README

The **SPIKE** program. A collaborative development for a FT-spectroscopy processing program.

Release

This is the version 0.99.2 - January 2019

MANY improvements and corrections have been made, and where available in the devel brach. There are now available in the default release.

Complete history in the release_notes.md file.

What is SPIKE?

SPIKE is a program that allows the processing, the display and the analysis of data-sets obtained from various Fourier-Transform spectroscopies. The name stands for **S**pectrometry **P**rocessing Innovative **KE**rnel.

It allows the processing of **1D** and **2D** FT spectroscopies, implementing Real, Complex and HyperComplex n-dimensionnal Fourier Transform, as well as many other functionalities.

It is written in python (tested in python 2.7 and 3.5) and can be used as a set of tools, using for instance jupyter notebook as an interactive front-end.

To our knowledge, it is the first program freely available allowing the processing, display and analysis of 2D-FT-ICR (Fourier Transform Ion Cyclotron Resonance).

It is still in very active development. Many features are missing, and many other while present, are not fully fixed. However, considering the amount of efforts already present in this code, we decided to make it available. We believe that even in this partial development stage, this program might prove useful for certain usages.

Documentation

Documentation is way behind the code, and certain parts are presenteing obsolete features. This is the main reason for geing 0.99 rather than 1.00 version!

You can find the Very preliminary documentation here

On the other hand, the in-line python documentation is rather complete and up to date.

Citing SPIKE

If you happen to use SPIKE successfully please cite it, and refer to this site, as well as the following possible references:

- first publication of the program itself rejected from Anal. Chem. Reviewer 1 said "too much NMR", Reviewer 2 said "too much MS", !!
 - 1. Chiron L., Coutouly M-A., Starck J-P., Rolando C., Delsuc M-A. SPIKE a Processing Software dedicated to Fourier Spectroscopies https://arxiv.org/abs/1608.06777 (2016)
- first version of the python set-up on which the current SPIKE is partially based
 - Tramesel, D., Catherinot, V. & Delsuc, M.-A. Modeling of NMR processing, toward efficient unattended processing of NMR experiments. *J Magn Reson* 188, 56–67 (2007).
- first version of the 2D FT-ICR-MS processing
 - 3. van Agthoven, M. A., Chiron, L., Coutouly, M.-A., Delsuc, M.-A. & Rolando, C. Two-Dimensional ECD FT-ICR Mass Spectrometry of Peptides and Glycopeptides. *Anal Chem* **84**, 5589–5595 (2012).
- presentation of the automation possibilities in NMR
 - Margueritte, L., Markov, P., Chiron, L., Starck, J.-P., Vonthron Sénécheau, C., Bourjot, M., & Delsuc, M.-A. (2018). Automatic differential analysis of NMR experiments in complex samples. Magn. Reson. Chem., 80(5), 1387. http://doi.org/10.1002/mrc.4683

ref 1) is a general purpose reference, the other ones are more specific.

SPIKE proposes the following features

FT analysis of 1D data-sets

• apodisation, phasing, modulus, ...

Analysis of 2D data-sets

- phase or amplitude modulation
- complex or hyper-complex algebra

Robust processing

- no limit in data-set size
- parallel processing of the heaviest processing
 - on multi-core desktop using standard python ressources
 - on large clusters, using **MPI** library

High-Level features

- noise reduction (filtering, Linear-Prediction, Cadzow, urQRd, sane, ...)
- automatic or manual baseline correction
- 1D and 2D Peak-Picker

Plugin architecture

- allow easy extension of the core program
- reduces cross dependences

Complete spectral display using matplotlib

- zoom, available in several units (depending on the spectroscopy : seconds, Hz, ppm, m/z, etc...)
- store to png or pdf

For the moment, SPIKE handles the following Spectroscopies

NMR

- 1D and 2D are fully supported
- no nD yet

FT-ICR

• 1D and 2D are fully supported

Orbitrap

• 1D only (!)

other spectroscopies are being considered

Files can be imported from

- NMR:
 - Bruker Topspin
 - NMRNoteBook
 - NPK Gifa
- FT-ICR:
 - Bruker Apex

- Bruker Solarix
- Orbitrap:
 - Thermofisher raw data
- any data in memory in a Numpy buffer.

