Data and text mining

Advance Access publication February 19, 2013

# IRootLab: a free and open-source MATLAB toolbox for vibrational biospectroscopy data analysis

Júlio Trevisan<sup>1,2,\*</sup>, Plamen P. Angelov<sup>1,2</sup>, Andrew D. Scott<sup>3</sup>, Paul L. Carmichael<sup>3</sup> and Francis L. Martin<sup>2</sup>

<sup>1</sup>School of Computing and Communications, Infolab21, Lancaster University, Lancaster LA1 4WA, UK, <sup>2</sup>Centre for Biophotonics, Lancaster Environment Centre, Lancaster University, Lancaster LA1 4YQ, UK and <sup>3</sup>Safety and Environmental Assurance Centre, Unilever Colworth Science Park, Bedfordshire MK44 1LQ, UK Associate Editor: Jonathan Wren

#### **ABSTRACT**

Summary: IRootLab is a free and open-source MATLAB toolbox for vibrational biospectroscopy (VBS) data analysis. It offers an object-oriented programming class library, graphical user interfaces (GUIs) and automatic MATLAB code generation. The class library contains a large number of methods, concepts and visualizations for VBS data analysis, some of which are introduced in the toolbox. The GUIs provide an interface to the class library, including a module to merge several spectral files into a dataset. Automatic code allows developers to guickly write VBS data analysis scripts and is a unique resource among tools for VBS. Documentation includes a manual, tutorials, Doxygen-generated reference and a demonstration showcase. IRootLab can handle some of the most popular file formats used in VBS.

License: GNU-LGPL.

Availability: Official website: http://irootlab.googlecode.com/.

Contact: juliotrevisan@gmail.com

Supplementary information: Supplementary data are available at Bioinformatics online.

Received on December 3, 2012; revised on February 8, 2013; accepted on February 13, 2013

# 1 INTRODUCTION

Vibrational biospectroscopy (VBS) is the application of infrared and Raman spectroscopy to biological studies and biomedical applications (Martin et al., 2010). In the past decades, vibrational spectroscopy data processing was developed on the grounds of signal processing, statistics and pattern recognition, resulting in a large number of computational methods that are routinely applied in the field. However, the field needs further development and establishment of data analysis strategies, partially owing to a continuous push from instrumental advancements, but also motivated by challenges in pre-processing, biomarker extraction and standardization of inter-experimental datasets, and inspired by the increasing power of computers (Trevisan et al., 2012).

VBS data analysis is largely carried out by commercial softwares such as CytoSpec (www.cytospec.com), Neurodeveloper (www.neurodeveloper.com), Unscrambler (www.camo.com), Pirouette (www.infometrix.com), OPUS (www.bruker.com) and

(www.renishaw.com). Popularly used commercial MATLAB toolboxes include the Neural Network and the Bioinformatics Toolbox. Others are free to use, but closed-source, such as PRTools (Duin et al., 2007) and GA ORS (Nikulin et al., 1998). While these are all high-quality software, the use of closed-source software contradicts the multidisciplinary aspect of the field, as researchers looking for new analysis strategies remain dependant from the software manufacturers. On the other hand, open-source softwares available are largely restricted to a given task, such as LibSVM (Chang and Lin, 2011), Functional Data Analysis toolbox (Ramsay et al., 2009) and various other resources available online.

IRootLab is a free/libre and open-source (FOSS) MATLAB toolbox created to fulfil the need for FOSS software encompassing every stage of VBS data analysis. It is a highly modular software that was created based on existing theory of data analysis (Alpaydin, 2004; Bishop, 2006; Duda et al., 2001; Guyon et al., 2006; Hastie et al., 2007; Kuncheva, 2004), and as such, applied to VBS (Griffiths and Haseth, 2007; Somorjai, 2009), bringing together families of methods such as outlier removal, pre-processing, feature extraction, feature selection, feature construction, classification, classifier aggregation, clustering; concepts such as cross-validation, random sub-sampling, peak detection, biomarker identification; a set of unique visualization options (Supplementary Fig. S1) and a MATLAB code generator. The basic modular design is simple, which makes IRootLab a flexible and intuitive resource for vibrational spectroscopy data analysis developers.

