be used):

```
API_LOCALHOST = FALSE
API_HOST = "host.address"
# host.address must be a resolveable network path to the hosting machine, either an IP
address or a "name.domain" path
```

Settings in other environment resolution files should, in most cases, not be changed.

References

Adusumilli, Ravali and Mallick, Parag. 2017. "Data Conversion with ProteoWizard msConvert." Methods in Molecular Biology (Clifton, N.J.) 1550: 339–68. https://doi.org/10.1007/978-1-4939-6747-6_23.