# The ten minute guide to mzTab

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## Introduction

The purpose of this guide is to give a quick introduction on how to use mzTab efficiently. It is targeted at both, developers and end-users alike. This guide is not intended to give a complete and detailed overview of mzTab but should only be a quick and easy to understand introduction. The complete format specification as well as example files can be found at <http://mztab.googlecode.com>.

## Basic structure

mzTab files can have four sections: The metadata section, the protein section, the peptide section, and the small molecule section (see Figure 1). All of these sections are optional and may not be present in every file.

Figure : Basic structure of an mzTab file.

All lines in an mzTab file start with a three letter code to identify the information held by the line:

**MTD** for metadata

**PRH** for the protein table header line (the column labels)

**PRT** for rows of the protein table

**PEH** for the peptide table header line (the column labels)

**PEP** for rows of the peptide table

**SMH** for small molecule table header line

**SML** for rows of the small molecule table

**COM** for comment lines

*For developers:*

mzTab is a tab separated file format. The three letter codes must be separated by a tab from the next field. Also, field names and values in the metadata section are separated by tabs as are the columns in the table based sections.

## Units in mzTab

Every identification and metadata field is assigned to a unit. A unit is only an identifier that groups these pieces of information together. The only limitation units have is that a protein must be unambiguously identified by its protein accession in the same unit. Units are not reported in a separate table or section but only exit through their Unit\_IDs, which are referenced by every metadata field and identification. One mzTab file can contain multiple units that can represent different concepts depending on the experimental design and the granularity needed by the data producer to communicate the results. Some examples:

- The results from one technical replicate (one LC-MS run) or the summary value after all replicates have been combined. Technical replicates can be explicitly reported in mzTab and grouped together by using special “Unit\_IDs” in case this level of detail is required. To report the results from an experimental setup (labelled as “EXP\_1”) containing the data from two replicates as well as the final combined results the resulting mzTab file would contain three units: the replicates as “EXP\_1\_rep[1]” and “EXP\_1\_rep[2]”, and the final results as “EXP\_1”.

- The summary results of all experiments from one dataset submitted to a proteomics repository. For example, the example file “PXD000002 submission” represents the final results from one single dataset but it contains three units: “PRIDE\_22142”, “PRIDE\_22143”, and “PRIDE\_22144”. Each of these represents the results of the corresponding PRIDE experiment (equivalent to one MS run). Thereby, a researcher can easily get an overview of the results of a whole submission by only looking at a single file. Another example of one unit in this context could be one complete PeptideAtlas build.

COM Example showing the usage of UNIT IDs  
MTD EXP\_1-title The first experiment in the file  
…  
MTD EXP\_2-title The second experiment in the file  
…  
PRH accession unit\_id description taxid species …  
PRT P02042 EXP\_1 Hemoglobin subunit delta 9606 Homo sapie…  
PRT P02042 EXP\_2 Hemoglobin subunit delta 9606 Homo sapie…  
  
PEH sequence accession unit\_id unique …  
PEP ABC P02042 EXP\_1 0 …  
PEP CDE P02042 EXP\_1 1 …  
PEP ABC P02042 EXP\_2 0 …  
PEP FGH P02042 EXP\_2 0 …

*For developers:*

Unit\_IDs are freely generated by the software that generated the mzTab file and should somehow sensibly identify the, for example, experiment. If mzTab files are generated by local software tools, these UNIT\_IDs can be any sensible identifier for the, for example, experiment. Unit\_IDs must only contain the following characters: ‘A’-‘Z’, ‘a’-‘z’, ‘0’-‘9’, and ‘\_’.

## Metadata in mzTab

The metadata section in mzTab files contains information about the units and consists of key - value pairs separated by a tab. The name of every field contains the Unit\_ID. Thereby, every field can be attributed to a unit. A complete list of available fields can be found in the specification document.