# **2 CLASS LIBRARY**

The majority of IRootLab code lines are attributed to its hierarchically organized object-oriented programming (OOP) classes. The basic framework is constituted of three branches: datasets, blocks and logs. A dataset is an object representing point spectra (with their respective classes, patient names, etc.), image maps, clustering data or classifier estimations. A block is the basis of data manipulation. It is an object that allows mainly two operations: training and using. Training involves the use of training data to modify a block (according to its specific idiosyncrasy). Using consists of inputting data into a block to get various results, including output data, estimation data, figures and HTML reports. Finally, logs represent the output of special blocks called

<sup>\*</sup>To whom correspondence should be addressed.

analysis sessions, which perform complex analyses and generate output that does not fit into the usual dataset class.

Additionally, there are a few complementary branches, from which the most important are *sub-dataset generation specs* (SGS), *feature sub-set grader* (FSG) and *peak detector*. SGS is an abstraction of the concept of random sub-sampling (e.g. cross-validation, repeated random sub-sampling); FSG is used to evaluate sub-sets of features in feature selection tasks, freeing the feature selection algorithm from having to implement its own evaluation; and *peak detection* (Coombes *et al.*, 2003) is used in visualizations and biomarker identification.

Blocks can be created and combined recursively (e.g. a special block called 'block cascade' is composed of other blocks, which in turn can be used as a component of a higher level block cascade) to build analyses as complex as desired.

Many complex operations are already offered as sub-classes of a special class of block called *analysis session*, including grid search optimization (Hsu *et al.*, 2010), repeated train-test using an SGS object to obtain performance estimation of a classifier and cross-calculation of Linear Discriminant Analysis (LDA) scores (Riding *et al.*, 2012).

### 3 GRAPHICAL USER INTERFACES

Figure 1 shows the interaction of IRootLab graphical user interfaces (GUIs) with the class library and their surrounding environment. IRootLab offers a flexible and intuitive GUI called *objtool* that is an interface to the class library. *Objtool* allows datasets and other objects present in the MATLAB workspace to be browsed and manipulated using new or existing blocks. *Objtool* can handle three types of TXT files, OPUS image maps and a native IRootLab MAT format.

Mergetool is another GUI that allows a collection of single-spectrum files to be merged together to form a dataset. Currently, mergetool can import three different types of

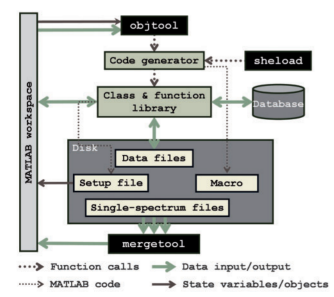


Fig. 1. IRootLab GUIs (objtool, mergetool, sheload) and their surrounding environment

single-spectrum files: Pirouette.dat, OPUS single-spectrum files and Wire TXT files. Finally, the *SHEload* GUI accesses an online MySQL database that is part of a chemical database project (Trevisan *et al.*, 2010), importing datasets from there.

## 4 CODE GENERATION

One of the major features of IRootLab is the ability of *objtool* to generate MATLAB code (similar to macro recording in Microsoft Word). This is a highly efficient resource for MATLAB scripting. In fact, a significant part of IRootLab was written using this resource. Code generation also keeps developers from having to consult the library documentation too often.

#### 5 DOCUMENTATION

IRootLab has a clear documentation project. The official documentation (available at http://bioph.lancs.ac.uk/irootlabdoc) is generated directly from the source code using Doxygen software. Apart from this, IRootLab has a manual and a series of tutorials (available at downloads area of the official website) that together provide a friendly start for new users. The official reference can be accessed from MATLAB in two different occasions: from the command line, by using the *help2* command, and from the GUIs, by pressing the F1 key (context-sensitive help). Additionally, a demonstration showcase was prepared (which can be opened by typing the command *browse\_demos* at MATLAB command line), and a number of sample datasets are shipped with the toolbox.