Usage

import spike

As a processing library

dd.pp(threshold=1E7)

SPIKE is primary meant for being used as a library, code can as simple as :

```
from spike.File import Solarix

dd = Solarix.Import_1D('FTICR-Files/ESI_pos_Ubiquitin_000006.d') # Import create a basic St.

dd.hamming().zf(2).rfft().modulus() # we have a simple piped processing scheme
    # here doing apodisation - zerofilling (doubling the size) - FT and modulus.

dd.unit = "m/z"

dd.display(zoom=(500,2000)) # display the spectrum for m/z ranging from 500 to 2000
```

peak-pick the spectrum in this range

insure spike is in your PYTHONPATH

```
dd.centroid()  # compute centroids

dd.display(zoom=(856.5, 858.5))  # and zoom on the isotopic peak
dd.display_peaks(zoom=(856.5, 858.5), peak_label=True)
```

For the moment, SPIKE does not provide a installation script. If you downloaded spike in a given directory, insure this directory is in your PYTHONPATH. To do so, either modify the \$PYTHONPATH environment variable, or add the following lines in the scripts that use SPIKE:

```
import sys
sys.path.append('the_dir_where_you_put_spike_distrib')
```

interactive mode

SPIKE allows to process datasets interactively from an jupyter (IPython) prompt, and is perfectly working in jupyter notebook or even jupyter lab

- Look at the examples files (eg_*.py and *.ipynb) for examples and some documentation. (* not fully up to data *)
- display is performed using the Matplotlib library.
- large 2D-FT-ICR are handled in batch using the processing.py batch program, controlled by parameter file called *.mscf

- The batch mode supports multiprocessing, both with MPI and natively on multi-core machines (still in-progress)
- large 2D-FT-ICR are stored in a hierarchical format, easyly displayed with an interactive program.
- data-sets are handled in the HDF5 standard file-format, which allows virtually unlimited file size ($tested\ up\ to\ 500\ Gb$).

running stand-alone programs

processing.py and visu2D.py are two stand alone programs, written on the top of SPIKE. - processing.py allowing the efficient processing of FT-ICR 2D datasets, with no limit on the size of the final file Produces multi-resolution files - visu2D.py is an interactive tool for visualizing 2D FT-ICR multi-resolution files

```
syntax:
```

```
python -m spike.processing param_file.mscf
or
python -m spike.visu2D param_file.mscf
typically, you want to add
python
import sys
sys.path.append('the dir where you put spike distrib')
```

to the header of these scripts, and launch them from the directory which contains SPIKE the distribution.

A more complete documentation is available here.

How do I get SPIKE?

SPIKE is written in pure Python, and relies on several external libraries. It is compatible and fully tested with both python 2.7 and python 3.5

It requires the following non-standard Python libraries :

- Numpy
- Scipy
- Matplotlib
- HDF5 / Pytables
- $\bullet~$ Qt / PySide optional used by visu2D
- MPI / mpi4py optionnal used for parallel processing of large FTICR 2D files

It has been successfully tested in the Enthought and Anaconda distributions.

To get it, you can simply - insall the above python distributions - download the latest stable version here: https://bitbucket.org/delsuc/spike/downloads - or hg clone the devel branch and keep it up-to-date

Origins

SPIKE is originated from the ** Gifa ** program, developed by M-A Delsuc and others in FORTRAN 77 since the late eighties. Gifa has known several mutations, and finally ended as a partial rewrite called **NPK**. The NPK program is based on some of the original FORTRAN code, wrapped in Java and Python, which allows to control all the program possibilities from the Python level. NPK is purely a computing kernel, with no graphical possibilities, and has been used as a kernel embedded in the commercial program NMRNoteBook, commercialized by NMRTEC.