COM Example showing the use of the metadata section  
MTD EXP\_1-title The unit’s / experiment’s title  
MTD EXP\_1-description This is just an example. No description needed...  
MTD EXP\_1-instrument[1]-source [MS, MS:1000073, ESI,]  
MTD EXP\_1-instrument[1]-analyzer [MS, MS:1000291, linear ion trap,]  
MTD EXP\_1-instrument[1]-detector [MS, MS:1000253, electron multiplier,]  
MTD EXP\_1-software[1] [MS, MS:1001207, Mascot, 2.3]

MTD EXP\_1-software[2] [MS, MS:1001561, Scaffold, 1.0]  
MTD EXP\_1-false-discovery-rate [MS, MS:1001364, pep:global FDR, 0.01]  
MTD EXP\_1-contact[1]-name James D. Watson  
MTD EXP\_1-contact[1]-affiliation Cambridge University, UK  
MTD EXP\_1-contact[1]-email watson@cam.ac.uk

## Proteins in mzTab

Protein identifications are reported in the protein section. The protein section is table based. The table header is identified by the prefix “PRH”, entries in the protein table are identified through “PRT”. The protein section must only be present once but can contain identifications from multiple units. Columns are separated by a tab.

COM Example of the protein section. Other sections are omitted  
PRH accession unit\_id description taxid species database …  
PRT P12345 EXP\_1 mAspAT 9986 Rabbit UniProtKB/SwissProt …  
PRT P02042 EXP\_2 Hemoglobin 9606 Human UniProtKB/SwissProt …

All columns in the protein section are mandatory, except the quantitative and optional columns. The full list of columns can be found in the specification document.

## Peptides in mzTab

The peptide section is similar to the protein section. It is table based, columns are separated by a tab and all columns are mandatory apart from quantitative and optional columns. The header of the peptide table is indicated by “PEH”, and entries in the table by “PEP”. The peptide section must also be present only once but may contain identifications from multiple units. The full list of columns can be found in the specification document.

COM Example of the peptide section. Other sections are omitted.  
PEH sequence accession unit\_id unique … search\_engine …  
PEP ABC P12345 EXP\_1 0 … [MS,MS:1001207,Mascot,] …  
PEP ABC P12345 EXP\_2 0 … [MS,MS:1001208,Sequest,] …

## Small Molecules in mzTab

The small molecule section is also a table based section (same rules apply). Small molecules are identified through an “identifier” in mzTab. This identifier can be any text that sensibly identifies the given small molecule in the given field of research. These identifiers should generally be entries in compound databases used in the respective field (for example, Human Metabolome Database entries, ChEBI identifiers, PubChem IDs or LIPID MAPS IDs). Apart from this identifier, small molecules can be assigned a chemical formula, SMILES and/or InChi identifier, a human readable description, a precursor *m/z* value, a charge state, retention time(s), a species, source database and search engine including score. We are aware, that these fields are not applicable to all fields of metabolomics, but we believe that they represent a sensible selection. A more detailed list of possible columns can be found in the specification document.

COM Example of the small molecule section. Other sections are omitted.  
SMH identifier unit\_id chemical\_formula description mass\_to\_charge charge …  
SML TG54:0 EXP\_1 H110O6C57 - 949.892 1

## Missing values

The table-based sections (protein, peptide, and small molecule) must not contain empty cells. In case a given property is not available for an entry “NA” should be reported.

Any calculation that results in “not a number” (NaN type) must also be reported using “NA”. If ratios are included and the denominator is zero, the “INF” value must be given. In some cases, there is ambiguity with respect to these cases: e.g. in spectral counting if no peptide-spectrum matches are observed for a given protein, it is open for debate as to whether its abundance is zero or missing (“NA”).

## Reliability score

All protein, peptide and small molecule identifications reported in an mzTab file should be assigned a reliability score (column “reliability” in all tables). The idea is to provide a way for researcher and/or MS proteomics or metabolomics repositories or data producers to score the reported identifications based on their own criteria. This score is completely resource-dependent and must not be seen as a comparable score between mzTab files generated from different resources. The criteria used to generate this score should be documented by the data providers. If this information is not provided by the producers of mzTab files, “NA“ must be provided as the value for each of the protein, peptide or small molecule identifications.

The reliability must be an integer between 1-3 and should be interpreted as follows:

1: high reliability

2: medium reliability

3: poor reliability

The idea behind this score is to mimic the general concept of “resource based trust”. For example, if one resource reports identifications with a given reliability this would be interpreted differently as an identification reported from another resource. If resources now report their reliabilities using this metric and document how this metric is generated, a user can base his own interpretation of the results based on his trust in the resource. Furthermore, approaches to make various, for example search engine scores comparable have failed so far. To prevent the notion that the reported scores represent comparable probabilities this very abstract metric was chosen.

## Quantitative Data

There are multiple quantification techniques available for MS-based experiments that often result in slightly different types of data. mzTab was not designed to capture any of these specific differences. The goal for mzTab was to provide a generic view on quantitative MS-based identification data that is applicable to as many different quantitation methods as possible.

