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Quantum Transport in Matter Interfaces and Correlated Electron systems

QTM measurement toolbox

Documentation on a Python framework for measurements

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${\bf QTM\ Measurement\ toolbox}$

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Introduction

The toolbox presented in this document has been developed in order to perform measurements in Python. The introduction of a measurement toolbox in this language is a natural step that follows from the bachelor programme of Applied Physics, where Python is introduced as main programming language.

Structure of the QTM toolbox

The QTM measurement toolbox consists of various scripts that interact with each other. Here, we clarify the structure of the software.

2.1 Program structure

Let's start with the different folders. In the root directory¹, one finds:

- The file QTM.py. This is the main program that the user executes. The main program sets up connections to all specified GPIB devices and loads the required modules to perform measurements. Depending on the user, the main program may also contain a list of measurement commands (in the LabVIEW software, these were so-called *batch files*).
- The folder functions. This folder contains Python scripts that hold all functions that can be used during the measurements. Some examples are move, sweep and measure. The functions are all stored in a Python file called qtmlab.py.
- The folder instruments. This folder contains definitions for each instrument that can be used for measurements. Essentialy, these files tell Python how to convert our "simple" commands to the actual commands that are sent through GPIB.

2.2 Data flow

The diagram in Fig. 2.1 shows the possible data flows in the QTM measurement toolbox.

As can be seen from the figure, every part of the toolbox can interact with other parts of the toolbox, and the user (via the **IPython console** is able to communicate directly to all modules, as long as the system is initialised².

¹The root directory is the main project folder. All other paths mentioned are relative to this folder.

²That is, all modules and instrument definitions are loaded and communication with instruments has been established.

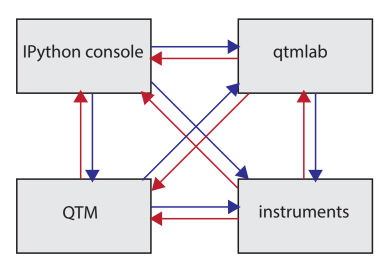


Figure 2.1: Data flow chart for the toolbox. Blue lines represent commands being issued, red lines represent data being returned.

Starting and preparing the toolbox

Before measurements can be performed, the toolbox must be initialised. In this chapter, we walk through the steps necessary to complete this process.

3.1 Launching Spyder

The easiest way to perform measurements is throughout the software **Spyder**. Spyder is an IDE (Integrated Development Environment). The layout is similar to the MATLAB software. Sypder can be found in the application menu of the measurement computer, in the 'Anaconda3' folder.

When launched, a window like Fig. 3.1 appears. The window consists of three smaller windows:

- 1. The **Editor**. The editor (shown at the left side of the program) enables the user to write scripts and edit existing scripts. A script can be executed by pressing the green 'Play' button in the menu bar, or by pressing <F5> on the keyboard.
- 2. The **Variable explorer** is shown at the top right panel. Here, used variables are presented. Note that not all variables (for example, instruments that we define) will be shown here. With the tabs at the bottom of the panel, one can also bring up the **File explorer**, **Help** or the **Profiler**.
- 3. The **IPython console** is the place where one can type code that will be executed when **<Enter>** is pressed. This resembles the 'Command window' of MATLAB.

3.2 Open the toolbox and change the working directory

The next step is to open the toolbox. For this, click the 'Open' button or type <Ctrl>+<0> and browse to the QTM.py file, then open it.

