PSI Recommendation
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mzTab: exchange format for proteomics and metabolomics results

Status of This Document

This document presents a draft specification for the mzTab data format developed by members of the Human Proteome Organisation (HUPO) Proteomics Standards Initiative (PSI) Proteomics Informatics (PI) Working Group. Distribution is unlimited.

Version of This Document

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Abstract

The Human Proteome Organisation (HUPO) Proteomics Standards Initiative (PSI) defines community standards for data representation in proteomics to facilitate data comparison, exchange and verification. The Proteomics Informatics Working Group is developing standards for describing the results of identification and quantification processes for proteins, peptides, small molecules and protein modifications from mass spectrometry. This document defines a tab delimited text file format to report proteomics and metabolomics results.

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1. Introduction

1.1 Background

This document addresses the systematic description of peptide, protein, and small molecule identification and quantification data retrieved from mass spectrometry (MS)-based experiments. A large number of software tools are available that analyze MS data and produce a variety of different output data formats. The HUPO Proteomics Standards Initiative (PSI) has developed several vendor-neutral data formats to overcome this heterogeneity of data formats for MS data. Currently, the PSI promotes the usage of three file formats to report an experiment's data: mzML to store the pure MS data (i.e. the spectra and chromatograms), mzIdentML to store (poly)peptide identifications and potentially inferred protein identifications, and mzQuantML to store quantitative data associated with these results. All three of these formats are XML-based and require sophisticated software to access the stored data.

While full, detailed representation of MS data including provenance is essential for researchers in the field, many downstream analysis use cases are only concerned with the results of the experiment in an easily accessible format. In addition, there is a trend for performing more integrated experimental workflows involving both proteomics and metabolomics data. Thus, the current lack of standardization in the field of metabolomics was taken into account in the development of the format presented here, and structures were developed that can report protein, peptide, and small molecule MS based data.

mzTab is intended as a lightweight supplement to the already existing standard file formats, providing a summary, similar to the supplementary table of results of a scientific publication.

mzTab files can contain protein, peptide, and small molecule identifications together with basic quantitative information. mzTab is not intended to store an experiment's complete data / evidence but only its final reported results. This format is also intended to provide local LIMS systems as well as MS proteomics repositories a simple way to share and combine basic information.

mzTab has been developed with a view to support the following general tasks (more specific use cases are provided in Section 2):

- T1. Facilitate the sharing of final experimental results, especially with researchers outside the field of proteomics that i) lack specialized software to parse the existing PSI's XML-based standard file formats, and ii) are only interested in the final reported results and not in all the details related to the data processing due to the inherent complexity of MS proteomics data. Furthermore, this should encourage the development of small innovative tools without the requirement of parsing huge XML files, which might be outside the scope of many bioinformaticians.
- T2. Export of results to external software, that is not able to parse proteomics/metabolomics specific data formats but can handle simple tab-delimited file formats. As a guideline the file format is designed to be viewable by programs such as Microsoft Excel® and Open Office Spreadsheet.
- T3. Contain the results of an experiment in a single file, and thus not require linking two files to retrieve identification and quantification results to again simplify the processing of the data.
- T4. Act as an output format of (web-) services that report MS-based results and thus can produce standardized result pages.
- T5. Allow the combination of MS-based proteomics and metabolomics experimental results within a single file.
- T6. Be able to link to the external experimental evidence (i.e. the mass spectra in different formats), following the same approach used in mzldentML and mzQuantML.

This document presents a specification, not a tutorial. As such, the presentation of technical details is deliberately direct. The role of the text is to describe the model and justify design decisions made. The document does not discuss how the models should be used in practice, consider tool support for data capture or storage, or provide comprehensive examples of the models in use. It is anticipated that tutorial material will be developed independently of this specification.

1.2 Document Structure

The remainder of this document is structured as follows. Section 2 lists use cases mzTab is designed to support. Section 3 describes the terminology used. Section 4 describes how the specification presented in Section 6 relates to other specifications, both those that it extends and those that it is intended to complement. Section 5 discusses the reasoning behind several design decisions taken. Section 6 contains the documentation of the file. Section 7 lists use cases that are currently not supported. Conclusions are presented in Section 8.

2. Use Cases for mzTab

The following cases of usage have driven the development of the mzTab data model, and are used to define the scope of the format in version 1.0.

- 1. mzTab files should be simple enough to make proteomics/metabolomics results accessible to people outside the respective fields. This should facilitate the sharing of data beyond the borders of the fields and make it accessible to non-experts.
- 2. mzTab files should contain sufficient information to provide an electronic summary of all findings in a proteomics/metabolomics study to permit its use as a standard documentation format for 'supplementary material' sections of publications in proteomics and metabolomics. It should thus be able to replace PDF tables as a way of reporting peptides and proteins and make published identification and quantification information more accessible.
- 3. When different samples and assays (including replicates) are reported in a single mzTab file, this file can be generated in two ways: 'Summary' mode, and 'Complete' mode. In 'Summary' full results per assay/replicate need not be included, only the final data for the experimental conditions analysed must be present. In 'Complete' mode, all the results per assay/replicate need to be detailed.
- 4. It should be possible to open mzTab files with "standard" software such as Microsoft Excel® or Open Office Spreadsheet. This should furthermore improve the usability of the format to people outside the fields of proteomics/metabolomics.
- 5. It should be possible to export proteomics data from, for example, mzldentML/mzQuantML files into mzTab to then load this data into, for example, statistical tools such as those provided through the R programming language. With the current formats, complex conversion software would be needed to make proteomics results available to such environments.
- 6. mzTab files should make MS derived results easily accessible to scripting languages allowing bioinformaticians to develop software without the overhead of developing sophisticated parsing code. Since mzTab files will be comparatively small, the data from multiple experiments can be processed at once without requiring special resource management techniques.
- 7. It should be possible to contain the complete final results of an MS-based proteomics/metabolomics experiment in a single file. This should furthermore reduce the complexity of sharing and processing an experiment's final results. mzTab files should be able to store quantitative values for protein, peptide, and small molecule identifications. Furthermore, mzTab files should contain basic protein inference information and modification position ambiguity information. Additionally, mzTab files should be able to report merged results from multiple search engines.
- 8. It should be useful as an output format by web-services that can then be readily accessed by tools supporting mzTab.
- 9. As mzTab files only contain an experiment's core results, all entries should link back to their source. Furthermore, it should be possible to directly link a given peptide / small molecule identification to its source spectrum in an external MS data file. The same referencing system as in mzIdentML/mzQuantML should be used.

3. Notational Conventions

The key words "MUST," "MUST NOT," "REQUIRED," "SHALL," "SHALL NOT," "SHOULD," "SHOULD NOT," "RECOMMENDED," "MAY," and "OPTIONAL" are to be interpreted as described in RFC-2119 (Bradner 1997).

4. Relationship to Other Specifications

The specification described in this document has not been developed in isolation; indeed, it is designed to be complementary to, and thus used in conjunction with, several existing and emerging models. Related specifications include the following:

- mzML (http://www.psidev.info/mzml). mzML is the PSI standard for capturing mass spectra / peak lists resulting from mass spectrometry in proteomics (Martens, L., et al. 2011). mzTab files MAY be used in conjunction with mzML, although it will be possible to use mzTab with other formats of mass spectra. This document does not assume familiarity with mzML.
- mzldentML (http://www.psidev.info/mzidentml). mzldentML is the PSI standard for capturing of peptide and protein identification data (Jones, A. R., et al. 2012). mzTab files MAY reference mzldentML files that then contain the detailed evidence of the reported identifications.
- mzQuantML (http://www.psidev.info/mzquantml). mzQuantML is the PSI standard for capturing quantitative proteomics data from mass spectrometry (Walzer, M. et al. 2013). mzTab files that report quantitative data MAY reference mzQuantML files for detailed evidence of the reported values.

4.1 The PSI Mass Spectrometry Controlled Vocabulary (CV)

The PSI-MS controlled vocabulary is intended to provide terms for annotation of mzML, mzIdentML, and mzQuantML files. The CV has been generated with a collection of terms from software vendors and academic groups working in the area of mass spectrometry and proteome informatics. Some terms describe attributes that must be coupled with a numerical value attribute in the CvParam element (e.g. MS:1001191 "p-value") and optionally a unit for that value (e.g. MS:1001117, "theoretical mass", units = "dalton"). The terms that require a value are denoted by having a "datatype" key-value pair in the CV itself: MS:1001172 "mascot:expectation value" value-type:xsd:double. Terms that need to be qualified with units are denoted with a "has_units" key in the CV itself (relationship: has_units: UO:0000221! dalton).

As recommended by the PSI CV guidelines, psi-ms.obo should be dynamically maintained via the psidev-ms-vocab@lists.sourceforge.net mailing list that allows any user to request new terms in agreement with the community involved. Once a consensus is reached among the community the new terms are added within a few business days. If there is no obvious consensus, the CV coordinators committee should vote and make a decision. A new psi-ms.obo should then be released by updating the file on the CVS server without changing the name of the file (this would alter the propagation of the file to the OBO website and to other ontology services that rely on file stable URI). For this reason an internal version number with two decimals (x.y.z) should be increased:

- x should be increased when a first level term is renamed, added, deleted or rearranged in the structure. Such rearrangement will be rare and is very likely to have repercussion on the mapping.
- y should be increased when any other term except the first level one is altered.
- z should be increased when there is no term addition or deletion but just editing on the definitions or other minor changes.

The following ontologies or controlled vocabularies specified below may also be suitable or required in certain instances:

- Unit Ontology (http://www.obofoundry.org/cgi-bin/detail.cgi?id=unit)
- ChEBI (http://www.ebi.ac.uk/chebi/)
- OBI (Ontology of Biological Investigations http://obi.sourceforge.net/)
- PSI Protein modifications workgroup http://psidev.cvs.sourceforge.net/psidev/psi/mod/data/PSI-MOD.obo
- Unimod modifications database http://www.unimod.org/obo/unimod.obo
- PRIDE Controlled Vocabulary (http://ebi-pride.googlecode.com/svn/trunk/pride-core/schema/pride_cv.obo)
- NEWT UniProt Taxonomy Database (http://www.ebi.ac.uk/ontology-lookup/browse.do?ontName=NEWT)
- BRENDA tissue/ enzyme source (http://www.brenda-enzymes.info/ontology/tissue/tree/update/update_files/BrendaTissueOBO).
- Cell Type ontology (http://obo.cvs.sourceforge.net/obo/obo/ontology/anatomy/cell_type/cell.obo).

5. Resolved Design and scope issues

There were several issues regarding the design of the format that were not clear cut, and a design choice was made that was not completely agreeable to everyone. So that these issues do not keep coming up, we document the issues here and why the decision that is implemented was made.

5.1 Handling updates to the controlled vocabulary

There is a difficult issue with respect to how software should encode CV terms, such that changes to the core can be accommodated. This issue is discussed at length in the mzML specification document (Martens, L *et al.* 2011), and mzTab follows the same convention. In brief, when a new term is required, the file producers must contact the CV working group (via the mailing list psidev-ms-vocab@lists.sourceforge.net) and request the new term. It is anticipated that problems may arise if a consumer of the file encounters a new CV term and they are not working from the latest version of the CV file. It has been decided that rather than aim for a workaround to this issue, it can be expected that data file consumers must ensure that the OBO file is up-to-date.

5.2 Use of identifiers for input spectra to a search

PSMs and small molecules MUST be linked to an identifier of the source spectrum (in an external file) from which the identifications are made by way of a reference in the spectra_ref attribute and via the ms_run element which stores the URL of the file in the location attribute.

It is advantageous if there is a consistent system for identifying spectra in different file formats. The following table is implemented in the PSI-MS CV for providing consistent identifiers for different spectrum file formats. This is the exact same approach followed in mzldentML and mzQuantML. Note, this table shows examples from the CV but will be extended. The CV holds the definite specification for legal encodings of spectrumID values.

ID Term Data type Co	omment
----------------------	--------

MS:1000768	Thermo	controllerType=xsd:nonNegativeInteger	controller=0 is usually the mass
	nativeID	controllerNumber=xsd:positiveInteger	spectrometer
ļ	format	scan=xsd:positiveInteger.	'
MS:1000769	Waters	function=xsd:positiveInteger	
	nativeID	process=xsd:nonNegativeInteger	
	format	scan=xsd:nonNegativeInteger	
MS:1000770	WIFF	sample=xsd:nonNegativeInteger	
	nativeID	period=xsd:nonNegativeInteger	
	format	cycle=xsd:nonNegativeInteger	
		experiment=xsd:nonNegativeInteger	
MS:1000771	Bruker/Agilent	scan=xsd:nonNegativeInteger	
	YEP nativeID		
	format		
MS:1000772	Bruker BAF	scan=xsd:nonNegativeInteger	
	nativeID		
	format		
MS:1000773	Bruker FID	file=xsd:IDREF	The nativeID must be the same as
	nativeID		the source file ID
140 4000774	format		
MS:1000774	multiple peak list nativeID	index=xsd:nonNegativeInteger	Used for referencing peak list files
	format		with multiple spectra, i.e. MGF,
	ioimat		PKL, merged DTA files. Index is the spectrum number in the file, starting
			from 0.
MS:1000775	single peak	file=xsd:IDREF	The nativeID must be the same as
1000773	list nativeID	IIIC-AGG.IDITEI	the source file ID. Used for
	format		referencing peak list files with one
	Tomat		spectrum per file, typically in a
			folder of PKL or DTAs, where each
			sourceFileRef is different
MS:1000776	scan number	scan=xsd:nonNegativeInteger	Used for conversion from mzXML,
	only nativeID		or a DTA folder where native scan
	format		numbers can be derived.
MS:1000777	spectrum	spectrum=xsd:nonNegativeInteger	Used for conversion from mzData.
	identifier		The spectrum id attribute is
	nativeID		referenced.
	format		
MS:1001530	mzML unique	xsd:string	Used for referencing mzML. The
	identifier	-	value of the spectrum id attribute is
			referenced directly.

Table 1 Controlled vocabulary terms and rules implemented in the PSI-MS CV for formulating the "nativeID" to identify spectra in different file formats.