Quantitative technologies generally result in some kind of abundance measurement of the identified analyte. Several of the available techniques furthermore allow/require multiple similar samples to be multiplexed and analyzed in a single MS run. When several biological samples are multiplexed these samples are referred to as “subsamples” in mzTab. Subsamples must furthermore be linked to the used labels in the metadata section of the mzTab file (see example below). In case a quantification method is used that does not lead to multiplexed biological samples, the generated quantification values are reported as subsample 1. Detailed information about how to report subsamples in the metadata section can be found in the specification document.

COM The following example shows how two different quantitative experiments  
COM can be reported in one mzTab file. Not all labels are shown  
…  
MTD EXP\_1-quantification\_method [MS,MS:1001837,iTraq,]  
MTD EXP\_1-sub[1]-description Healthy human liver tissue  
MTD EXP\_1-sub[1]-quantification\_reagent [PRIDE,PRIDE:0000114,iTRAQ reagent 114,]  
MTD EXP\_1-sub[2]-description Human hepatocellular carcinoma sample.  
MTD EXP\_1-sub[2]-quantification\_reagent [PRIDE,PRIDE:0000115,iTRAQ reagent 115,]  
…  
MTD EXP\_2-quantification\_method [MS,MS:100999,SILAC,]  
MTD EXP\_2-sub[1]-description Healthy rat liver tissue  
MTD EXP\_2-sub[1]-quantification\_reagent [PRIDE,PRIDE:0000325,SILAC heavy,]  
MTD EXP\_2-sub[2]-description Intoxicated rat liver.  
MTD EXP\_2-sub[2]-quantification\_reagent [PRIDE,PRIDE:0000326,SILAC light,]  
…  
PRH accession unit\_id … protein\_abundance\_sub[1] … protein\_abundance\_sub[2] …  
PRT P12345 EXP\_1 … 1 … 0.82749  
PRT P15151 EXP\_2 … 2.42114 … 1  
…

MS2 spectral counting-based approaches can be reported using optional columns in the peptide table as well as the protein table as they only result in one single value per analyte. In case the approach used also generates standard deviation and standard errors the quantification results may also be reported using the subsample 1 columns. MS label-free quantification techniques do not require any additional support in mzTab as they simply need to report abundance values per sample in a straight-forward manner. CV parameter accessions may be used as optional column names following the following format: opt\_cv\_{accession}.

COM Example showing how emPAI values are reported in an additional column using  
COM MS CV parameter “emPAI value” (MS:1001905)  
…  
PRH accession … opt\_cv\_MS:1001905  
PRT P12345 … 0.658

## Protein Inference

There are multiple approaches to report protein inference. mzTab is designed to only hold experimental results which in proteomics experiments can be very complex. At the same time, for down-stream statistical analysis there is a need to simplify this problem. It is not possible to model detailed protein inference data without a significant level of complexity at the file format level. Therefore, it was decided to “mention” the protein inference problem in mzTab files but not provide detailed information on how it was resolved. Protein entries in mzTab files contain the field “ambiguity\_members”. The protein accessions listed in this field should identify proteins that could also be identified through the same (sub-)set of peptides but were not chosen as the primary identification. The members of the ambiguity group are not reported in the peptide table for the respective unit.

COM In the following example only one peptide was identified that can be attributed to  
COM multiple proteins. The choice which one to pick as primary accession depends on COM the resource generating the mzTab file.  
...  
PRH accession unit\_id … ambiguity\_members …  
PRT P19012 EXP\_1 P13646, P08779, P02533, Q7Z3Z0, Q7Z3Y9, Q7Z3Y8 …  
...  
PEH sequence accession unit\_id …  
PEP ALEEANADLEVK P19012 EXP\_1 …

## Advanced topics

There are several other features in mzTab that could not be introduced here. Detailed information about these features can be found in the specification document such as:

* Reporting post-translational modifications including modification position ambiguity.
* Reporting results from multiple search engines.
* Reporting replicates in mzTab.
* Merging mzTab files.
* Referencing external spectra.
* Referencing external resources such as mzIdentML or mzQuantML files.
* Adding optional columns.

An up-to-date list of example files can be found at <http://code.google.com/p/mztab/wiki/ExampleFiles>. The specification document can be found at <http://code.google.com/p/mztab/>.