

# mzTab cheat sheet

| Data types     |  |
|----------------|--|
| Type           | Format   |
| Parameter      | [{CV label}, {accession}, {name}, {value}]<br>[MS, MS:1001171, Mascot:score, 40.21]<br>[, ,A user parameter, The value]                    |
| Parameter List | "{" separated<br>[MS, MS:1001171, Mascot:score, 40.21] [, ,Another parameter,]   |
| String List    | " " separated<br>first string second string  |
| Modification   | {position} {reliability score} - {Modification identifier}<br>1 (0.5) -UNIMOD:4<br>10 (0.8)  11 (0.2) -CHEMMOD:+57.0215<br>23-CHEMMOD:+NA4 |
| Spec Ref       | ms_file[1-n]:{SPEC REF}<br>ms_file[2]:index=7 ms_file[2]:index=9   |

| MTD - Metadata section (Key-value)          |                |   |
|---|----------------|---|
| Field                                       | Type           | Description   |
| {UNIT_ID}-title                             | String         | The unit's title                                    |
| {UNIT_ID}-description                       | String         | The unit's description                              |
| {UNIT_ID}-sample_processing[1-n]            | Parameter List | Description of the sample processing.               |
| {UNIT_ID}-instrument[1-n]-source            | Parameter      | The instrument's source                             |
| {UNIT_ID}-instrument[1-n]-analyzer          | Parameter      | The instrument's analyzer                           |
| {UNIT_ID}-instrument[1-n]-detector          | Parameter      | The instrument's detector                           |
| {UNIT_ID}-software[1-n]                     | Parameter      | Analysis software used in the order it was used.    |
| {UNIT_ID}-false_discovery_rate              | Parameter List | False discovery rate(s)for the experiment.          |
| {UNIT_ID}-publication                       | String List    | Publication ids (pubmed / doi).                     |
| {UNIT_ID}-contact[1-n]-name                 | String         | Contact name.                                       |
| {UNIT_ID}-contact[1-n]-affiliation          | String         | Contact affiliation.                                |
| {UNIT_ID}-contact[1-n]-email                | String         | Contact's e-mail address.                           |
| {UNIT_ID}-uri                               | URI            | Points to the unit's source data.                   |
| {UNIT_ID}-mod                               | Parameter List | Modifications reported in the unit.                 |
| {UNIT_ID}-mod-probability_method            | Parameter      | Method used to report modification probabilities.   |
| {UNIT_ID}-quantification_method             | Parameter      | Quantification method used.                         |
| {UNIT_ID}-protein-quantification_unit       | Parameter      | Unit of protein quantification results.             |
| {UNIT_ID}-peptide-quantification_unit       | Parameter      | Unit of peptide quantification results.             |
| {UNIT_ID}-ms_file[1-n]-format               | Parameter      | Data format of the external MS data file.           |
| {UNIT_ID}-ms_file[1-n]-location             | URI            | Location of the external MS data file.              |
| {UNIT_ID}-ms_file[1-n]-id_format            | Parameter      | Identifier format of the external MS data file.     |
| {UNIT_ID}-custom                            | Parameter      | Additional parameters.                              |
| {UNIT_ID}-({SUB_ID})-species[1-n]           | Parameter      | Species of the unit / subsample.                    |
| {UNIT_ID}-({SUB_ID})-tissue[1-n]            | Parameter      | Tissue of the unit / subsample.                     |
| {UNIT_ID}-({SUB_ID})-cell_type[1-n]         | Parameter      | Cell type of the unit / subsample.                  |
| {UNIT_ID}-({SUB_ID})-disease[1-n]           | Parameter      | Disease state of the unit / subsample.              |
| {UNIT_ID}-({SUB_ID})-description            | String         | Description of the subsample.                       |
| {UNIT_ID}-({SUB_ID})-quantification_reagent | Parameter      | Quantification reagent used to label the subsample. |
| {UNIT_ID}-({SUB_ID})-custom                 | Parameter      | Additional parameters for the subsample.            |