In mzTab, the spectra_ref attribute should be constructed following the data type specification in Table 1. As an example, to reference the third spectrum (index = 2) in an MGF (Mascot Generic Format) file:

```
MTD ms_run[1]-format [MS, MS:1001062, Mascot MGF file, ]
MTD ms_run[1]-id_format [MS, MS:1000774, multiple peak list nativeID format, ]
...

PSH sequence ... spectra_ref ...
PSM NILNELFQR ... ms_run[1]:index=2 ...
```

Example: Reference the spectrum with identifier "scan=11665" in an mzML file.

```
MTD    ms_file[1]-format    [MS, MS:1000584, mzML file, ]
MTD    ms_run[1]-id_format    [MS, MS:1001530, mzML unique identifier, ]
...
PSH    sequence    ...    spectra_ref    ...
PSM    NILNELFQR    ...    ms_run[1]:scan=11665    ...
```

5.3 Recommendations for reporting replicates within experimental designs

Modeling the correct reporting of technical/biological replicates within experimental designs is supported in mzTab using an adaptation of the system originally developed for mzQuantML comprising four components described below (Figure 1). These components have various cross-references and MUST be used in different types of mzTab file, as described in Section 5.4:

- Study variable The variables about which the final results of a study are reported, which may have been derived following averaging across a group of replicate measurements (assays). In files where assays are reported, study variables have references to assays. The same concept has been defined by others as "experimental factor".
- MS run An MS run is effectively one run (or set of runs on pre-fractionated samples) on an MS instrument, and is referenced from assay in different contexts.
- Assay The application of a measurement about the sample (in this case through MS)

 producing values about small molecules, peptides or proteins. One assay is typically
 mapped to one MS run in the case of label-free MS analysis or multiple assays are
 mapped to one MS run for multiplexed techniques, along with a description of the label
 or tag applied.
- Sample a biological material that has been analysed, to which descriptors of species, cell/tissue type etc. can be attached. In all of types of mzTab file, these MAY be reported in the metadata section as sample[1-n]-description. Samples are NOT MANDATORY in mzTab, since many software packages cannot determine what type of sample was analysed (e.g. whether biological or technical replication was performed).

Clear definitions of biological and technical replicates are difficult to provide as these are somewhat dependent upon the biological domain. However, we use the following general definitions in mzTab.

- Biological replicates are where different samples have been analysed by MS.
- Technical replicates are where same samples are analysed multiple times by (LC)-MS.

Note: there is deliberately no attempt to define the boundary of the term "sample".

If sample level information is provided optimally, it is expected that *n* biological replicates can be mapped to sample[1-n]; *m* technical replicate measurements of sample 1 SHOULD be mapped to assay[1-m] referencing sample[1] (for example). However, an open challenge

remains since analysis software is often not aware of whether replicates (multiple MS runs) are originally biological or technical in nature. As such, the default behavior for mzTab exporters from quantitative software is to exclude sample level information and report quantitative data for assay[1-n] and/or study_variable[1-n] depending on whether it is a 'Complete' or 'Summary' file. Additional annotation software would typically be required to add the sample-level information, as provided (often manually) by the user.

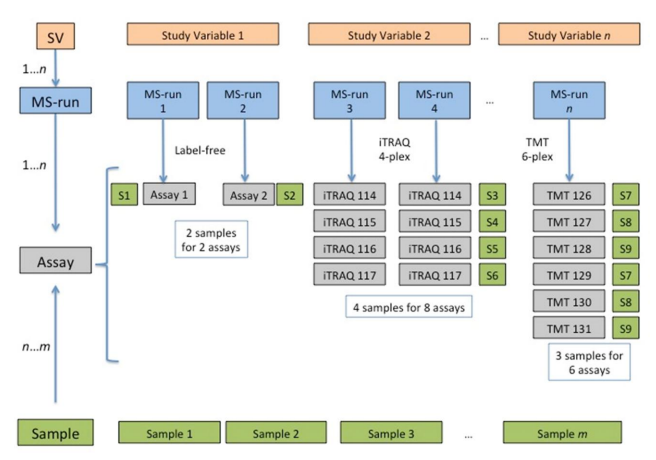


Figure 1. Diagram summarizing the relation between Study Variables (SVs), MS runs, assays and samples.

5.4 mzTab types 'Identification' and 'Quantification'

There are two types of mzTab files which MUST be specified by the mandatory meta-value 'mzTab-type' ('Identification' or 'Quantification'). 'Identification' MUST be used to report raw peptide, protein and small molecule identifications. The type 'Quantification' MUST be used for quantification results (which optionally might contain identification results about the quantified protein/peptide or small molecules). 'Quantification' files MUST always report quantification data on the level of study variables and MAY report quantification data on the level of assays. In contrast, 'Identification' files MAY contain neither study variables nor assays but only report identifications on the level of MS runs. Of course, 'Identification' files SHOULD include information about study variables and assays if this information is available.

Providing metadata on samples is not mandatory in both mzTab types as most software for quantification and identification can't readily export this information.

5.5 mzTab modes 'Summary' and 'Complete'

There are two modes of reporting of 'Identification' and 'Quantification' type results in mzTab files which MUST be specified by the mandatory meta value 'mzTab-mode' ('Summary' and 'Complete'). The 'Summary' mode is used if final results are provided (e.g. quantification data at the level of study variables). The 'Complete' mode is used if all quantification data is provided (e.g. quantification on the assay level and on the study variable level).

The MANDATORY fields in the Metadata Section 'mzTab-mode' and 'mzTab-type' MUST therefore be present to indicate which type of file it is. In Table 2, the columns that MUST be included in a 'Summary' file are indicated and in Table 3 the columns that MUST be included in addition in a 'Complete' file. It must be highlighted that all the columns in Tables 2 and 3 MUST be present in a "Complete" file.

In general, "null" values SHOULD not be given within any column of a "Complete" file if the information is available.

Section type	Identification	Quantification
(Cardinality) Metadata Section (1)	mzTab-version mzTab-mode mzTab-type description ms_run[1-n]-location fixed_mod[1-n] (if PSM section is present) variable_mod[1-n] (if PSM section is present)	mzTab-version mzTab-mode mzTab-type description ms_run[1-n]-location fixed_mod[1-n] (if PSM section is present) variable_mod [1-n] (if PSM section is present) protein-quantification_unit (if protein section is present) peptide-quantification_unit (if peptide section is present) smallmolecule-quantification_unit (if small molecule section is present) study_variable[1-n]-description
Protein Section (01)	accession description taxid species, database database_version search_engine best_search_engine_score ambiguity_members modifications protein_coverage	accession description taxid species database database_version search_engine best_search_engine_score ambiguity_members modifications protein_coverage protein_abundance_study_variable[1-n] protein_abundance_std_error_study_variable[1-n]
Peptide Section (01)	Peptide section is NOT RECOMMENDED to be used in identification only files.	sequence accession unique database database_version search_engine best_search_engine_score modifications retention_time retention_time_window charge

Г		
		mass_to_charge
		peptide_abundance_study_variable[1-n]
		peptide_abundance_stdev_study_variable[1-n]
		peptide_abundance_std_error_study_variable[1-n]
PSM Section (01)	sequence	sequence
· • · · · · · · · · · · · · · · · · · ·	PSM ID	PSM ID
	accession	accession
	unique	unique
	database	database
	database_version	database_version
	search engine	search engine
	search_engine_score	search_engine_score
		modifications
	modifications	
	spectra_ref	spectra_ref
	retention_time	retention_time
	charge	charge
	exp_mass_to_charge	exp_mass_to_charge
	calc_mass_to_chargepre	calc_mass_to_charge
	post	pre
	start	post
	end	start
		end
SmallMolecule	identifier	identifier
	chemical_formula	chemical_formula
Section (01)	smiles	smiles
	inchi_key	inchi_key
	description	description
	exp_mass_to_charge	exp_mass_to_charge
	_ ·	•
	calc_mass_to_charge	calc_mass_to_charge
	charge	charge
	retention time	retention time
	taxid	taxid
	species	species
	database	database
	database_version	database_version
	spectra_ref	spectra_ref
	search_engine	search_engine
	best_search_engine_score	best_search_engine_score
	modifications	modifications
		smallmolecule_abundance_assay[1-n] (if assays
		are reported)
		smallmolecule_abundance_study_variable[1-n] (if
		study variables are reported)
		smallmolecule_stdev_study_variable[1-n] (if study
		variables are reported)
		smallmolecule_std_error_study_variable[1-n] (if
		study variables are reported)
		study variables are reported)

Table 2. Mandatory columns in mzTab 'Summary' files. Where noted, these columns are mandatory for every study_variable[1-n] or every ms_run[1-n] reported in the file. Note — any Quantification file type MAY include any of the Columns or Sections required for an Identification file type.

Section type (Cardinality)	Identification	Quantification
Metadata Section (1)	software[1-n] fixed_mod[1-n] variable_mod[1-n]	software[1-n] fixed_mod[1-n] variable_mod [1-n] quantification_method assay[1-n]-ms_run_ref assay[1-n]-quantification_reagent study_variable[1-n]-assay_refs
Protein Section (01)	search_engine_score_ms_run[1-n] num_psms_ms_run[1-n] num_peptides_distinct_ms_run[1-n] num_peptide_unique_ms_run[1-n]	search_engine_score_ms_run[1-n] protein_abundance_assay[1-n]
Peptide Section (01)	Peptide section is NOT RECOMMENDED to be used in identification only files.	search_engine_score_ms_run[1-n] peptide_abundance_assay[1-n]

spectra_ref (if MS ² based quantification employed)
search_engine_score_ms_run[1-n]

Table 3. Mandatory columns in mzTab 'Complete' files. In addition, 'Complete' files MUST also have all the items that are MANDATORY in a 'Summary' file (Table 2 above). Where noted, these columns are mandatory for every assay[1-n], ms_run[1-n] or study_variable[1-n] reported in the file.

The rest of the fields are optional for both 'Complete' and 'Summary' files (Table 4).

Section type	Identification	Quantification
		- Cadiminoution
(Cardinality)	mzTab-ID	maTob ID
Metadata Section (1)	title	mzTab-ID title
	sample_processing[1-n]	sample_processing[1-n]
	instrument[1-n]-name	instrument[1-n]-name
	instrument[1-n]-source	instrument[1-n]-name
	instrument[1-n]-analyzer	instrument[1-n]-analyzer
	instrument[1-n]-detector	instrument[1-n]-detector
	software[1-n]-setting	software[1-n]-setting
	false discovery rate	false_discovery_rate
	publication[1-n]	publication[1-n]
	contact-name[1-n]	contact-name[1-n]
	contact-affiliation[1-n]	contact-affiliation[1-n]
	contact-email[1-n]	contact-email[1-n]
	uri	uri
	fixed_mod[1-n]-site	fixed_mod[1-n]-site
	fixed_mod[1-n]-position	fixed_mod[1-n]-position
	variable_mod[1-n]-site	variable_mod[1-n]-site
	variable_mod[1-n]-position	variable_mod[1-n]-position
	ms_run[1-n]-format	ms_run[1-n]-format
	ms_run[1-n]-id_format	ms_run[1-n]-id_format
	ms_run[1-n]-fragmentation_method	ms_run[1-n]-fragmentation_method
	custom	custom
	sample[1-n]-species	sample[1-n]-species
	sample[1-n]-tissue	sample[1-n]-tissue
	sample[1-n]-cell_type	sample[1-n]-cell_type
	sample[1-n]-disease	sample[1-n]-disease
	sample[1-n]-description	sample[1-n]-description
	sample[1-n]-custom	sample[1-n]-custom
	assay[1-n]-ms_run_ref	assay[1-n]-quantification_mod[1-n]
	assay[1-n]-sample_refs	assay[1-n]-quantification_mod[1-n]-position
	study_variable[1-n]-description	assay[1-n]-quantification_mod[1-n]-site
	study_variable[1-n]-sample_refs	assay[1-n]-sample_refs
	study_variable[1-n]-assay_refs	study_variable[1-n]-sample_refs
	cv[1-n]-label	cv[1-n]-label
	cv[1-n]-full_name	cv[1-n]-full_name
	cv[1-n]-version	cv[1-n]-version
	cv[1-n]-url	cv[1-n]-url
	colunit_protein	colunit_protein
	colunit_peptide	colunit_peptide
	colunit_psm	colunit_psm
	colunit_small_molecule	colunit_small_molecule
Protein Section (01)	opt_global_*	opt_global_*
i iotemi oection (o1)	go_terms	go_terms
	reliability	reliability
	uri	num_psms_ms_run[1-n]
		num peptides distinct ms run[1-n]
		num_peptide_unique_ms_run[1-n]
		uri
Peptide Section (01)		opt_global_*
- Spilac Coolion (U.1)	Peptide section is NOT RECOMMENDED to be	reliability
	used in identification only files.	uri
PSM Section (01)	opt_global_*	opt_global_*
1 3m 30000011 (01)	11 -0 12 -0	1 -0

	reliability	reliability
	uri	uri
SmallMolecule Section (01)	opt_global_* reliability uri	opt_global_* reliability uri

Table 4. Optional fields in mzTab 'Complete' and 'Summary' files.

5.6 Recommendations for reporting protein inference

There are multiple approaches to how protein inference can be reported. mzTab is designed to only hold experimental results, which in proteomics experiments can be very complex. At the same time, for downstream 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 have only limited support for protein inference/grouping reporting in mzTab files. Protein entries in mzTab files contain the field ambiguity_members. The protein accessions listed in this field should identify proteins that were also identified through the same set of peptides or spectra, or proteins supported by a largely overlapping set of evidence, and could also be a viable candidate for the "true" identification of the entity reported. It is RECOMMENDED that "subset proteins" that are unlikely to have been identified SHOULD NOT be reported here. The mapping of a single peptide-spectrum match (PSM) to multiple accessions is supported through the reporting of the same PSM on multiple rows of the PSM section, as exemplified below.

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

5.7 Recommendations for reporting quantification results

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 – for example in label-based techniques, such as SILAC/N¹⁵ where quantification occurs on MS¹ data or in tagbased techniques, such as iTRAQ/TMT where quantification occurs in MS² data.

One measurement of a small molecule, peptide or protein is mapped to the concept of assay for both multiplexed techniques and label-free techniques in Complete files. Each assay MUST have a reference to the quantification reagent/label used ("unlabelled" in the label-free case and the "light" channel in SILAC/N¹⁵) and each assay MUST have a reference to the ms_run[1_n] from which it originated. As such, in multiplexed techniques where *n* reagents are used within one analysis, assay[1-n] MUST reference the same ms_run.

If the data exporter wishes to report only final results for 'Summary' files (i.e. following averaging over replicates), then these MUST be reported as quantitative values in the

columns associated with the study_variable[1-n] (e.g. protein_abundance_study_variable[1]). mzTab allows the reporting of abundance, standard deviation, and standard error for any study_variable. The unit of values in the abundance column MUST be specified in the metadata section of the mzTab file. The reported values SHOULD represent the final result of the performed data analysis. The exact meaning of the values will thus depend on the used analysis pipeline and quantitation method and is not expected to be comparable across multiple mzTab files.

