# MSnbase input/output capabilities

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#### Abstract

This vignette describes MSnbase's input and output capabilites.

Keywords: Mass Spectrometry (MS), proteomics, infrastructure, IO.

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#### 1 Overview

MSnbase's aims are to facilitate the reproducible analysis of mass spectrometry data within the R environment, from raw data import and processing, feature quantification, quantification and statistical analysis of the results. Data import functions for several formats are provided and intermediate or final results can also be saved or exported. These capabilites are presented below.

# 2 Data input

Raw data Data stored in one of the published XML-based formats. i.e. mzXML (Pedrioli et al., 2004), mzData (Orchard et al., 2007) or mzML (Martens et al., 2010), can be imported with the readMSData method, which makes use of the mzR package to create MSnExp objects. The files can be in profile or centroided mode. See ?readMSData for details.

**Peak lists** Peak lists in the mgf format<sup>1</sup> can be imported using the read-MgfData. In this case, the peak data has generally been pre-processed by other software. See ?readMgfData for details.

Quantitation data Third party software can be used to generate quantitative data and exported as a spreadsheet (generally comma or tab separated format). This data as well as any additional metadata can be imported with the readMSnSet function. See ?readMSnSet for details.

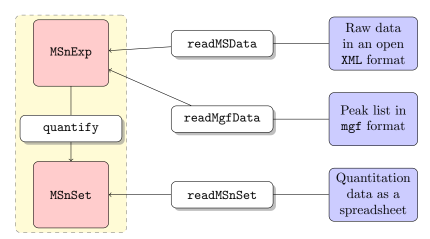


Figure 1: Illustration of MSnbase input capabilities. The white and red boxes represent R functions/methods and objects respectively. The blue boxes represent different disk storage formats.

# 3 Data output

RData files R objects can most easily be stored on disk with the save function. It creates compressed binary images of the data representation that can later be read back from the file with the load function.

Peak lists MSnExp instances as well as individual spectra can be written as mgf files with the writeMgfData method. Note that the metadata in the original R object can not be included in the file. See ?writeMgfData for details.

Quantitation data Quantitation data can be exported to spreadsheet files with the write.exprs method. Feature metadata can be appended to the feature intensity values. See ?writeMgfData for details.

<sup>&</sup>lt;sup>1</sup>http://www.matrixscience.com/help/data\_file\_help.html#GEN

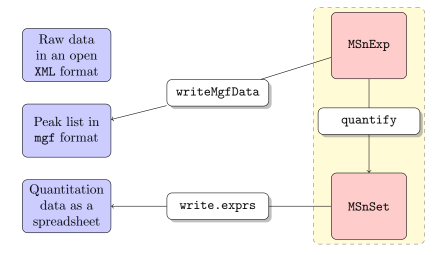


Figure 2: Illustration of MSnbase output capabilities. The white and red boxes represent R functions/methods and objects respectively. The blue boxes represent different disk storage formats.

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