Just as MATLAB, Python has a *working directory*. It can only find functions, modules and files within its installation folder or this specific working directory. Our program is located in a different folder, and although the default working directory can be chosen to be our toolbox folder, it might be

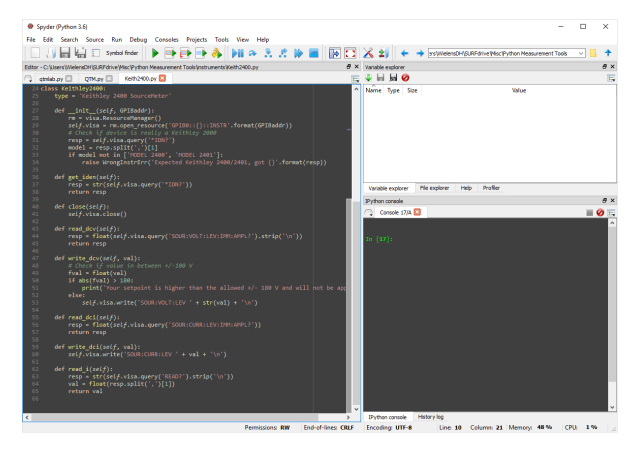


Figure 3.1: The user interface of Sypder. The **editor** is placed at the left side of the window. The right side is divided into two different regions, the **Variable explorer** (or **File explorer**, **Help** or **Profiler** window) and the **IPython console**

that one has to adjust it manually.

To change the working directory, click on the 'Folder' icon next to the textbox that contains the path of the current working directory in the top right corner of the window. Choose the folder that contains the QTM.py file.

3.3 Prepare the toolbox

In this section, we prepare the toolbox by telling it what instruments we will use and which GPIB addresses are used by these instruments. Then, we tell our toolbox what values of instruments should be recorded during measurements. Finally, we run our - just modified - QTM.py file to initialise our system.

3.3.1 Adding instruments

To add instruments, we modify the QTM.py file. In the **Setup** section, we first add all different *types* of instruments that will be used¹.

For example, if we want to use two Keithley 2400 SourceMeters, a Keithley 2000 Multimeter and a LakeShore 332 Temperature controller, we modify the # Import device definitions section as follows:

```
from instruments.Keith2400 import *
from instruments.Keith2000 import *
from instruments.Lake332 import *
```

Importing the modules means that Python adds the modules to its *namespace*, which essentially means that all functions within these modules can be accessed and used from now on.

3.3.2 Connecting to instruments

Now that the definitions for the instruments have been imported, we can bind instruments. For every instrument, we give it a name and then use the corresponding *Class* from the imported module. As argument, we specify the GPIB address. The general syntax is of the form

```
1 <devicename> = <DeviceClass>(<GPIBaddress>)
```

For our example, we would add the following lines of code to the # Connect to devices section:

```
keithBG = Keithley2400(20)
keithTG = Keithley2400(22)
lake = Lake332(12)
```

3.3.3 Define measurement variables

In this step, we tell the toolbox what variables of which devices will be recorded during a measurement. All variables are stored into a single *dictionary*, called meas_dict. The structure of the dictionary must be as follows:

 $^{^{1}}$ This means that if you use 4 sr830 lock-in amplifiers, you only have to import the corresponding instrument module once

```
1 meas_dict = {
2   '<storagename>' : {
3    'dev' = : <devicename>,
4    'var' = '<variable>'
5    },
6    ...
7  }
```

The **<storagename>** will be used in the header of a datafile when measurements are performed and may be chosen by the user. The **<devicename>** should be the exact name of the device as defined in the previous section. Note that there are no quotes around the variable name. The **<variable>** holds the name of the variable that should be read.

To see what variables can be measured, either look further into this manual (chapter 6) or manually open the files that define the instruments (so, in the functions folder). Every function that starts with read_ can be measured.

Note that the prefix read_itself is omitted here!

For our example, we would change the dictionary to:

```
meas_dict = {
           'keithBG.dcv' : {
                   'dev' : kb,
3
                    'var' : 'dcv'
                   },
           'keithBG.i' : {
                    'dev' : kb,
                    'var' : 'i'
                    },
           'keithTG.dcv' : {
10
                    'dev' : kt,
                    'var' : 'dcv'
12
                    },
13
           'keithTG.i' : {
14
                    'dev' : kt,
15
                    'var' : 'i'
16
                    }
17
           'lake.TA' : {
18
                    'dev' : lake,
19
                    'var' : 'temp'
20
21
                    }
22
           }
```

3.4 Start the toolbox

With that, our setup process is complete. Run the QTM.py file by clicking on the Run button (green Play button) or by pressing <F5>.