[Need to insert examples from label-free, MS1 label, MS2 tag, spectral counting and SRM here]

5.8 Reporting modifications and amino acid substitutions

Modifications are defined in the meta-data section and reported in the modification columns of the protein, peptide or PSM section.

Defining modifications in the meta-data section:

The meta values "fixed_modification[1-n]" and "variable_modification[1-n]" describe all search modifications used to identify peptides and proteins of the mzTab file (e.g. carbamidomethylation, oxidation, labels/tags). This is the minimal information that MUST be provided for Complete Identification or Quantification files.

In addition, for each assay the optional meta-data assay[1-n]-quantification_mod* MAY be specified that allows to define details of modifications associated with the quantification reagent (e.g. SILAC label).

Reporting of modifications in columns of the protein, peptide and PSM sections:

Fixed modifications or modifications specified as quantification_modification in the metadata Section SHOULD NOT be reported in protein (PRT) and peptide rows (PEP). In contrast, all variable modifications plus fixed modifications like those induced by the quantification reagents MUST be reported in peptide spectrum match rows (PSM).

Modifications or substitutions are modelled using a specific modification object with the following format:

{position}{Parameter}-{Modification or Substitution identifier}|{neutral loss}

The number of modification (or substitution) objects MUST correspond to the number of identified modifications (or substitutions) on a given peptide or PSM. It is also expected that modifications SHOULD be reported for proteins using the same format. However, it is recognised that some export software may not be able to do this. If software cannot determine protein-level modifications, "null" MUST be used. If the software has determined that there are no modifications to a given protein "0" MUST be used.

{position} is mandatory. However, if it is not known (e.g. MS1 Peptide Mass Fingerprinting), 'null' must be used Terminal modifications in proteins and peptides MUST be reported with the position set to 0 (N-terminal) or the amino acid length +1 (C-terminal) respectively. N-terminal modifications that are specifically on one amino acid MUST still be reported at the position 0. This object allows modifications to be assigned to ambiguous locations, but only at the PSM and Peptide level. Ambiguity of modification position MUST NOT be reported at the

Protein level. In that case, the modification element can be left empty. Ambiguous positions can be reported by separating the {position} and (optional) {cvParam} by an '|' from the next position. Thereby, it is possible to report reliabilities / scores / probabilities etc. for every potential location.

```
Here only the modification field is given:

3-MOD:00412, 8-MOD:00412

TESTPEPTIDES with two known phosphorylation sites
3|4-MOD:00412, 8-MOD:00412

First phosphorylation site can be either on S or T
3|4|8-MOD:00412, 3|4|8-MOD:00412

Three possible positions for two phosphorylation sites
```

{Parameter} is optional. It MAY be used to report a numerical value e.g. a probability score associated with the modification or location.

```
Reporting the first two possible sites for the phosphorylation with given probability score Here only the modification field is given:

3[MS,MS:1001876, modification probability, 0.8]|4[MS,MS:1001876, modification probability, 0.2]
MOD:00412, 8-MOD:00412
```

This option is not allowed though:

```
(3|4) [MS,MS:1001876, modification probability, 0.8] |7[MS,MS:1001876, modification probability, 0.2]-MOD:00412
```

(Modification or Substitution identifier) for proteins and peptides modifications SHOULD be reported using either UNIMOD or PSI-MOD accessions. As these two ontologies are not applicable to small molecules, so-called CHEMMODs can also be defined. Two types of CHEMMODs are allowed: specifying a chemical formula or specifying a given m/z delta. Additionally, it is possible to report substitutions of amino acids using SUBST:{amino acid}. In these cases, the "sequence" column MUST contain the original, unaltered sequence. The list of allowed {Modification or Substitution identifier}s therefore is:

```
CHEMMOD:+NH4
CHEMMOD:-18.0913
UNIMOD:18
MOD:00815
SUBST:{amino acid}
```

CHEMMODs SHOULD NOT be used for protein/peptide modifications if the respective entry is present in either the PSI-MOD or the UNIMOD ontology. Furthermore, mass deltas SHOULD NOT be reported if the given delta can be expressed through a known and unambiguous chemical formula.

All (identified) variable modifications as well as fixed modifications MUST be reported for every identification.

{neutral loss} is optional. Neutral losses are reported as cvParams. They are reported in the same way that modification objects are (as separate, comma-separated objects in the modification column). The position for a neutral loss MAY be reported.

```
PEH sequence ... modifications ...

COM Phosphorylation with a neutral loss:

PEP EISILACEIR ... 3-UNIMOD:21,3-[MS, MS:1001524, fragment neutral loss, 63.998285],7-UNIMOD:4

...

COM Neutral loss without an associated modification:

PEP EISILACEIR ... [MS, MS:1001524, fragment neutral loss, 63.998285],7-UNIMOD:4 ...
```

5.9 Encoding missing values, zeroes, nulls, infinity and calculation errors

In the table-based sections (protein, peptide, and small molecule) there MUST NOT be any empty cells. In case a given property is not available "null" MUST be used. This is, for example, the case when a URI is not available for a given protein (*i.e.* the table cell MUST NOT be empty but "null" has to be reported). If ratios are included and the denominator is zero, the "INF" value MUST be used. If the result leads to calculation errors (for example 0/0), this MUST be reported as "not a number" ("NaN"). 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 ("null").

5.10 Number of peptides reported

There are columns allowed in the protein section to report the number of peptides supporting a given protein identification, which are MANDATORY for Complete Identification files.

- num_psms_ms_run[1_n]
 - The count of the total significant PSMs that can be mapped to the reported protein
- num_peptides_distinct_ms_run[1_n]
 - The count of the number of different peptide sequences that have been identified above the significance threshold. Different modifications or charge states of the same peptide are not counted.
- num_peptides_unique_ms_run[1_n]
 - The number of peptides that can be mapped uniquely to the protein reported. If ambiguity members have been reported, the count MUST be derived from the number of peptides that can be uniquely mapped to the group of accessions, since the assumption is that these accessions are supported by the same evidence.

The idea of these three columns is to give the researcher a quick overview of how well a given protein identification is supported by peptide identifications for a given ms_run reported. The num_psms column also provides the opportunity for reporting pseudo-quantitative (label-free) values from approaches in which no explicit quantification has been performed.

5.11 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). This reliability only applies to the identification reliability but not to modification and or quantification reliabilities. The idea is to provide a way for researcher and/or MS proteomics or metabolomics repositories to score the reported identifications based on their own criteria. This score is completely resource-dependent and MUST NOT be interpreted 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, "null" MUST be provided as the value for each of the protein, peptide or small molecule identification.

The reliability value, if provided, 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 was 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 – depending on who is responsible for the given resource and how it is build. 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 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. Resources MUST explicitly specify how these reliability scores are calculated and what metric they represent.

5.12 Comments on Specific Use Cases

Many special use cases for mzTab were considered during its development. Each of these use cases has a corresponding example file that exercises the relevant part of the format and provides a reference implementation example (see supporting documentation). Authors of software that create mzTab are encouraged to examine the examples that accompany this format release before implementing the writer.

5.12.1 Multiple database search engines

Proteomics groups now commonly analyze MS data using multiple search engines and combine results to improve the number of peptide and protein identifications that can be made. The output of such approaches can be represented in mzTab as follows: mzTab files SHOULD only contain the "final" protein list generated by any such workflow. Any protein, peptide, and small molecule can be associated with any number of search engines as well as multiple search engine scores. Thus, it is possible to report which element was identified by which search engine together with the resulting scores.

5.12.2 Adding optional columns

Additional columns MAY be added to the end of rows in all the table-based sections (protein, peptide, PSM and small molecule). These columns represent information not included by default in the currently defined fields and differ from the specification of optionality with regards to columns that MUST be present in Summary or Complete files (Tables 2 and 3).

These column headers MUST start with the prefix "opt_" followed by the identifier of the object they reference: assay, study variable, MS run or "global" (if the value relates to all replicates). Column names MUST only contain the following characters: 'A'-'Z', 'a'-'z', '0'-'9', '_', '-', '[', ']', and ':'. CV parameter accessions MAY be used for optional columns following the format: opt_{OBJECT_ID}_cv_{accession}_{parameter name}. Spaces within the parameter's name MUST be replaced by '_'.

The information stored within an optional column is completely up to the resource that generates the file. It MUST not be assumed that optional columns having the same name in different mzTab files contain the same type of information. CV parameter accessions MAY be

used as optional column names according to the following convention: opt_{OBJECT_ID}_cv_{accession}_{parameter name}. Spaces within the parameter's name MUST be replaced by '_'.

```
COM Example showing how emPAI values are reported in an additional column from MS run 1 using COM MS CV parameter "emPAI value" (MS:1001905) ...

PRH accession ... opt_ms_run[1]_cv_MS:1001905_emPAI_value

PRT P12345 ... 0.658
```

5.12.3 Referencing external resources (i.e. mzldentML or mzQuantML files)

In mzTab all identifications MAY reference external resources that contain detailed evidence for the identification. This link is stored in the "uri" column of the respective table. This field MUST NOT be used to reference an external MS data file. MS data files should be referenced using the method described in Section 5.2.

Where these URIs point to depends on the resource that generated the mzTab file. If, for example, PeptideAtlas was exporting data in the mzTab format the URI would be expected to point to the identification's entry within the respective PeptideAtlas build. mzTab files originating from an mzIdentML file MAY reference the mzIdentML file using the URI column. In case quantitative values are reported coming from an mzQuantML file, the mzQuantML file SHOULD be referenced as it contains the reference to the underlying mzIdentML file.

5.12.4 Reporting sequence ambiguity

In MS based proteomics approaches, some amino acids cannot be unambiguously identified. To report such ambiguous amino acid identifications, the following symbols SHOULD be used:

```
Asparagine or aspartic acid B
Glutamine or glutamic acid Z
Leucine or Isoleucine J
Unspecified or unknown amino acid X
```

5.12.5 Reporting decoy peptide identifications

To report the results of a target-decoy search, decoy identifications MAY be labeled using the optional column "opt_global_cv_MS:1002217_decoy_peptide". The value of this column MUST be a Boolean (1/0).

5.13 Other supporting materials

The following example instance documents are available and between them cover all the use cases supported.

All example files can be downloaded from:

http://code.google.com/p/mztab/wiki/ExampleFiles

- a) mztab_SILAC_example.txt (hand crafted) mzTab file showing how SILAC data can be reported.
- b) mztab_itraq_example.txt (hand crafted) mzTab file showing how iTRAQ data can be reported.
- c) mztab merged example.txt merged version of the example file a and b.
- d) PRIDE_Exp_Complete_Ac_16649.xml-mztab.txt file generated using the mztab-exporter (converted PRIDE experiment accession 16649) containing iTRAQ data.
- e) mztab_lipidomics_example.txt Example containing MS lipidomics data produced by the Lipid Data Analyzer tool (http://genome.tugraz.at/lda/lda_download.shtml).

- f) PXD000002_mztab.txt.gz Summary file of ProteomeXchange submission PXD000002 (the complete submission can be found at ftp://ftp.pride.ebi.ac.uk/2012/03/PXD000002/).
- g) CPTAC_Progenesis_label_free_mzq.txt Label free example. Created by an exporter from an mzQuantML file.

6. Format specification

This section describes the structure of an mzTab file.

Field separator

The column delimiter is the Unicode Horizontal Tab character (Unicode codepoint 0009).

• File encoding

The UTF-8 encoding of the Unicode character set is the preferred encoding for mzTab files. However, parsers should be able to recognize commonly used encodings.

Case sensitivity

All column labels and field names are case-sensitive.

• Line prefix

Every line in an mzTab file MUST start with a three letter code identifying the type of line delimited by a Tab character. The three letter codes are as follows:

- 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
- PSH for the PSM table header (the column labels)
- PSM for rows of the PSM table
- SMH for small molecule table header line (the column labels)
- SML for rows of the small molecule table
- COM for comment lines

Header lines

Each table based section (protein, peptide, PSM and small molecule) MUST start with the corresponding header line. These header lines MUST only occur once in the document since each section also MUST only occur once.

Dates

Dates and times MUST be supplied in the ISO 8601 format ("YYYY-MM-DD", "YYYY-MM-DDTHH:MMZ" respectively).

• Decimal separator

In mzTab files the dot (".") MUST be used as decimal separator. Thousand separators MUST NOT be used in mzTab files.

Comment lines and empty lines

Comment lines can be placed anywhere in an mzTab file. These lines must start with the three-letter code COM and are ignored by most parsers. Empty lines can also occur anywhere in an mzTab file and are ignored.

Params

mzTab makes use of CV parameters. As mzTab is expected to be used in several experimental environments where parameters might not yet be available for the generated scores etc. all parameters can either report CV parameters or user parameters that only contain a name and a value.

Parameters are always reported as [CV label, accession, name, value]. Any field that is not available MUST be left empty.

```
[MS, MS:1001207, Mascot,]
[MS, MS:1001171, Mascot:score, 40.21]
[,,A user parameter, The value]
```

In case, the name of the param contains commas, quotes MUST be added to avoid problems with the parsing: [label, accession, "first part of the param name, second part of the name", value].

```
[MOD, MOD:00648, "N,O-diacetylated L-serine",]
```

Sample IDs

To be able to supply metadata specific to each sample, ids in the format sample[1-n] are used.

```
MTD sample[1]-species [NEWT, 9606, Homo sapiens (Human), ]
```

Assay IDs

To be able to supply metadata specific to each assay, ids in the format assay[1-n] are used.

```
MTD assay[1]-quantification_reagent [MS,MS:1002038,unlabeled sample,]
```

Study variable IDs

To be able to supply metadata specific to each study variable (grouping of assays), ids in the format study_variable[1-n] are used.

```
MTD study variable[1]-description Group B (spike-in 0.74 fmol/uL)
```

6.1 Sections

mzTab files can contain five different sections. The MANDATORY metadata section is made up of key-value pairs. The other four sections are OPTIONAL: protein, peptide, PSM and small molecule section are table-based.

Every section in an mzTab file MUST only occur once if present. If the PSM, Peptide and Protein Sections are present, the information MUST be consistent between these sections. Field names with indices in square brackets MUST be numbered sequentially and non-decreasing (starting at the first value indicated in the bracket; single integer steps).