However, NPK was showing many weaknesses, mostly due to the 32bits organization, and a poor file format. So, when a strong scientific environment became available in Python, a rewrite in pure Python was undertaken. To this initial project, called NPK-V2, many new functionalities were added, and mostly the capability to work in other spectroscopies than NMR.

At some point in 2014, we chose to fork NPK-V2 to SPIKE, and make it public.

Developing for SPIKE

SPIKE is an open-source program, this means that external contributions are welcomed. If you believe your improvement is useful for other people, please submit a pull request. Note that pull request should be associated to the devel branch. This branch is devoted to new features not fully tested yet and still susceptible of changes, while the default branch is meant for stable code.

plugins

If you consider adding some new feature, it is probably a good idea to implement it as a plugin. The code contains already quite a few plugins, some are quite sophisticated - see Peaks.py for instance which implements a 1D and 2D peak picker, as well as a centroid evaluation and a full listing capability.

You can check also fastclean.py for a very simple plugin, or wavelet.py for a plugin relying on an external library which has to be installed.

Some Good Practice

- Spike contains many tools, most of the basic function for data interaction are found in the NPKData.py master file; utilities are also scattered in the util module. Use then, life will be easier for the users.
- Please write tests, even for the plugins! We use standard python unittest, so nothing fancy. All the tests are run automatically every night (code is Tests.py), so it will detect rapidly all potential problem.
- push your pull requests to the devel branch default is for the stable releases.

Organisation of the Code

The main program is NPKData.py, which defines NPKData object on which everything is built.

Spectroscopies are defined in the FTICR.py and Orbitrap.py code, which sub class NPKData. It is prototyped as an NMR data-set. This set-up is temporary.

Many programs contain routines tests (in an object unittest) that also serve as an example of use. The code goes through extensive tests daily, using the unittest Python library. However, many tests rely on a set of tests data-sets which is more than 1Go large, and not distributed here.

Main programs:

a small description of the files:

- NPKData.py the main library, allows all processing for NMR experiments (1D, 2D and 3D) to be used as a library, in a stand-alone program or in IPvthon interactive session
- FTICR.py an extension of NPKData for processing FT-ICR datasets (1D and 2D)
- Orbitrap.py an extension of NPKData for processing Orbitrap datasets (1D)
- processing.py a stand alone program, written on the top of FTICR.py, allowing the efficient processing of FT-ICR 2D datasets, with no limit on the size of the final file Produces multi-resolution files syntax:

python -m spike.processing param_file.mscf

 $\bullet\,$ visu2D.py an interactive tool for visualizing 2D FT-ICR multi-resolution files

python -m spike.visu2D param_file.mscf

Directories

- Algo contains algorithms to process data-sets (MaxEnt, Laplace, etc...) not everything active!
- Display
 a small utility to choose either for regular Matplotlib display of fake
 no-effect display (for tests)
- File Importers for various file format for spectrometry, as well as the HDF5 SPIKE native format.
- plugins

 Tools automatically plugged in NPK kernel : display utilities, urQRd algorithm and various other tools.
- Miscellaneous "en vrac"
- *Visu* utilities for the Visu2D program
- *util* set of low-level tools used all over in the code
- v1
 a library implementing a partial compatibility with the NPKV V1 program
- • $SPIKE_usage_eg$ example of Python programs using the various libraries available
- example of configuration files
 - process eg.mscf
 - test.mscf
- \bullet and various utilities
 - NPKConfigParser.py reads .mscf files
 - NPKError.py generates error msg
 - QC.pyQuality Check
 - Tests.py runs all tests
 - dev_setup.pyrolls a new version

- version.py defines version number
- __init__.pydefines library
- repylint
- To_Do_list.txt
- QC.txt
- Release.txt

Authors and Licence

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