# 6 CONCLUSION

Although the most valuable contributions of IRootLab are its framework and its code generation, there are currently more than 200 implemented OOP classes representing methods, algorithms, concepts and visualizations. One of the advantages of FOSS software is that any user can potentially expand the toolbox to include the methods necessary to their own analysis, should these methods not be already present, and then share the additions, creating a synergic effort among the scientific community. Expansion is facilitated by IRootLab modular structure, which allows developers to target or inherit a specific OOP class. IRootLab is a unique and innovative contribution to the interdisciplinary research field of VBS, providing researchers with a valuable tool for the development of VBS data analysis.

### **ACKNOWLEDGEMENT**

IRootLab uses a number of third-party packages, whose developers we would like to thank: Chih-Chung Chang, Chih-Jen Lin, Jacob Filik, James O. Ramsay, Michael Nasse, Paul Bassan and Yannick Maret. We would also like to thank our colleagues Abdullah A. Ahmadzai, Imran I. Patel, Jemma G. Kerns and Valon Llabjani for their participation in the conception.

Funding: This work was funded by Unilever as 'part of Unilever's ongoing effort to develop novel ways of delivering consumer

safety'. We would also like to thank Rosemere Cancer Foundation for their support over the years.

Conflict of Interest: none declared.

# **REFERENCES**

- Alpaydin, E. (2004) Introduction to Machine Learning. MIT Press, Cambridge, MA, USA.
- Bishop, C.M. (2006) Pattern Recognition and Machine Learning. Springer, Cambridge, UK.
- Chang, C.-C. and Lin, C.-J. (2011) LIBSVM: a library for support vector machines. ACM TIST, 2, Article no. 27.
- Coombes, K.R. et al. (2003) Quality control and peak finding for proteomics data collected from nipple aspirate fluid by surface-enhanced laser desorption and ionization. Clin. Chem., 49, 1615–1623.
- Duda,R.O. et al. (2001) Pattern Classification. 2nd edn. John Wiley & Sons, New York.
- Duin,R.P.W. et al. (2007) PRTools4—a MATLAB toolbox for pattern recognition. Pattern Recognit., 1–61.
- Griffiths, P.R. and Haseth, J.A. (2007) Fourier Transform Infrared Spectroscopy. 2nd edn. Wiley, Hoboken, NJ, USA.
- Guyon, I. et al. (2006) Feature Extraction—Foundations and Applications. Springer, New York.

- Hastie, T. et al. (2007) The Elements of Statistical Learning. 2nd edn. Springer, New York.
- Hsu,C. et al. (2010) A practical guide to support vector classification. Bioinformatics, 1, 1–16.
- Kuncheva, L.I. (2004) Combining Pattern Classifiers: Methods and Algorithms. Wiley, Hoboken, NJ, USA.
- Martin,F.L. et al. (2010) Distinguishing cell types or populations based on the computational analysis of their infrared spectra. Nat. Prot., 5, 1748–1760.
- Nikulin, A.E. et al. (1998) Near-optimal region selection for feature space reduction: novel preprocessing methods for classifying MR spectra. NMR Biomed., 11, 209–16.
- Ramsay, J. et al. (2009) Functional Data Analysis with R and MATLAB. Springer, New York.
- Riding, M.J. et al. (2012) Concentration-dependent effects of carbon nanoparticles in gram-negative bacteria determined by infrared spectroscopy with multivariate analysis. Environ. Pollut., 163C, 226–234.
- Somorjai, R.L. (2009) Creating robust, reliable, clinically relevant classifiers from spectroscopic data. *Biophys. Rev.*, 1, 201–211.
- Trevisan, J. et al. (2010) Syrian hamster embryo (SHE) assay (pH 6.7) coupled with infrared spectroscopy and chemometrics towards toxicological assessment. Analyst, 135, 3266–3272.
- Trevisan, J. et al. (2012) Extracting biological information with computational analysis of Fourier-transform infrared (FTIR) biospectroscopy datasets: current practices to future perspectives. Analyst, 137, 3202–3215.