6.2 Metadata Section

The metadata section can provide additional information about the dataset(s) reported in the mzTab file. All fields in the metadata section are optional apart from five exceptions:

- "mzTab-version" MUST always be reported.
- "mzTab-mode" MUST always be reported. Two modes are possible: 'Summary' and 'Complete'.
- "mzTab-type" MUST always be reported. Two types are possible: 'Quantification' or 'Identification'. Any analyses generating both quantification and identification results MUST be flagged as 'Quantification'.
- "description" MUST always be reported.
- "ms_run-location[1-n]" MUST always be reported.

In addition, various other metadata parameters are REQUIRED for different file types, as defined above and in Tables 2 and 3.

The fields in the metadata section should be reported in order of the various fields listed here. The field's name and value MUST be separated by a tab character:

```
MTD publication [PRIDE, PRIDE:00000029, PubMed, 12345]
```

In the following list of fields any term encapsulated by {} is meant as a variable which MUST be replaced accordingly.

6.2.1 mzTab-version

Description:	The version of the mzTab file.			
Туре:	String			
! !		Summary	Complete	
Mandatory	Quantification	√	√	
	Identification	√	√	
Example:	MTD mzTab-v	ersion	1.0	

6.2.2 mzTab-mode

Description:	The results included in an mzTab file can be reported in 2 ways: 'Complete' (when results for each assay/replicate are included) and 'Summary', when only the most representative results are reported.		
Type:	Enum		
Mandatory	Quantification Identification	ımmary Complete ✓ ✓ ✓	
Example:	MTD mzTab-mode Complete MTD mzTab-mode Summary		

6.2.3 mzTab-type

Description:	The results included in an mzTab file MUST be flagged as 'Identification' or 'Quantification' - the latter encompassing approaches that are quantification only or quantification and identification.				
Type:	Enum				
Mandatory	Summa Quantification ✓ Identification ✓	ry Complete ✓ ✓			
Example:	MTD mzTab-type Quantification MTD mzTab-type Identification				

6.2.4 mzTab-ID

Description:	The ID of the mzTab file.				
Type:	String				
	S	Summary	Complete		
Mandatory	Quantification				
	Identification				
Example:	MTD mzTab-ID	PRIDE_1	234		

6.2.5 title

Description:	The file's human readable title.
Туре:	String

		Summary	Complete	
Mandatory	Quantification			
	Identification			
Example:	MTD title	My first	-	eriment

6.2.6 description

Description:	The file's human readable description.			
Type:	String			
		Summary	Complete	
Mandatory Quantification Quantification				
	Identification	✓	✓	
Example:	MTD descri	otion An	experime	nt investigating the effects of Il-6.

6.2.7 sample_processing[1-n]

Description:	A list of parameters describing a sample processing step. The order of the data_processing items should reflect the order these processing steps were performed in. If multiple parameters are given for a step these MUST be separated by a " ".			
Type:	Parameter List			
Mandatory	Summary Complete Quantification Identification			
Example:	MTD sample_processing[1] [SEP, SEP:00173, SDS PAGE,] MTD sample_processing[2] [SEP, SEP:00142, enzyme digestion,] [MS, MS:1001251, Trypsin,]			

6.2.8 instrument[1-n]-name

Description:	The name of the instrument used in the experiment. Multiple instruments are numbered 1n.				
Туре:	Parameter				
Mandatani	Summary Complete				
Mandatory	Quantification				
	MTD instrument[1]-name [MS, MS:1000449, LTQ Orbitrap,]				
Example:	<pre> MTD instrument[2]-name [MS, MS:1000031, Instrument model, name of the instrument not included in the CV]</pre>				

6.2.9 instrument[1-n]-source

	The instrument's source used in the experiment. Multiple instruments are numbered 1n.					
}- <u></u>	Parameter					
Туре:	raiailletei					
	Summary Complete					
Mandatory	Quantification					
	Identification					
	MTD instrument[1]-source [MS, MS:1000073, ESI,]					
Example:						
•	MTD instrument[2]-source [MS, MS:1000598, ETD,]					

6.2.10 instrument[1-n]-analyzer

Description:	The instrument's analyzer type used in the experiment. Multiple instruments					
Description.	are enumerated 1n.					
Type:	Parameter					
	Summary Complete					
Mandatory	Quantification					
	Identification					
_	MTD instrument[1]-analyzer [MS, MS:1000291, linear ion trap,]					
Example:	MTD instrument[2]-analyzer [MS, MS:1000484, orbitrap,]					

6.2.11 instrument[1-n]-detector

Description:	The instrument's detector type used in the experiment. Multiple instruments					
Description.	are numbered 1n.					
Type:	Parameter					
	Summary Complete					
Mandatory	Quantification					
	Identification					
_	MTD instrument[1]-detector [MS, MS:1000253, electron multiplier,]					
Example:						
	MTD instrument[2]-detector [MS, MS:1000348, focal plane collector,]					

6.2.12 software[1-n]

Description:	Software used to analyze the data and obtain the results reported. The parameter's value SHOULD contain the software's version. The order (numbering) should reflect the order in which the tools were used.			
Type:	Parameter			
Summary Complete				
Mandatory Quantification ✓				
	Identification	,	/	
Example:		. ,		, Mascot, 2.3]
LXample.	MTD software[2]	[MS, MS:	L001561,	, Scaffold, 1.0]

6.2.13 software[1-n]-setting

Description:	A software setting used. This field MAY occur multiple times for a single software. The value of this field is deliberately set as a String, since there currently do not exist cvParams for every possible setting.			
Type:	String			
	Summa	y Complete		
Mandatory	Quantification			
	Identification			
Example:			ent tolerance = 0.1 Da	
Lvailibie.	MTD software[2]-setting Parent tolerance = 0.5 Da			

6.2.14 false_discovery_rate

Description:	The file's false discovery rate(s) reported at the PSM, peptide, and/or protein level. False Localization Rate (FLD) for the reporting of modifications can also be reported here. Multiple parameters MUST be separated by " ".		
Type:	Parameter List		
	Summary Complete		
Mandatory Quantification Quantification			
	Identification		
Example:	MTD false_discovery_rate [MS, MS:1001364, pep:global FDR, 0.01]		
Lampie.	[MS, MS:1001214, prot:global FDR, 0.08]		

6.2.15 publication[1-n]

Description:	A publication associated with this file. Several publications can be given by indicating the number in the square brackets after "publication". PubMed ids must be prefixed by "pubmed:", DOIs by "doi:". Multiple identifiers MUST be separated by " ".			
Type:	String			
Mandatory	Quantification Summary Identification	Complete		
Example:		ubmed:21063943 doi:10.1007/978-1-60761-987-1_6 ubmed:20615486 doi:10.1016/j.jprot.2010.06.008		

6.2.16 contact[1-n]-name

Description:	The contact's name. Several contacts can be given by indicating the number in the square brackets after "contact". A contact has to be supplied in the format [first name] [initials] [last name] (see example).		
Туре:	String		
	Summary	Complete	
Mandatory	Quantification Identification		
}	MTD contact[1]-name	James D. Watson	
Example:	MTD contact[2]-name	Francis Crick	

6.2.17 contact[1-n]-affiliation

Description:	The contact's affiliation.		
Type:	String		
	Sum	mary Complete	
Mandatory	Quantification		
,	Identification		
Evample	MTD contact[1]-a	ffiliation Car	mbridge University, UK
Example:	MTD contact[2]-a	ffiliation Car	mbridge University, UK

6.2.18 contact[1-n]-email

Description:	The contact's e-mail address.
Type:	String
	Summary Complete
Mandatory	Quantification
	Identification
	MTD contact[1]-email watson@cam.ac.uk
Example:	
	MTD contact[2]-email crick@cam.ac.uk

6.2.19 uri

Description:	A URI pointing to the file's source data (e.g., a PRIDE experiment or a PeptideAtlas build).		
Type:	URI		
	Summary Complete		
Mandatory	Quantification		
	Identification		
Example: MTD uri http://www.ebi.ac.uk/pride/url/to/experiment			
Example:	MTD uri http://proteomecentral.proteomexchange.org/cgi/GetDataset		

6.2.20 fixed_mod[1-n]

Description:	A parameter describing a fixed modifications searched for. Multiple fixed modifications are numbered 1n.		
Type:	Parameter		
	Summary	ry Complete	
Quantification (\checkmark) ¹		✓	
Mandatory	Identification (✓) ¹	✓	
	mandatory if PSM section is present		
Example:		[UNIMOD, UNIMOD:4, Carbamidomethyl,]	
Lample.	MTD fixed_mod[2] [U	[UNIMOD, UNIMOD:35, Oxidation,]	

6.2.21 fixed_mod[1-n]-site

Description:	A string describing a fixed modifications site. Following the unimod convention, modification site is a residue (e.g. "M"), terminus ("N-term" or "C-term") or both (e.g. "N-term Q" or "C-term K").
Type:	String

	Summary Complete
Mandatory	Quantification
	Identification
Example:	MTD fixed_mod[1] [UNIMOD, UNIMOD:35, Oxidation,] MTD fixed_mod[1]-site M MTD fixed_mod[2] [UNIMOD, UNIMOD:1, Acetyl,] MTD fixed_mod[2]-site N-term MTD fixed_mod[3] [UNIMOD, UNIMOD:2, Amidated,] MTD fixed_mod[3]-site C-term

6.2.22 fixed_mod[1-n]-position

Description:	A string describing the term specifity of a fixed modification. Following the unimod convention, term specifity is denoted by the strings "Anywhere", "Any N-term", "Any C-term", "Protein N-term", "Protein C-term".		
Type:	String		
Mandatory	Summary Complete Quantification Identification		
Example:	MTD fixed_mod[1] [UNIMOD, UNIMOD:35, Oxidation,] MTD fixed_mod[1]-site M MTD fixed_mod[2] [UNIMOD, UNIMOD:1, Acetyl,] MTD fixed_mod[2]-site N-term MTD fixed_mod[2]-position Protein N-term MTD fixed_mod[3] [UNIMOD, UNIMOD:2, Amidated,] MTD fixed_mod[3]-site C-term MTD fixed_mod[3]-position Protein C-term		

6.2.23 variable_mod[1-n]

Description:	A parameter modifications	A parameter describing a variable modifications searched for. Multiple variable modifications are numbered 1 n.				
Type:	Parameter					
	S	Summary	Complete			
Mondotory	Quantification	(✓) ¹	✓			
Mandatory	Identification	(√) ¹	✓			
	mandatory if PSM section is present					
Example:	MTD variable_r					
LAAIIIPIE.	MTD variable m	mod[1]	[UNIMOD,	UNIMOD:35,	Oxidation,]

6.2.24 variable_mod[1-n]-site

Description:	A string describing a variable modifications site. Following the unimod convention, modification site is a residue (e.g. "M"), terminus ("N-term" or "C-term") or both (e.g. "N-term Q" or "C-term K").		
Type:	String		
Mandatory	Summary Complete Quantification Identification		
Example:	MTD variable_mod[1] [UNIMOD, UNIMOD:35, Oxidation,] MTD variable_mod[1]-site M MTD variable_mod[2] [UNIMOD, UNIMOD:1, Acetyl,] MTD variable_mod[2]-site N-term MTD variable_mod[3] [UNIMOD, UNIMOD:2, Amidated,] MTD variable_mod[3]-site C-term		

6.2.25 variable_mod[1-n]-position

Description: A string describing the term specifity of a variable modification. Following the

	unimod convention, term specifity is denoted by the strings "Anywhere", "Any N-term", "Any C-term", "Protein N-term", "Protein C-term".		
Type:	String		
	Summary Complete		
Mandatory	Quantification		
	Identification		
Example:			

6.2.26 quantification_method

Description:	The quantification method used in the experiment reported in the file.			
Type:	Parameter			
		Summary	Complete	
Mandatory	Quantification		✓	
	Identification			
Example:	MTD quantifi	cation_me	thod [MS	, MS:1001837, iTRAQ quantitation analysis,]

6.2.27 protein-quantification_unit

Description:	Defines what type of units is reported in the protein quantification fields.			
Type:	Parameter			
	Summary Complete			
Mandatani	Quantification $(\checkmark)^1$ $(\checkmark)^1$			
Mandatory	Identification			
	¹ mandatory if protein section is present			
Example:	MTD protein-quantification_unit [PRIDE, PRIDE:0000395, Ratio,]			

6.2.28 peptide-quantification_unit

Description:	Defines what type of units is reported in the peptide quantification fields.			
Type:	Parameter			
	Summary Complete			
Mondotory	Quantification $(\checkmark)^1$ $(\checkmark)^1$			
Mandatory	Identification			
	mandatory if peptide section is present			
Example:	MTD peptide-quantification_unit [PRIDE, PRIDE:0000395, Ratio,]			

6.2.29 small_molecule-quantification_unit

Description:	Defines what type of units is reported in the small molecule quantification fields.				
Type:	Parameter				
	Summary Complete				
Mandatory	Quantification $(\checkmark)^1$ $(\checkmark)^1$				
Manuator y	Identification				
	¹ mandatory if small molecule section is present				
Example:	MTD small_molecule-quantification_unit [PRIDE, PRIDE:0000395, Ratio,]				

6.2.30 ms_run[1-n]-format

Description:	A parameter specifying the data format of the external MS data file.		
Type:	Parameter		
Mandatory	Summary Complete		
Ivialidatol y	Quantification		

	Identification	
Francis	MTD ms_run[1]-format [MS, MS:1000584, mzML fi	le,]
Example:	 MTD ms run[2]-format [MS, MS:1001062, Mascot N	MGF file,]

6.2.31 ms_run[1-n]-location

Description:	Location of the external data file. If the actual location of the MS run is unknown, a "null" MUST be used as a place holder value.		
Type:	URL		
Mandatory	Summary Complete Quantification		
Example:	MTD ms_run_location[1] file://C:\path\to\my\file MTD ms run location[2] ftp://ftp.ebi.ac.uk/path/to/file		

6.2.32 ms_run[1-n]-id_format

Description:	Parameter specifying the id format used in the external data file.		
Type:	Parameter		
	Sum	mary Complete	
Mandatory	Quantification		
,	Identification		
	MTD ms_run[1]-id	l_format [MS, N	MS:1000530, mzML unique identifier,]
Example:	 MTD ms_run[2]-id	l_format [MS, N	MS:1000774, multiple peak list
	_	_	<pre>nativeID format,]</pre>

6.2.33 ms_run[1-n]-fragmentation_method

Description:	A list of " " separated parameters describing all the types of fragmentation used in a given ms run.		
Type:	Parameter List		
	Summary Complete		
Mandatory	Quantification Quanti		
	Identification		
MTD ms_run[1]-fragmentation_method [MS, MS:1000133, CID,]			
Example:			
	MTD ms run[2]-fragmentation method [MS, MS:1000422, HCD,]		