Executing commands and measurements

With our toolbox ready, we can execute commands to change setpoints, to get the current value of a device, and to perform measurements.

4.1 Individual commands

To run individual commands, head over to the **IPython console**. From the IPython console, you can either directly control devices or issue move, sweep, ... commands.

Some commands (such as readings and measurements) return values, some commands do not. To see whether a command is finished, simply wait until the IPython console presents a new In [<i>] statement.

Typical commands that can be issued - after preparing and executing our QTM.py - from the console are

```
In [1]: keithTG.read_dcv()
2 Out[1]: 0.0

In [2]: keithBG.read_i()
5 Out[2]: -1.94671e-11

In [3]: keithBG.write_dcv(2.1)

In [4]: qtmlab.move(keithBG, 'dcv', 0, 0.5)
```

Note that we can directly interact with devices, but that functions from qtmlab have the module name as prefix for the command. This is because of the way how we imported the modules, which, in turn,

is necessary for the toolbox to work as we intended.

4.2 Batch file

In the LabVIEW software, the possibility existed to execute batch files: files that contain a sequence of commands that will all be executed sequentially. In the Python toolbox, we can write our 'batch commands' in the QTM.py file directly.

In the QTM.py file, you'll find a section called #%% Batch commands. Simply enter all commands that must be executed sequentially in the file and run the code to start your batch file¹!

 $^{^{1}}$ Note that it might be useful to add some print commands to tell yourself what measurement is running / finished.

List of functions

In this section all functions within the qtmlab module will be listed.

5.1 move

The move command can be used to *move* a variable (setpoint) to a specific value at a specified rate.

The general syntax is

```
qtmlab.move(<device>, '<variable>', <setpoint>, <rate>)
```

An example would be

```
1 qtmlab.move(keithBG, 'dcv', 10, 0.5)
```

The function accepts the following arguments:

• <device>: the device identifier. Should be the variable name as defined in QTM.py.

• <variable>: the variable that will be moved. Note that a variable can only be moved if

it contains both a read and write function in the instrument definition.

The <variable> does not need a read or write prefix here!

• <setpoint>: the setpoint that the device will move to. Must be provided as float.

• **<rate>**: the rate at which the device will move to the setpoint.

Must be provided as float.

5.2 measure

The measure command can be used to acquire data of all devices that are specified in the measurement dictionary meas_dict in the QTM.py file.

The general syntax is

```
1 qtmlab.measure()
```

The function will return a NumPy array containing all the values of all variables.

The function does not require any inputs. However, it may look that md must be passed as argument. This is automatically done in the function definitions and when the qtmlab module is imported.

5.3 sweep

The sweep command can be used to measure datapoints while one device is constantly changing its setpoint in between the measurements. The device that is being swept will move to a point, measure data, move to the next point, etc.

The general syntax is

An example would be

```
qtmlab.sweep(keithBG, 'dcv', -10, 10, 0.5, 21, 'Gatesweep.csv', 'Vbg(V)')
```

The function accepts the following arguments:

• <device>: the device identifier. Should be the variable name as defined in QTM.py.

• <variable>: the variable that will be moved. Note that a variable can only be moved if

it contains both a read and write function in the instrument definition.

The <variable> does not need a read or write prefix here!

• <start>: the starting point of the variable that is swept. Must be provided as float.

• <stop>: the end point of the variable that is swept. Must be provided as float.

• <rate>: the rate at which the device will move to the setpoint.

Must be provided as float.

 \bullet $\mbox{ ``npoints'}:$ the number of data points that will be acquired during the sweep.

Must be provided as int.