| PRH/PRT - Protein section (Table based) |                   |  |
|---|-------------------|--|
| Column                                  | Type              | Description  |
| accession                               | String            | The protein's accession.                           |
| unit_id                                 | String            | The unit's id.                                     |
| description                             | String            | Human readable description (i.e. the name)         |
| taxid                                   | Integer           | NEWT taxonomy for the species.                     |
| species                                 | String            | Human readable species                             |
| database                                | String            | Name of the protein database.                      |
| database_version                        | String            | Version of the protein database.                   |
| search_engine                           | Parameter List    | Search engine(s) identifying the protein.          |
| search_engine_score                     | Parameter List    | Search engine(s) reliability score(s).             |
| reliability                             | Integer (1-3)     | Identification reliability.                        |
| num_peptides                            | Integer           | Number of PSMs assigned to the protein.            |
| num_peptides_distinct                   | Integer           | Distinct (sequence + modifications) # of peptides. |
| num_peptides_unambiguous                | Integer           | Distinct number of unambiguous peptides.           |
| ambiguity_members                       | String List (“,”) | Alternative protein identifications.               |
| modifications                           | Modification List | Modifications identified in the protein.           |
| uri                                     | URI               | Location of the protein's source entry.            |
| go_terms                                | String List (“,”) | List of GO terms for the protein.                  |
| protein_coverage                        | Double (0-1)      | Amount of protein sequence identified.             |
| protein_abundance_sub[1-n]              | Double            | Protein abundance in the subsample.                |
| protein_abundance_stdev_sub[1-n]        | Double            | Standard deviation of the protein abundance.       |
| protein_abundance_std_error_sub[1-n]    | Double            | Standard error of the protein abundance.           |
| opt_*                                   | Column            | Additional columns must start with “opt_”          |

| PEH/PEP - Peptide section (Table based) |                   |   |
|---|-------------------|---|
| Column                                  | Type              | Description                                   |
| sequence                                | String            | The peptide's sequence.                       |
| accession                               | String            | The protein's accession.                      |
| unit_id                                 | String            | The unit's id.                                |
| unique                                  | Boolean (1/0)     | Peptide is unique for the protein.            |
| database                                | String            | Name of the sequence database.                |
| database_version                        | String            | Version (and optionally # of entries).        |
| search_engine                           | Parameter List    | Search engine(s) that identified the peptide. |
| search_engine_score                     | Parameter List    | Search engine(s) score(s) for the peptide.    |
| reliability                             | Integer(1-3)      | Identification reliability for the peptide.   |
| modifications                           | Modification List | Modifications identified in the peptide.      |
| retention_time                          | Double List (“,”) | Time points in seconds. Semantics may vary.   |
| charge                                  | Double            | Precursor ion's charge.                       |
| mass_to_charge                          | Double            | Precursor ion's $m/z$ .                       |
| uri                                     | URI               | Location of the PSM's source entry.           |
| spectra_ref                             | Spec Ref          | Spectra identifying the peptide.              |
| peptide_abundance_sub[1-n]              | Double            | Peptide abundance in the subsample.           |
| peptide_abundance_stdev_sub[1-n]        | Double            | Peptide abundance standard deviation.         |
| peptide_abundance_std_error_sub[1-n]    | Double            | Peptide abundance standard error.             |
| opt_*                                   | Column            | Optional columns must start with “opt_”.      |

### SMH/SML Small molecule section (Table based)

| Column                                     | Type              | Description  |
|--|-------------------|--|
| identifier                                 | String            | The small molecule's identifier.                   |
| unit_id                                    | String            | The unit's id.                                     |
| chemical_formula                           | String            | Chemical formula of the identified compound.       |
| smiles                                     | String            | Molecular structure in SMILES format.              |
| inchi_key                                  | String            | InChi Key of the identified compound.              |
| description                                | String            | Human readable description (i.e. the name)         |
| mass_to_charge                             | Double            | The precursor ion's $m/z$ .                        |
| charge                                     | Double            | The precursor ion's charge.                        |
| retention_time                             | Double List (“,”) | List of time points in seconds.                    |
| taxid                                      | Integer           | NEWT taxonomy id of the species.                   |
| species                                    | String            | Human readable name of the species.                |
| database                                   | String            | Name of the used database.                         |
| database_version                           | String            | Version (and optionally # of compounds)            |
| reliability                                | Integer (1-3)     | The identification reliability.                    |
| uri  | URI               | The source entry's location.                       |
| spectra_ref                                | Spec Ref          | References to spectra identify the small molecule. |
| search_engine                              | Parameter List    | Search engine(s) identifying the small molecule.   |
| search_engine_score                        | Parameter List    | Search engine(s) identifications score(s).         |
| modifications                              | Modification List | Modifications identified on the small molecule.    |
| smallmolecule_abundance_sub[1-n]           | Double            | Abundance in the subsample.                        |
| smallmolecule_abundance_stdev_sub[1-n]     | Double            | Standard deviation of the abundance.               |
| smallmolecule_abundance_std_error_sub[1-n] | Double            | Standard error of the abundance.                   |
| opt_*                                      | Column            | Optional columns must start with “opt_”            |