6.2.34 custom

Description:	Any additional parameters describing the analysis reported.			
Type:	Parameter			
		Summary	Complete	
Mandatory	Quantification			
	Identification			
Example:	MTD custom	[,,MS ope	rator, Fl	orian]

6.2.35 sample[1-n]-species

Description:	The respective species of the samples analysed.		
Type:	Parameter		
	Summary Complete		
Mandatory	Quantification		
	Identification		
	COM Experiment where all samples consisted of the same two species		
	MTD sample[1]-species [NEWT, 9606, Homo sapiens (Human),]		
Example:	MTD sample[2]-species [NEWT, 12059, Rhinovirus,]		
Lxailipie.			
	COM Experiment where different two samples from different species (combinations)		

COM	were analysed as biological replicates.
MTI	sample[1]-species [NEWT, 9606, Homo sapiens (Human),]
MTI	sample[1]-species [NEWT, 573824, Human rhinovirus 1,]
MTI	sample[2]-species [NEWT, 9606, Homo sapiens (Human),]
MTI	sample[2]-species [NEWT, 12130, Human rhinovirus 2,]

6.2.36 sample[1-n]-tissue

Description:	The respective tissue(s) of the sample.		
Type:	Parameter		
	Summary	Complete	
Mandatory	Quantification		
	Identification		
Example:	MTD sample[1]-tissue	[BTO, BTO	0:0000759, liver,]

6.2.37 sample[1-n]-cell_type

Description:	The respective cell type(s) of the sample.		
Type:	Parameter		
	Summary Complete		
Mandatory	Quantification		
	Identification		
Example:	MTD sample[1]-cell_type [CL, CL:0000182, hepatocyte,]		

6.2.38 sample[1-n]-disease

Description:	The respective di	ease(s) of	the sample.
Type:	Parameter		
	Summa	y Complete	
Mandatory	Quantification		
	Identification		
Evample	MTD sample[1]-dise	se [DOID, D	OID:684, hepatocellular carcinoma,]
Example:	MTD sample[1]-disea	se [DOID, D	OID:9451, alcoholic fatty liver,]

6.2.39 sample[1-n]-description

Desc	cription:	A human readable description of the sample.
Туре	e:	String
		Summary Complete
Man	datory	Quantification
	,	Identification
Evar	mple:	MTD sample[1]-description Hepatocellular carcinoma samples.
Exai	ilibie.	MTD sample[2]-description Healthy control samples.

6.2.40 sample[1-n]-custom

	Description:	Parameters describing the sample's additional properties.		
	Туре:	Parameter		
[Summary Complete		
Ì	Mandatory	Quantification		
		Identification		
[Example:	MTD sample[1]-custom [,,Extraction date, 2011-12-21]		
l.,	Lxampie.	MTD sample[1]-custom [,,Extraction reason, liver biopsy]		

6.2.41 assay[1-n]-quantification_reagent

Description:	The reagent used to label the sample in the assay. For label-free analyses the "unlabeled sample" CV term SHOULD be used. For the "light" channel in label-based experiments the appropriate CV term specifying the labelling channel should be used.
Type:	Parameter
Mandatory	Summary Complete Quantification (✓) ¹ ✓

	Identification 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2
Example:	MTD assay[1]-quantification_reagent [PRIDE,PRIDE:0000114,iTRAQ reagent,114] MTD assay[2]-quantification_reagent [PRIDE,PRIDE:0000115,iTRAQ reagent,115] OR MTD assay[1]-quantification_reagent [MS,MS:1002038,unlabeled sample,] OR
	MTD assay[1]-quantification_reagent [PRIDE, PRIDE:0000326, SILAC light] MTD assay[2]-quantification_reagent [PRIDE, PRIDE:0000325, SILAC heavy]

6.2.42 assay[1-n]-quantification_mod[1-n]

Description:	A parameter describing a modification associated with a quantification_reagent. Multiple modifications are numbered 1n.		
Type:	Parameter		
	Summary Complete		
Mandatani	Quantification		
Mandatory	Identification 1 1		
	¹ not recommended for identification only files		
Example:	MTD assay[2]-quantification_mod[1] [UNIMOD, UNIMOD:188, Label:13C(6),]		

6.2.43 assay[1-n]-quantification_mod[1-n]-site

A string describing the modifications site. Following the unimod convention, modification site is a residue (e.g. "M"), terminus ("N-term" or "C-term") or both (e.g. "N-term Q" or "C-term K").		
String		
Summary Complete		
Quantification		
Identification 1 1 1 1 1 1 1 1 1		
MTD assay[2]-quantification_mod[1] [UNIMOD, UNIMOD:188, Label:13C(6),]		
MTD assay[2]-quantification_mod[2] [UNIMOD, UNIMOD:188, Label:13C(6),]		
<pre>MTD assay[2]-quantification_mod[1]-site R MTD assay[2]-quantification_mod[2]-site K</pre>		

6.2.44 assay[1-n]-quantification_mod[1-n]-position

Description:	A string describing the term specifity of the modification. Following the unimod convention, term specifity is denoted by the strings "Anywhere", "Any N-term", "Any C-term", "Protein N-term", "Protein C-term".		
Type:	String		
	Summary Complete		
Mandatani	Quantification		
Mandatory	Identification 1 1		
	¹ not recommended for identification only files		
	MTD assay[2]-quantification_mod[1] [UNIMOD, UNIMOD:188, Label:13C(6),]		
Example:	MTD assay[2]-quantification_mod[2] [UNIMOD, UNIMOD:188, Label:13C(6),]		
	MTD assay[2]-quantification_mod[1]-site R		
	MTD assay[2]-quantification_mod[2]-site K		
	MTD assay[2]-quantification_mod[1]-position Anywhere		
	MTD assay[2]-quantification mod[2]-position Anywhere		

6.2.45 assay[1-n]-sample_ref

Description:	An association from a given assay to the sample analysed.
Type:	{SAMPLE_ID}
Mandatory	Summary Complete Quantification

	Ide	ntification				
i – .	MTD	assay[1]	-sample_r	ef samp		٦
Example:	MTD	assay[2]	-	ef samp	ple[2]	١

6.2.46 assay[1-n]-ms_run_ref

Description:	An association from a given assay to the source MS run.					
Type:	{MS_RUN_ID}					
	Summary Complete					
Mondotory	Quantification $(\checkmark)^1$ \checkmark					
Mandatory	Identification $(\checkmark)^1$ $(\checkmark)^1$					
	mandatory if assays are reported					
Example:	MTD assay[1]-ms_run_ref ms_run[1]					

6.2.47 study_variable[1-n]-assay_refs

Description:	Comma-separated references to the IDs of assays grouped in the study variable.					
Type:	{ASSAY_ID},					
	Summary Complete					
Mandatory	Quantification (\checkmark) ¹ \checkmark					
Ivialidatol y	Identification					
	mandatory if both assays and study variables are reported					
Example:	MTD study_variable[1]-assay_refs assay[1], assay[2], assay[3]					

6.2.48 study_variable[1-n]-sample_refs

Description:	Comma-separated references to the samples that were analysed in the study variable.				
Type:	{SAMPLE_ID}, {SAMPLE_ID}				
	Summary Complete				
Mandatory	Quantification				
Example:	MTD study_variable[1]-sample_refs sample[1]				

6.2.49 study_variable[1-n]-description

Description:	A textual description of the study variable.					
Type:	String					
	Summary Complete					
	Quantification (\checkmark) ¹ \checkmark					
Mandatory	Identification $(\checkmark)^1$ $(\checkmark)^1$					
	¹ mandatory of study variables reported					
Example:	MTD study_variable[1]-description Group B (spike-in 0.74 fmol/uL)					

6.2.50 cv[1-n]-label

Description:	A string describing the labels of the controlled vocabularies/ontologies used in						
Description.	the mzTab file						
Type:	String						
		Summary	Complete				
Mandatory	Quantification						
	Identification						
_	MTD cv[1]-1a	abel MS					
Example:							
1							

6.2.51 cv[1-n]-full_name

Description:	A string describing the full names of the controlled vocabularies/ontologies	
Description.	used in the mzTab file	

Туре:	String		
		Summary	Complete
Mandatory	Quantification		
,	Identification		
_	MTD cv[1]-f	ull_name	MS
Example:			

6.2.52 cv[1-n]-version

Description:	A string describing the version of the controlled vocabularies/ontologies used in the mzTab file						
Type:	String						
Mandatory	Summary Complete Quantification Identification						
Example:	MTD cv[1]-version 3.54.0						

6.2.53 cv[1-n]-url

Description:	A string containing the URLs of the controlled vocabularies/ontologies used in the mzTab file				
Type:	String				
Mandatory	Quantification Identification	Summary	Complete		
Example:	MTD cv[1]-url http://psidev.cvs.sourceforge.net/viewvc/psidev/psi/psi-ms/mzML/controlledVocabulary/psi-ms.obo				

6.2.54 colunit-protein

Description:	Defines the unit for the data reported in a column of the protein section. The format of the value has to be {column name}={Parameter defining the unit} This field MUST NOT be used to define a unit for quantification columns. The unit used for protein quantification values MUST be set in protein-quantification_unit.
Туре:	String
Multiplicity:	0*
Example:	MTD

6.2.55 colunit-peptide

Description:	Defines the used unit for a column in the peptide section. The format of the value has to be {column name}={Parameter defining the unit} This field MUST NOT be used to define a unit for quantification columns. The unit used for peptide quantification values MUST be set in peptide-quantification_unit.					
Туре:	String					
Mandatory	Quantification Identification	Summary	Complete			

Example: MTD	colunit-peptide retention_time=[UO,UO:0000031, minute,]
--------------	---

6.2.56 colunit-psm

Description:	Defines the used unit for a column in the PSM section. The format of the value has to be {column name}={Parameter defining the unit} This field MUST NOT be used to define a unit for quantification columns. The unit used for peptide quantification values MUST be set in peptide-quantification_unit.								
Type:	String								
Mandatory	Summary Complete Quantification Identification								
Example:	MTD colunit-psm retention_time=[UO,UO:0000031, minute,]								

6.2.57 colunit-small_molecule

Description:	Defines the used unit for a column in the small molecule section. The format of the value has to be {column name}={Parameter defining the unit} This field MUST NOT be used to define a unit for quantification columns. The unit used for small molecule quantification values MUST be set in small_molecule-quantification_unit.								
Type:	String								
Mandatory	Summary Complete Quantification Identification								
Example:	MTD columit-small_molecule retention_time=[U0,U0:0000031, minute,]								

6.3 Protein Section

The protein section is table-based. The protein section MUST always come after the metadata section. All table columns MUST be tab-separated. There MUST NOT be any empty cells. Missing values MUST be reported using "null". Most columns are mandatory. The order of columns is not specified although for ease of human interpretation, it is RECOMMENDED to follow the order specified below.

6.3.1 accession

Description:	The accession of the protein in the source database. A protein accession MUST be unique within one mzTab file. If different quantification values are required for the same underlying accession, for example if differentially modified forms of a protein have been quantified, a the suffix [1-n] SHOULI be appended to the accession e.g. P12345[1], P12345[2].					
Type:	String					
		Summary	Complete			
Mandatory	Quantification	✓	✓			
	Identification	✓	✓			
	PRH accession	on				
Example:	PRT P12345					
l	PRT P12346					

6.3.2 description

Description:	The protein's name and or description line.
Type:	String
Mandatory	Summary Complete

ĺ		(Quantification	✓	✓		
1		I	dentification	✓	✓		
ſ		PR	H accession	n d	lescriptio	n	
1	Example:	PR	T P12345	Aspa	rtate ami	notransferase, mitochondrial	
1	• •	PR			transferr	in	

6.3.3 taxid

Description:	The NCBI/NEWT taxonomy id for the species the protein was identified in.						
Type:	Integer	Integer					
	Sum	mary	Complete				
Mandatory	Quantification	/	√				
	Identification	/	✓				
	PRH accession	tax	id				
Example:	PRT P12345	101	16				
	PRT P12346	101	16				

6.3.4 species

Description:	The human r the NCBI ent	The human readable species the protein was identified in - this SHOULD be the NCBI entry's name.						
Туре:	String							
Mandatory	Summary Complete Quantification Identification Summary Complete Quantification Identification Ident							
Example:	PRH accession PRT P12345 PRT P12346		16 Rattı	ies us norvegicus (Rat) us norvegicus (Rat)				

6.3.5 database

Description:	The protein database used for the search (could theoretically come from a different species). Wherever possible the Miriam (http://www.ebi.ac.uk/miriam) assigned name SHOULD be used.							
Type:	String							
		Summary	Complete					
Mandatory	Quantification	√	√					
	Identification	✓						
l	PRH accession		1	i i				
Example:	PRT P12345	101		us norvegicus (Rat) UniProtKB				
İ	PRT P12346	101	16 Rattı	us norvegicus (Rat) UniProtKB				

6.3.6 database_version

Description:	The protein database's version – in case there is no version available (custom build) the creation / download (e.g., for NCBI nr) date SHOULD be given. Additionally, the number of entries in the database MAY be reported in round brackets after the version in the format: {version} ({#entries} entries), for example "2011-11 (1234 entries)".							
Type:	String							
		Summary	Complete					
Mandatory	Quantification	√	√					
	Identification ✓ ✓							
	PRH accession				database	database_version		
Example:	PRT P12345	101		us norvegicus (Rat)				
l	PRT P12346	101	.16 Ratt	us norvegicus (Rat)	UniProtKB	2011 11		

6.3.7 search_engine

Description: A " " delimited list of search engine(s) used to identify this protein. Search engines MUST be supplied as parameters.						
Type:	Parameter List					
Mandatory	Summary Complete					
Mandatory	Quantification ✓ ✓ ✓					

I		Ider	ntification	✓	✓	
-		COM	In this	example	the first	protein was identified by Mascot and Sequest while
COM the second protein was only identified by Mascot.						
1	Example:	PRH	accessio	on s	earch engi	ne
İ	•	PRT	P12345	[MS,MS:1001	207, Mascot,] [MS, MS: 1001208, Sequest,]
ı		PRT	P12346	[MS,MS:1001	207, Mascot,]

6.3.8 best_search_engine_score

:	A " " delimited list of the best search engine score(s) for the given protein across all replicates reported. Scores SHOULD be reported using CV parameters whenever possible.			
Type:	Parameter	List		
Mondotory	Quantification	Summary	Complete	
Mandatory	Identification	√	✓	
	PRH accessi	on be	est_search	engine_score_ms_run[1]
Example:	 PRT P12345 PRT P12346			171, Mascot score, 50] [MS, MS:1001155, Sequest:xcorr, 2]