• <filename >: the filename of the dataset that will be saved. The path can be relative (as

in the example. In that case, the file will be stored in the working directory of Python. The path can also be absolute, i.e. D:\Data\User\Folder\data.csv

for which the data will be stored in the specified path.

ullet <sweepdev> : OPTIONAL The name specified here will serve as the name of the swept

variable in the header of the data file. When not specified, 'sweepdev' will

be used.

List of instrument commands and variables

In this section all instruments and the available commands that can be issued for each instrument will be listed.

6.1 Keithley 2400 SourceMeter

To connect to a Keithley 2400 / 2401 SourceMeter, use

```
from instruments.Keith2400 import *
keithBG = Keithley2400(<GPIBaddress>)
```

The following commands can be issued to a Keithley 2400 / 2401 SourceMeter.

• get_iden(): returns the identification string of the device as a string.

• close(): closes the GPIB connection to the device.

ullet read_dcv(): reads the DC voltage when in $\it Voltage\ source\ mode$ and returns the value

in Volts as float.

• write_dcv(val): sets the DC voltage to val (must be a float) when in Voltage source mode.

Note that the voltage is limited to ± 180 V by the code.

• read_dci(): reads the DC current when in Current source mode and returns the value

in Ampères as float.

• write_dci(val): sets the DC current to value (must be a float)when in Current source mode.

• read_i(): reads the current and returns the value in Volts as float.

6.2 Keithley 2000 MultiMeter

To connect to a Keithley 2000 MultiMeter, use

```
1 from instruments.Keith2000 import *
2 keithR = Keithley2000(<GPIBaddress>)
```

The following commands can be issued to a Keithley 2000 Multimeter.

• get_iden(): returns the identification string of the device as a string.

• close(): closes the GPIB connection to the device.

• read_dcv(): reads the DC voltage and returns the value in Volts as float.

6.3 LakeShore 332 Temperature Controller

To connect to a LakeShore 331 / 332 Temperature Controller, use

```
1 from instruments.Lake332 import *
2 lake = Lake332(<GPIBaddress>)
```

The following commands can be issued to a LakeShore 331 / 332 Temperature Controller.

• get_iden(): returns the identification string of the device as a string.

• close(): closes the GPIB connection to the device.

• read_temp(): returns the current temperature in Kelvin as float.

• write_PID(P, I, D): sets the PID values of the controller to P, I and D.

These values should all be of the type float

• write_setp(val): sets the temperature setpoint in Kelvin as to val (a float).

• write_range(val): changes the setpoint range. Inputs can be of the following form:

['Off', 'off', 0] ['Low', 'low', 1] ['Medium', 'medium', 2] ['High', 'high', 3]

6.4 Stanford Research 830 Lock-In Amplifier

To connect to a Stanford Research 830 Lock-In Amplifier, use

```
1 from instruments.sr830 import *
2 lake = sr830(<GPIBaddress>)
```

The following commands can be issued to a Stanford Research 830 Lock-In Amplifier.

• get_iden(): returns the identification string of the device as a string.

• close(): closes the GPIB connection to the device.

read_x(): returns the X reading as a float.
read_y(): returns the Y reading as a float.
read_r(): returns the R reading as a float.
read_theta(): returns the θ reading as a float.
read_freq(): returns the frequency as a float.

• write_freq(val) : sets the frequency to val (a float).

• read_amp(): returns the amplitude of the Sine Out as float.

• write_amp(val) : sets the Sine Out amplitude to val (a float).

• read_phase(): returns the phase as float.

• write_phase(val) : sets the phase to val (a float).

• read_sens(): returns the sensitivity as int.

• write_sens(val) : sets the sensitivity to val (an int).

For all four DAC outputs, one can use the following commands (here shown for DAC1 - just replace 1 by the number of the output that is used).

• read_dac1(): returns the DAC1 reading as float.

• write_dac1(val): sets the DAC1 output to val (a float).