6.3.9 search_engine_score_ms_run[1-n]

Description:	A " " delimited list of search engine score(s) for the given protein. Scores SHOULD be reported using CV parameters whenever possible.			
Туре:	Parameter	List		
Mandatory	Quantification Identification	Summary	Complete ✓	
Example:	PRH accession PRT P12345 PRT P12346	[M	 MS:1001	ne_score_ms_run[1] 171,Mascot score,50] [MS,MS:1001155,Sequest:xcorr,2] 171,Mascot score,47.2]

6.3.10 reliability

Description:	The reliability of the given protein identification. This must be supplied by the resource and has to be one of the following values: 1: high reliability 2: medium reliability 3: poor reliability Important: An identification's reliability is resource-dependent.				
Type:	Integer			Tonasin, is researed aspendent.	
Mandatory	Quantification Identification	Summary	Complete		
Example:	PRH accession PRT P12345 PRT P12346	on re 3 1	eliability	 	

6.3.11 num_psms_ms_run[1-n]

Description:	The total number of PSMs that were used to identify this protein from a given						
Description.	ms_run.						
Type:	Integer						
		Summary	Complete				
Mandatory	Quantification						
	Identification		✓				
	COM P12345 i	s identif	ied throu	gh ABCM, ABCM+Oxidation, CDE, CDE			
Example:	 PRH accessic	on nu	m psms ms	run[1]			
	PRT P12345	4					

6.3.12 num_peptides_distinct_ms_run[1-n]

Description:	The number of distinct peptide sequences identifying this protein in a given ms_run. Distinct peptides are defined based on their sequence, ignoring different modifications or charge states.			
Type:	Integer			
		Summary	Complete	
Mandatory	Quantification			
Example:				gh ABCM, ABCM+Oxidation, CDE, CDE s distinct ms run[1]
	PRT P12345	3	m_bebride	 2_d12c1Hcc_H2_tan[1]

6.3.13 num_peptides_unique_ms_run[1-n]

Description:	The number of peptides that are mapped uniquely to this protein and the other ambiguity members in this ms_run.			
Type:	Integer			
Mandatory	Summary Complete Quantification			
Example:	COM P12345 is identified through ABCM, ABCM+Oxidation, CDE, CDE COM ABCM is only from P12345, CDE from P12345 and P12346 PRH accession num_peptides_unique_ms_run[1] PRT P12345 2			

6.3.14 ambiguity_members

Description:	A comma-delimited list of protein accessions. This field should be set in the representative protein of the ambiguity group (the protein identified through the accession in the first column). The accessions listed in this field should identify proteins that could also be identified through these peptides (e.g. "same-set proteins") but were not chosen by the researcher or resource, often for arbitrary reasons. It is NOT RECOMMENDED to report subset proteins as ambiguity_members, since the proteins reported here, together with the representative protein are taken to be a group that cannot be separated based on the peptide evidence.				
Type:	String List				
Mandatory	Summary Complete Quantification Identification Summary Complete Complet				
Example:	COM P12345, P12347, and P12348 can all be identified through the same peptides PRH accession ambiguity_members PRT P12345 P12347,P12348				

6.3.15 modifications

Modification location scores cannot be supplied at the Protein level. Furthermore, in case a position is unknown no position information MAY be supplied.	Description:	Furthermore, in case a position is unknown no position information MAY be
--	--------------	---

Terminal modifications MUST be reported at position 0 or protein size + 1 respectively.

Valid modification identifiers are either PSI-MOD or UNIMOD accession (including the "MOD:" / "UNIMOD:" prefix) or CHEMMODS. CHEMMODS have the format CHEMMOD:+/-{chemical formula or *m/z* delta}. Valid CHEMMODS are for example "CHEMMOD:+NH4" or "CHEMMOD:-10.1098". CHEMMODs MUST NOT be used if the modification can be reported using a PSI-MOD or UNIMOD accession. Mass deltas MUST NOT be used for CHEMMODs if the delta can be expressed through a known chemical formula.

Neutral losses MAY be reported as cvParams. If a neutral loss is not associated with an existing modification it is reported as separated commaseparated entry. Otherwise, the neutral loss MUST be reported after the modification it is associated with and separated by a '|' from the modification. Additionally, it is possible to report substitutions of amino acids using SUBST:{amino acid}.

If different modifications are identified from different ms_runs, a superset of the identified modifications SHOULD be reported here. Detailed modification mapping to individual ms_runs is provided through the PSM table.

If protein level modifications are not reported, a "null" MUST be used. If protein level modifications are reported but not present on a given protein, a "0" MUST be reported.

6.3.16 uri

Description:	A URI pointing to the protein's source entry in the unit it was identified in (e.g., the PRIDE database or a local database / file identifier).				
Type:	URI				
Mandatory	Summary Complete Quantification Identification				
Example:	PRT accession uri PRH P12345 http://www.ebi.ac.uk/pride/url/to/P12345				

6.3.17 go_terms

Description:	A ']'-delimited list of GO accessions for this protein.				
Type:	String List				
	Sum	mary	Complete		
Mandatory	Quantification				
,	Identification				
Example:	PRT accession	go	terms		
Example.	PRH P12345	GO	:0006457	GO:0005759 GO:0005886 GO:0004069	

6.3.18 protein_coverage

Description:	A value between 0 and 1 defining the protein coverage.				
Type:	Double				
		Summary	Complete		
Mandatory	Quantification	-	√		
,	Identification		✓		
Example:	PRT accession	on pr	otein_cov	erage	
Example.	PRH P12345	0.	4		

6.3.19 protein_abundance_assay[1-n]

Description:	The protein's abundance as measured in the given assay through whatever technique was employed.			
Type:	Double			
Mandatory	Summary Complete Quantification (✓) ✓ Identification mandatory if quantification data is provided for assays			
Example:	PRT accession protein_abundance_assay[1] protein_abundance_assay[2] PRH P12345 0.4			

6.3.20 protein_abundance_study_variable[1-n]

Description:	The protein's abundance as measured in the given Study Variable, for					
Description.	example mean or median of quantitative values reported in Assays.					
Type:	Double					
		Summary	Complete			
Mandatory	Quantification	✓	✓			
	Identification					
	PRT accession protein_abundance_study_variable[1] protein_abundance_study_variable[2]					
Example:	protein_abundance_study_variable[2]					
İ	PRH P12345	0.	. 4	0.2		

6.3.21 protein_abundance_stdev_study_variable[1-n]

Description:	The standard deviation of the protein's abundance. If a protein's abundance is given for a certain study variable, the corresponding standard deviation column MUST also be present (in case the value is not available "null" should be used).				
Type:	Double				
Mandatory	Summary Complete Quantification (✓)¹ (✓)¹				

	Identification
	¹ mandatory if protein abundance study variable reported
Evample:	PRT accession protein_abundance_stdev_study_variable[1]
Lxample.	PRH P12345 0.4

6.3.22 protein_abundance_std_error_study_variable [1-n]

Description:	The standard error of the protein's abundance. If a protein's abundance is given for a certain study variable, the corresponding standard error column MUST also be present (in case the value is not available "null" should be used).			
Type:	Double			
Mandatory	Summary Complete Quantification (\(\sigma\)^1 (\(\sigma\)^1 Identification			
Example:	PRT accession protein_abundance_sub[1] protein_abundance_std_error_sub[1] PRH P12345 0.4 0.03			

6.3.23 opt_global_*

Description:	Additional columns can be added to the end of the protein table. These column headers MUST start with the prefix "opt_" followed by the identifier of the object they reference: assay, study variable, MS run or "global" (if the value relates to all replicates). Column names MUST only contain the following characters: 'A'-'Z', 'a'-'z', '0'-'9', '_', '-', '[', ']', and ':'. CV parameter accessions MAY be used for optional columns following the format: opt_{OBJECT_ID}_cv_{accession}_{parameter}. Spaces within the parameter's name MUST be replaced by '_'.				
Туре:	Column				
Mandatory	Summary Complete Quantification Identification Identi				
Example:	RT accession opt_assay[1]_my_value opt_global_another_value RH P12345 My_value about assay[1] some other value that is across reps				

6.4 Peptide Section

The peptide section is table based. The peptide section must always come after the metadata section and or protein section if these are present in the file. All table columns MUST be tab separated. There MUST NOT be any empty cells. Missing values MUST be reported using "null". Most columns are mandatory. The order of columns is not specified although for ease of human interpretation, it is RECOMMENDED to follow the order specified below.

6.4.1 sequence

Description:	The peptide	e's seque	ence		
Туре:	String				
		Summary	Complete		
	Quantification	√	√		
Mandatory	Identification	1	1		
	¹ Not recommend	ded in identific	cation only file		
	PEH sequenc	е			
Example:	PEP KVPQVST	PTLVEVSR			
	PEP EIEILAC	EIR			

6.4.2 accession

Description	The protein's accession the peptide is associated with. In case no protein
Description.	section is present in the file or the peptide was not assigned to a protein the

	field should be filled with "null". If the peptide can be assigned to more than one protein, multiple rows SHOULD be provided for each peptide to protein			
	l '. '. '	G IOWS C	or loot be provided for each peptide to protein	
	mapping.			
Туре:	String			
	Summary	Complete		
1	Quantification ✓	✓		
Mandatory	Identification 1	1		
	¹Not recommended in identification only files			
Example:		accession		
LAGITIPIE.	PEP KVPQVSTPTLVEVSR	P02768		

6.4.3 unique

Description:	Indicates whether the peptide is unique for this protein in respect to the						
•	searched database.						
Type:	Boolean (0/1)						
	Summary	Complete					
	Quantification ✓	✓					
Mandatory	Identification 1	1					
	¹ Not recommended in identific	ation only files					
	PEH sequence	accession	unique				
Example:	PEP KVPQVSTPTLVEVSR	P02768	0				
•	PEP VFDEFKPLVEEPQNLIK	P02768	1				

6.4.4 database

Description:	The protein database used for the search (could theoretically come from a different species) and the peptide sequence comes from.					
Type:	String					
Mandatory	Summary Cor Quantification Identification ¹ Not recommended in identification	nplete √ 1 only files				
Example:	PEP KVPQVSTPTLVEVSR P0	cession unique database 2768 O UniProtKB 2768 1 UniProtKB				

6.4.5 database_version

Description:	The protein database's version – in case there is no version available (custom build) the creation / download (e.g., for NCBI nr) date should be given. Additionally, the number of entries in the database MAY be reported in round brackets after the version in the format: {version} ({#entries} entries), for example "2011-11 (1234 entries)".					
Type:	String					
		ımmary Compl	ete			
Manadatan	Quantification	✓ ✓				
Mandatory	Identification	1 1				
	¹ Not recommended in identification only files					
	PEH sequence	acce	ssion	unique database database_version		
Example:	PEP KVPQVSTPTLV	VEVSR P027	68 0	0 UniProtKB 2011_11		
1-	PEP VFDEFKPLVE	EPQNLIK P027	68 1	1 UniProtKB 2011_11		

6.4.6 search_engine

	Description:	A " " delimited list of search engine(s) used to identify this peptide. Search engines must be supplied as parameters.						
1	уре:	F	Parameter List					
N	M andatory	Summary Complete Quantification						
- 1			Identification					

	¹ Not	recommended in identifica	tion c	only files]
	PEH	sequence		search_engine	
Example:	PEP	KVPQVSTPTLVEVSR		[MS, MS:1001207, Mascot,] [MS, MS:1001208, Sequest,]	
•	PEP	VFDEFKPLVEEPQNLIK		[MS, MS:1001207, Mascot,]	

6.4.7 best_search_engine_score

Description:	A " " delimited list of best search engine score(s) for the given peptide across all replicates. Scores SHOULD be reported using CV parameters whenever possible.				
Type:	Parameter List				
	Summary Complete				
Mandatory	Quantification ✓ Identification 1 1				
	¹ Not recommended in identification only files				
Example:	PEH sequence best_search_engine_score PEP KVPQVSTPTLVEVSR [MS,MS:1001155,Sequest:xcorr,2] PEP VFDEFKPLVEEPQNLIK [MS,MS:1001171,Mascot score,47.2]				

6.4.8 search_engine_score_ms_run[1-n]

Description:	A " " delimited list of search engine score(s) for the given peptide from a given MS run. Scores SHOULD be reported using CV parameters whenever possible.					
Type:	Parameter List					
Mandatory	Summary Complete Quantification Identification Not recommended in identification only files					
Example:	PEH sequence search_engine_score_ms_run[1] PEP KVPQVSTPTLVEVSR [MS,MS:1001155,Sequest:xcorr,2] PEP VFDEFKPLVEEPQNLIK [MS,MS:1001171,Mascot score,47.2]					

6.4.9 reliability

Description:	The reliability of the given peptide identification. This must be supplied by the resource and has to be one of the following values: 1: high reliability 2: medium reliability 3: poor reliability Important: An identification's reliability is resource dependent.					
Type:	Integer					
Mandatory	Summary Complete Quantification Identification 1 1 Not recommended in identification only files					
Example:	PEH sequence reliability PEP KVPQVSTPTLVEVSR 3 PEP VFDEFKPLVEEPQNLIK 1					

6.4.10 modifications

	The peptide's modifications or substitutions. To further distinguish peptide
	terminal modifications, these SHOULD be reported at position 0 or <i>peptide</i>
	size + 1 respectively. For detailed information see the modifications section in
Description:	the protein table. If substitutions are reported, the "sequence" column MUST
-	contain the original, unaltered sequence. Note that in contrast to the PSM
	section, fixed modifications or modifications caused by the quantification
	reagent i.e. the SILAC labels/tags SHOULD NOT be reported. It is thus also

	expected that modification reliability scores will typically be reported at the PSM-level only.					
Type:	String					
Mandatory	Summary Quantification ✓ Identification 1 ¹Not recommended in identifi	Complete / 1 cation only file	es			
Example:	PEH sequence PEP KVPQVSTPTLVEVSR PEP VFDEFKPLVEEPQNLII	10-1	fications IOD:00412			

6.4.11 retention_time

	A ' '-separated list of time points. Semantics may vary on how retention times are reported. For quantification approaches, different exporters MAY wish to export the retention times of all spectra used for quantification (e.g. in MS ²					
Description:	approaches) or the centre point of the feature quantified for MS ¹ approaches. It is assumed that the reported value(s) are for a given "master" peptide from one assay only (and the unlabeled peptide in label-based approaches). If the exporter wishes to export values for all assays, this can be done using optional columns. Retention time MUST be reported in seconds. Otherwise, units MUST be reported in the Metadata Section ("colunit-peptide").					
Туре:	Double List					
Mandatory	Summary Complete Quantification					
Example:	PEH sequence retention_time PEP KVPQVSTPTLVEVSR 10.2 PEP VFDEFKPLVEEPQNLIK 15.8					

6.4.12 retention_time_window

Description:	exporter wishes to export values for all assays, this can be done using optional					
		columns. Retention time windows MUST be reported in seconds. Otherwise,				
	units MUST be reported in the Metadata Section ("colunit-peptide").					
Type:	Double List					
		Summary	Complete			
Mondotoni	Quantification	✓	✓			
Mandatory	Identification	1	Т			
	¹ Not recommended in identification only files					
Example:	PEH sequence retention_time_window					
Example.	PEP KVPQVSTPT	LVEVSR	112	3.2 1145.3		

6.4.13

6.4.14 charge

Description:	The charge assigned by the search engine/software. In case multiple charge states for the same peptide are observed these should be reported as distinct entries in the peptide table. In case the charge is unknown "null" MUST be used.
Type:	Integer

			Summary	Comp	olete				 	
	- 1 [Quantification	✓	✓	•					
Mandatory	IJΓ	Identification	1	1						
	1	¹ Not recommended in identification only files								
	P	PEH sequence	9		charge					
Example:	P	PEP KVPQVSTI	PTLVEVSR		2					
	P	PEP VFDEFKP	LVEEPQNLIK	· ···	3					

6.4.15 mass_to_charge

Description:	The precursor's experimental mass to charge (<i>m/z</i>). It is assumed that the reported value is for a given "master" peptide from one assay only (and the unlabeled peptide in label-based approaches). If the exporter wishes to export values for all assays, this can be done using optional columns.					
Type:	Double					
Mandatory	Summary Complete Quantification					
	¹Not recommended in identification only files PEH seguence mass to charge					
Example:	PEH sequence mass_to_charge PEP KVPQVSTPTLVEVSR 1234.4 PEP VFDEFKPLVEEPQNLIK 123.4					

6.4.16 uri

Description:	A URI pointing to the peptide's entry in the experiment it was identified in (e.g., the peptide's PRIDE entry).				
Type:	URI				
		Summary	Complete		
Mandatary	Quantification	1	4		
Mandatory	Identification	'	'		
	¹ Not recommend	ed in identific	cation only fil	es	
_	PEH sequence	9	uri		
Example:	PEP KVPQVSTI	PTLVEVSR		p://www.ebi.ac.uk/pride/link/to/peptide	
	PEP VFDEFKPI	LVEEPQNLIK	htt	p://www.ebi.ac.uk/pride/link/to/peptide	

6.4.17 spectra_ref

Description:	Reference to spectra in a spectrum file. It is expected that spectra_ref SHOULD only be used for MS ² -based quantification approaches, in which retention time values cannot identify the spectra used for quantitation. The reference must be in the format ms_run[1-n]:{SPECTRA_REF} where SPECTRA_REF MUST follow the format defined in 5.2. Multiple spectra MUST be referenced using a " " delimited list.					
Type:	String					
Mandatory	Summary Complete Quantification (\checkmark) ² Identification 1 1 Not recommended in identification only files 2 Mandatory only if MS2 based quantification is used					
Example:	PEH sequence spectra_ref PEP KVPQVSTPTLVEVSR ms_run[1]:index=5 PEP VFDEFKPLVEEPQNLIK ms_run[2]:index=7 ms_run[2]:index=9					

6.4.18 peptide_abundance_assay[1-n]

Description:	The peptide's abundance in the given assay.						
Туре:	Double	Double					
		Summary	Complete				
Mandatory	Quantification		✓				
wandator y	Identification	1	1				

	¹ Not recommended in identification only files ² If quantification data is reported on assays level				
Example:	PEH	sequence		peptide_abundance_assay[1] peptide_abundance_assay[2]	
	PEP	KVPQVSTPTLVEVSR		0.4	

6.4.19 peptide_abundance_study_variable[1-n]

Description:	The peptide's abundance in the given study variable, for example calculated as an average of assay values.					
Type:	Double					
Mandatory	Summary Complete Quantification Identification Summary Complete Quantification Identification Ident					
Mandator y	Not recommended in identification only files mandatory if study variables are reported					
Example:	PEH sequence peptide_abundance_study_variable[1] PEP KVPQVSTPTLVEVSR 0.4					

6.4.20 peptide_abundance_stdev_study_variable[1-n]

Description:	The standard deviation of the peptide's abundance for a given study variable.					
Туре:	Double					
Mandatory	Summary Complete Quantification (\checkmark) ² (\checkmark) ² Identification 1 1					
manadioi y	¹ Not recommended in identification only files ² mandatory if peptide_abundance_study_variable reported					
Example:	PEH sequence peptide_abundance_sub[1] peptide_abundance_stdev_sub[1] PEP KVPQVSTPTLVEVSR 0.4 0.2					

6.4.21 peptide_abundance_std_error_sub[1-n]

Description:	The standard error of the peptide's abundance for a given study variable.					
Туре:	Double					
	Summary Complete					
	Quantification $(\checkmark)^2$ $(\checkmark)^2$					
Mandatory	Identification 1 1					
	Not recommended in identification only files					
	² mandatory if peptide_abundance_study_variable reported					
Example:	PEH sequence peptide abundance sub[1] peptide abundance std error sub[1]					
Ехапіріе.	PEP KVPQVSTPTLVEVSR 0.4 0.2					

6.4.22 opt_global_*

Description:	Additional columns can be added to the end of the peptide table. These column headers MUST start with the prefix "opt_" followed by the identifier of the object they reference: assay, study variable, MS run or "global" (if the value relates to all replicates). Column names MUST only contain the following characters: 'A'-'Z', 'a'-'z', '0'-'9', '_', '-', '[', ']', and ':'. CV parameter accessions MAY be used for optional columns following the format: opt_{OBJECT_ID}_cv_{accession}_{parameter name}. Spaces within the parameter's name MUST be replaced by ' '.					
Туре:	Column					
Mandatory	Summary Complete Quantification					
Example:	PRT accession opt_assay[1]_my_value opt_global_another_value PRH P12345 My value about assay[1] some other value that is across reps					

6.5 PSM Section

The PSM section is table-based. The PSM section MUST always come after the metadata section, peptide section and or protein section if they are present in the file. All table columns MUST be tab separated. Missing values MUST be reported using "null". Most columns are mandatory. The order of columns is not specified although for ease of human interpretation, it is RECOMMENDED to follow the order specified below.

6.5.1 sequence

Description:	The peptide's sequence corresponding to the PSM					
Type:	String					
	Summary		Complete			
Mandatory	Quantification	✓	√			
	Identification	✓	√			
Example:	PSH sequence	9				
	PSM KVPQVSTI	PTLVEVSR				
	PSM EIEILACE	EIR				

6.5.2 **PSM_ID**

Description:	multiple pro	A unique identifier for a PSM within the file. If a PSM can be matched to multiple proteins, the same PSM should be represented on multiple rows with different accessions and the same PSM_ID.									
Type:	Integer	iteger									
Mandatory	Quantification Identification	Summary ✓	Complete ✓								
Example:	PSH sequence	PTLVEVSR 2 PO	1 P02	ccession 768 							

6.5.3 accession

Description:	PSM) is as peptide wa PSM can b	The protein's accession the corresponding peptide sequence (coming from the PSM) is associated with. In case no protein section is present in the file or the peptide was not assigned to a protein the field should be filled with "null". If the PSM can be assigned to more than one protein, the same PSM should be represented on multiple rows with the same unique identifier.									
Type:	String										
		Summary	Complete								
Mandatory	Quantification	✓	✓								
	Identification	✓	✓								
Example:	PSH sequence		accession								
p.o.	PSM KVPQVST	PTLVEVSR	PU2768								

6.5.4 unique

Description:		ndicates whether the peptide sequence (coming from the PSM) is unique for his protein in respect to the searched database.									
Туре:	Boolean (0	Boolean (0/1)									
		Summary	Complete								
Mandatory	Quantification <		√								
	Identification ✓		✓								
	PSH sequenc	PSH sequence		n u	nique						
Example:	PSM KVPQVSTPTLVEVSR		P02768	0							
•	PSM VFDEFKP	LVEEPQNLIK	P02768	1							

6.5.5 database

Description: The protein database used for the search (could theoretically come from a

	different sp	different species) and the peptide sequence comes from.										
Туре:	String	String										
		Summary	Complete									
Mandatory	Quantification	✓	✓									
	Identification ✓		✓									
	PSH sequenc	е	accessior	n unique database …								
Example:	PSM KVPQVSTPTLVEVSR		P02768	0 UniProtKB								
•	PSM VFDEFKP	LVEEPQNLIK	P02768	1 UniProtKB								

6.5.6 database_version

Description:	build) the c Additionally brackets af	The protein database's version – in case there is no version available (custom build) the creation / download (e.g., for NCBI nr) date should be given. Additionally, the number of entries in the database MAY be reported in round brackets after the version in the format: {version} ({#entries} entries), for example "2011-11 (1234 entries)".										
Туре:	String											
	† ! !	Summary	Complete									
Mandatory	Quantification	✓										
	Identification	✓	✓									
	PSH sequence		accession		unique database database_version							
Example:	PSM KVPQVSTI	PTLVEVSR	P02768	0	UniProtKB 2011_11							
İ	PSM VFDEFKP	P02768	1	UniProtKB 2011_11								

6.5.7 search_engine

Description:	A " " delimited list of search engine(s) used to create the PSM. Search engines									
	must be supplied as parameters.									
Type:	Parameter List	arameter List								
	Summary	Complete								
Mandatory	Quantification <	✓								
	Identification ✓	✓								
	PSH sequence		rch_engine							
Example:	PSM KVPQVSTPTLVEVSR	-	,MS:1001207,Mascot,] [MS,MS:1001208,Sequest,]							
· -	PSM VFDEFKPLVEEPQNL:	[K [MS	[MS, MS:1001207, Mascot,]							

6.5.8 search_engine_score

Description:	A " " delimi	A " " delimited list of search engine score(s) for the given PSM.									
Туре:	Parameter	Parameter List									
		Summary	Complete								
Mandatory	Quantification	✓	✓								
,	Identification	✓	✓								
	PSH sequenc	е	bes	t_search_engine_score							
Example:	PSM KVPQVSTPTLVEVSR			,MS:1001155,Sequest:xcorr,2]							
	PSM VFDEFKP	LVEEPQNLIK	[MS	,MS:1001171,Mascot score,47.2]							

6.5.9 reliability

Description:	The reliability of the given PSM. This must be supplied by the resource and has to be one of the following values: 1: high reliability 2: medium reliability 3: poor reliability Important: An identification's reliability is resource dependent.									
Туре:	Integer									
Mandatory	Summary Complete Quantification Identification Identi									
Example:	PSH sequence reliability PSM KVPQVSTPTLVEVSR 3									

,			 	
	PSM	VFDEFKPLVEEPQNLIK	 1	

6.5.10 modifications

Description:	distinguish position 0 o modificatior "sequence" Note that in and fixed m	The peptide's (coming from the PSM) modifications or substitutions. To further distinguish peptide terminal modifications, these SHOULD be reported at position 0 or <i>peptide size</i> + 1 respectively. For detailed information see the modifications section in the protein table. If substitutions are reported, the "sequence" column MUST contain the original, unaltered sequence. Note that in contrast to the PRT and PEP section all modifications (variable and fixed modifications, including those induced by quantification reagents) MUST BE reported in the PSM section.									
Type:	String										
Mandatory	Quantification Identification	Summary ✓	Complete ✓								
Example:	PSH sequence PSM KVPQVSTE PSM VFDEFKPI		10[1	ifications MS,MS:100xxxx,Probability Score Y,0.8]-MOD:00412 L							

6.5.11 retention_time

Description:	allowed in c the PSM. It	ase mul MUST b	tiple spe e reporte	ectrum. A ' '-separated list of multiple time points is ctra were combined by the search engine to make ed in seconds. Otherwise, the units MUST be ction ('columnit_psm').
Type:	Double List			
Mandatory	Quantification Identification	Summary ✓	Complete ✓	
Example:	PSH sequence PSM KVPQVSTE PSM VFDEFKPI		10.2	-

6.5.12 charge

Description:	The ch	The charge assigned by the search engine/software.										
Type:	Intege	nteger										
			Summary	Com	olete							
Mandatory	Quantifi	Quantification <		✓								
	Identific	ation	✓	✓	<i>'</i>							
	PSH sequence			char	rge							
Example:	PSM KV	PQVSTE	TLVEVSR		2							
•	PSM VF	DEFKPI	VEEPQNLIK		3							

6.5.13 exp_mass_to_charge

Description:	The PSM's	The PSM's experimental mass to charge (<i>m/z</i>).						
Type:	Double	Double						
		Summary	Complete	Complete				
Mandatory	Quantification	✓	✓					
	Identification	✓	✓					
	PSH sequence		mas	s_to_charge				
Example:	PSM KVPQVSTP	TLVEVSR	123	34.4				
	PSM VFDEFKPL	VEEPQNLIK	123	. 4		<u></u>		

6.5.14 calc_mass_to_charge

Description:	The PSM's calculated (theoretical) mass to charge (<i>m/z</i>).						
Type:	Double						
		Summary	Complete				
Mandatory	Quantification	✓	✓				
,	Identification	✓	✓				

Ī		PSH	sequence	 mass_to_charge	
1	Example:	PSM	KVPQVSTPTLVEVSR	 1234.4	
İ	•	PSM	VFDEFKPLVEEPQNLIK	 123.4	

6.5.15 uri

Description:	tion: A URI pointing to the PSM's entry in the experiment it was identified in (e.g					
	the peptide	's PRIDE	entry).			
Type:	URI					
		Summary	Complete			
Mandatory	Quantification					
	Identification					
	PSH sequenc	е	uri	=		
Example:	PSM KVPQVST	PTLVEVSR		<pre>tp://www.ebi.ac.uk/pride/link/to/peptide</pre>		
•	PSM VFDEFKP	LVEEPQNLIK	htt	<pre>tp://www.ebi.ac.uk/pride/link/to/peptide</pre>		

6.5.16 spectra_ref

Description:	format ms_ the format delimited li	to a spectrum in a spectrum file. The reference must be in the run[1-n]: {SPECTRA_REF} where SPECTRA_REF MUST follow defined in 5.2. Multiple spectra MUST be referenced using a " " ist for the (rare) cases in which search engines have combined ectra to make identifications.				
Туре:	String					
Mandatory	Quantification Identification	Summary ✓	Complete ✓			
Example:			ms_	ctra_ref run[1]:index=5 run[2]:index=7 ms run[2]:index=9		

6.5.17 pre

Description:	Amino acid preceding the peptide (coming from the PSM) in the protein sequence. If unknown "null" MUST be used, if the peptide is N-terminal "-" MUST be used.						
Type:	String						
		Summary	Complete				
Mandatory	Quantification	✓	✓				
	Identification	✓	✓				
	PSH sequence		pre	post			
Example:		PTLVEVSR		D			
	PSM VFDEFKPI	LVEEPQNLIK	R	L			

6.5.18 post

Description:	Amino acid following the peptide (coming from the PSM) in the protein sequence. If unknown "null" MUST be used, if the peptide is C-terminal "-" MUST be used.						
Type:	String						
		Summary	Complete				
Mandatory	Quantification	✓	✓				
	Identification	✓	✓				
Example:	PSH sequence		pre	post			
	PSM KVPQVST	PTLVEVSR	K	D			
*	PSM VFDEFKP:	LVEEPQNLIK	R	L			

6.5.19 start

Description:	The start position of the peptide (coming from the PSM) within the protein, counting 1 as the N-terminus of the protein.	
Type:	String	
Mandatory	Summary Complete	

		Qι	antification	✓	✓			
		Ide	entification	✓	✓	•		
[PSH	sequence	Э		start	end	
	Example:	PSM	KVPQVST	PTLVEVSR		45	57	
<u> </u>	•	PSM	VFDEFKP:	LVEEPQNLIK		34	46	

6.5.20 end

Description: The end position of the peptide (coming from the PSM) within									
Description.	The end position of the peptide (coming from the PSM) within the prote counting 1 as the N-terminus of the protein.								
Type:	String	String							
	Summary	Complete							
Mandatory	Quantification ✓	✓							
	Identification ✓	✓							
_	PSH sequence	sta	art end						
Example:	PSM KVPQVSTPTLVEVSR	45	57						
· •	PSM VFDEFKPLVEEPQNL:	IK 34	46						

6.5.21 opt_global_*

Description:	Additional columns can be added to the end of the PSM table. The neaders MUST start with the prefix "opt_" followed by the identified object they reference: assay, study variable, MS run or "global" (if relates to all replicates). Column names MUST only contain the for characters: 'A'-'Z', 'a'-'z', '0'-'9', '_', '-', '[', ']', and ':'. CV parameter MAY be used for optional columns following the format: opt_{OBJECT_ID}_cv_{accession}_{parameter name}. Spaces with parameter's name MUST be replaced by '_'.	of the the value llowing accessions
Type:	Column	
Mandatory	Summary Complete Quantification Identification	
Example:	PSH sequence opt_assay[1]_my_value opt_global_another_value PSM PEPTIDER My value about assay[1] some other value that	is across reps

6.6 Small Molecule Section

The small molecule section is table-based. The small molecule section MUST always come after the metadata section, peptide section and or protein section if they are present in the file. All table columns MUST be Tab separated. There MUST NOT be any empty cells. Missing values MUST be reported using "null". Most columns are mandatory. The order of columns is not specified although for ease of human interpretation, it is RECOMMENDED to follow the order specified below.

6.6.1 identifier

Description:	A list of " " separated possible identifiers for these small molecules. The database identifier must be preceded by the resource description followed by a colon (in case this is not already part of the identifier format).						
Type:	String List						
		Summary	Complete				
Mandatory	Quantification	✓	✓				
	Identification	✓	✓				
	SMH identif	ier "					
Example:	SML CID:000	27395	•				
	SML HMDB:HM	DB12345					

6.6.2 chemical_formula

Description	The chemical formula of the identified compound.
Description:	This should be specified in Hill notation (EA Hill 1900), i.e. elements in the

	omitted. Ele "CO" vs. "C Charge sta positive and	ements s co"). The te is repo d negativ	hould be chemical orted by the e mode r	ically all other elements. Counts of one may be capitalized properly to avoid confusion (e.g., formula reported should refer to the neutral form. he charge field. This permits the comparison of esults. ne would be encoded by the string "C8H15NO6"
}	+	· acciyi	<u> </u>	no would be encoded by the earning control to
Туре:	String	i dooty is	,	no would be discussed by the caming controlled
	+	Summary	Complete	
	+			
Type: Mandatory	String			
	String Quantification	Summary ✓ ier	Complete ✓	al_formula

6.6.3 smiles

Description:	The molecules structure in the simplified molecular-input line-entry system (SMILES). If there are more than one SMILES for a given small molecule, use the " " separator.					
Type:	String List					
		Summary	Complete	9		
Mandatory	Quantification	✓	✓			
	Identification	✓	✓			
Example:	SMH identif	ier "	. chemical	al_formula smiles		
Example.	SML CID:000	27395	. C17H20N4	V402 C1=CC=C(C=C1)CCNC(=0)CCNNC(=0)C2=CC=NC=C2		

6.6.4 inchi_key

Description:	The standard IUPAC International Chemical Identifier (InChI) Key of the given substance. If there are more than one InChI identifier for a given small molecule, use the " " separator.						
Type:	String List						
		Summary	Complete				
Mandatory	Quantification	✓	✓				
	Identification	✓	✓				
Example:	SMH identif:			_formula inchi_key			
Lxample.	SML CID:0002	27395	. C17H20N4	02 QXBMEGUKVLFJAM-UHFFFAOYSA-N			

6.6.5 description

Description:	The small molecule's description / name.				
Type:	String				
		Summary	Complete		
Mandatory	Quantification	✓	✓		
	Identification	✓	✓		
Example:	SMH identif:	ier de	scription		
схапіріе.	SML CID:000	27395 N-	(2-phenyl	ethyl)-3-[2-(pyridine-4-carbonyl)hydrazinyl]propanamide	

6.6.6 exp_mass_to_charge

	Description:	The small mo	The small molecule's experimental mass to charge (<i>m/z</i>).				
	Туре:	Double	Double				
ľ		S	ummary	Complete			
l	Mandatory	Quantification	✓	✓			
L		Identification	✓	✓			
[Example:	SMH sequence		mass	s_to_charge		
į,	<u>-ханіріс.</u>	SMM CID:000273	395	123	34.4		

6.6.7 calc_mass_to_charge

Description:	The small molecule's precursor's calculated (theoretical) mass to charge ratio.	
Type:	Double	

			Summary	Complete	
Mandatory	Qua	ntification	✓	✓	
,	Iden	tification	✓	✓	
	SMH	identif	ier "	. mass to	
Example:	SML	CID:0002	27395	1234.5	

6.6.8 charge

Description:	The charge as	The charge assigned by the search engine/software.					
Type:	Integer						
	Su	ımmary	Complete				
Mandatory	Quantification	✓	✓				
	Identification	✓	✓				
Example:	SMH identifier		charge				
Ехапіріе.	SML CID:0002739	95	. 2				

6.6.9 retention_time

Description:	A ' '-separated list of time points. Semantics may vary. This time should refer to the small molecule's retention time if determined or the mid point between the first and last spectrum identifying the small molecule. It MUST be reported in seconds. Otherwise, the corresponding unit MUST be specified in the Metadata Section ('columnit_smallmolecule').					
Type:	Double List					
Mandatory	Quantification Identification	mmary Complete ✓ ✓				
Example:	SMH identifier SML CID:0002739!		tion_time 11.5			

6.6.10 taxid

Description:	The taxonomy id coming from the NEWT taxonomy for the species (if applicable).					
Type:	Integer					
		Summary	Complete			
Mandatory	Quantification	✓	✓			
	Identification	✓	✓			
Example:	SMH identif	ier	taxid			
Example.	SML CID:000	27395	null			

6.6.11 species

Description:	The species as a human readable string (if applicable).					
Туре:	String	String				
Mandatory	Sum Quantification	mary Complete				
marraator y	Identification	✓				
Example:	SMH identifier SML CID:00027395	specie null	es 			

6.6.12 database

Description:	Generally references the used spectral library (if applicable).					
Туре:	String	String				
	Sum	mary Complete				
Mandatory	Quantification	√				
	Identification	✓				
Evample:	SMH identifier	datab	ase			
Example:	SML CID:00027395	name	of used database			

6.6.13 database_version

Description: Either the version of the used database if available or otherwise the date of

Type:	creation. Additionally, the number of entries in the database MAY be reported in round brackets after the version in the format: {version} ({#entries} entries), for example "2011-11 (1234 entries)". String				
Mandatory	Quantification Identification	Summary ✓	Complete ✓		
Example:	SMH identifi SML CID:0002		databa 2011-1	ase version 12-22	

6.6.14 reliability

Description:	The reliability of the given small molecule identification. This must be supplied by the resource and has to be one of the following values: 1: high reliability 2: medium reliability 3: poor reliability Important: An identification's reliability is resource dependent.				
Type:	Integer				
Mandatory	Summary Complete Quantification Identification				
Example:	SMH identifier reliability SML CID:00027395 3				

6.6.15 uri

Description:	A URI pointing to the small molecule's entry in the experiment it was identified in (e.g., the small molecule's PRIDE entry).					
Type:	URI	URI				
		Summary	Complete			
Mandatory	Quantification	-				
	Identification					
Example:	SMH identifie	J	uri			
⊏xample.	SML CID:00027	7395	http://www	v.ebi.ac.uk/pride/link/to/identification		

6.6.16 spectra_ref

Description:	Reference to a spectrum in a spectrum file. The reference must be in the format ms_run[1-n]:{SPECTRA_REF} where spectra_ref MUST follow the format defined in 5.2. Multiple spectra can be referenced using a " " delimited list.				
Type:	String				
		Summary	Complete		
Mandatory	Quantification	✓	✓		
	Identification	✓	✓		
Example:	SMH identif:		spectra_re		
Lvailible.	SML CID:0002	27395	ms run[1]	:index=1002	

6.6.17 search_engine

Description:	A " " delimited list of search engine(s) used to identify this small molecule. Search engines must be supplied as parameters.				
Туре:	Parameter	List			
		Summary	Complete		
Mandatory	Quantification	✓	✓		
	Identification	✓	✓		
Example:	SMH identifi		search_en		
Lxailipie.	SML CID:0002	27395	[MS, MS:1	001477, SpectraST,]	

6.6.18 best_search_engine_score

Description:	A " " delimited list of best search engine score(s) across replicates for the given small molecule. Scores SHOULD be reported using CV parameters whenever possible.			
Type:	Parameter	List		
		Summary	Complete	
Mandatory	Quantification	√	✓	
	Identification	√	√	
Evample:	SMH identif	ier	search_en	gine_score
Example:	SML CID:0002	27395	[MS, MS:1	001419, SpectraST:discriminant score F, 0.7]

6.6.19 search_engine_score_ms_run[1-n]

	A " " delimited list of search engine score(s) in each MS run for the given small molecule. Scores SHOULD be reported using CV parameters whenever possible.				
Type:	Parameter List				
Mandatory	Summary Complete Quantification Identification Not recommended in identification only files				
Example:	SMH identifier search_engine_score SML CID:00027395 [MS, MS:1001419, SpectraST:discriminant score F, 0.7]				

6.6.20 modifications

Description:	The small molecule's modifications or adducts. The position of the modification must be given relative to the small molecule's beginning. The exact semantics of this position depends on the type of small molecule identified. In case the position information is unknown or not applicable it should not be supplied. For detailed information see protein table.				
Type:	String				
Mandatory	Summary Complete Quantification ✓ ✓ Identification ✓ ✓				
Example:					

6.6.21 smallmolecule_abundance_assay[1-n]

Description:	The small molecule's abundance in the given assays.				
Type:	Double				
	Summary Complete				
Mandatory	Quantification $(\checkmark)^1$ Identification				
	mandatory if assays are reported				
Example:	SMH identifier smallmolecule_abundance_assay[1]				
Example.	SML CID:00027395 0.3				

6.6.22 smallmolecule_abundance_study_variable[1-n]

Description:	The small molecule's abundance in the given study variables.					
Type:	Double	Double				
Mandatory		Summary	Complete			
Manuator y	Quantification	✓	✓			

	Ider	ntification	
Evample:	SMH	identifier	 smallmolecule_abundance_study_variable[1]
Lampie.	SML	CID:00027395	 0.3

6.6.23 smallmolecule_abundance_stdev_study_variable [1-n]

Description:	The standard deviation of the small molecule's abundance in the given study variable.
Type:	Double
Mandatory	Summary Complete Quantification Identification In case the abundance for a respective study variable is given the standard deviation column MUST also be present (in case the value is not available "null" MUST be used).
Example:	SMH identifier smallmolecule_abundance_sub[1] smallmolecule_abundance_stdev_sub[1] SML CID:00027395 0.3 0.04

6.6.24 smallmolecule_abundance_std_error_study_variable[1-n]

Description:	The standard error of the small molecule's abundance in the given study variable.
Туре:	Double
Mandatory	Summary Complete Quantification Identification In case the abundance for a respective study variable is given the standard error column MUST also be present (in case the value is not available "null" MUST be used).
Example:	SMH identifier smallmolecule_abundance_std_error_sub[1] SML CID:00027395 0.04

6.6.25 opt_global_*

Description:	Additional columns can be added to the end of the small molecule table. These column headers MUST start with the prefix "opt_" followed by the identifier of the object they reference: assay, study variable, MS run or "global" (if the value relates to all replicates). Column names MUST only contain the following characters: 'A'-'Z', 'a'-'z', '0'-'9', '_', '-', '[', ']', and ':'. CV parameter accessions MAY be used for optional columns following the format: opt_{OBJECT_ID}_cv_{accession}_{parameter name}. Spaces within the parameter's name MUST be replaced by '_'.
Туре:	Column
Mandatory	Summary Complete Quantification Identification
Example:	SMH identifier … opt_assay[1]_my_value opt_global_another_value SML CID:00027395 … My value some other value

7. Non-supported use cases

There are a number of use cases that were discussed during the development process and it was decided that they are not explicitly supported in mzTab version 1.0. They may be implemented in future versions of the standard.

- Sequence Tag approaches.
- Grouped modification position scoring systems.

8. Conclusions

This document contains the specifications for using the mzTab format to represent results from peptide, small molecule and protein identification pipelines, in the context of a proteomics investigation. This specification constitutes a proposal for a standard from the Proteomics Standards Initiative. These artefacts are currently undergoing the PSI document process, which will result in a standard officially sanctioned by PSI.

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