### Summary

The summary file contains summary information for all the raw files processed with a single MaxQuant run. The summary information consists of some MaxQuant parameters, information of the raw file contents, and statistics on the peak detection. Based on this file a quick overview can be gathered on the quality of the data in the raw file.

The last row in this file contains the summary information for each column on each of the processed files.

Name	Separator	Description
Raw file		The raw file processed.
Experiment		Experiment name assigned to this LC-MS run in the experimental design.
Fraction		Fraction assigned to this LC-MS run in the experimental design.
Enzyme		The protease used to digest the protein sample.
Enzyme mode		The protease used to digest the protein sample.
Enzyme first search		The protease used for the first search.
Enzyme mode first search		The protease used for the first search.
Use enzyme first search		Marked with '+' when a different protease setup was used for the first search.
Variable modifications		The variable modification(s) used during the identification of peptides.
Fixed modifications		The fixed modification(s) used during the identification of peptides.
Multi modifications		The multi modification(s) used during the identification of peptides.
Variable modifications first search		The variable modification(s) used during the first search.
Use variable modifications first search		Marked with '+' when different variable modifications were used for the first search.
Requantify		The number of labels used.
Multiplicity		The number of labels used.
Max. missed cleavages		The maximum allowed number of missed cleavages.
Max. labeled AAs		The maximum allowed of labeled amino acids in a peptide amino acid sequence.
Labels0		The labels used in the labeling experiment. Allowed values for X: 0=light; 1=medium; 2=heavy label partner.
Labels1		The labels used in the labeling experiment. Allowed values for X: 0=light; 1=medium; 2=heavy label partner.
Labels2		The labels used in the labeling experiment. Allowed values for X: 0=light; 1=medium; 2=heavy label partner.
LC-MS run type		The type of LC-MS run. Usually it will be 'Standard' which refers to a conventional shotgun proteomics run with data-dependent MS/MS.
Time-dependent recalibration		When marked with '+', time-dependent recalibration was applied to improve the data quality.
MS		The number of MS spectra recorded in this raw file.
MS/MS		The number of MS/MS spectra recorded in this raw file.
MS3		The number of MS3 spectra recorded in this raw file.
MS/MS Submitted		The number of tandem MS spectra submitted for analysis.
MS/MS Submitted (SIL)		The number of tandem MS spectra submitted for analysis, where the precursor ion was detected as part of a labeling cluster.
MS/MS Submitted (ISO)		The number of tandem MS spectra submitted for analysis, where the precursor ion was detected as an isotopic pattern.
MS/MS Submitted (PEAK)		The number of tandem MS spectra submitted for analysis, where the precursor ion was detected as a single peak.
MS/MS Identified		The total number of identified tandem MS spectra.
MS/MS Identified (SIL)		The total number of identified tandem MS spectra, where the precursor ion was detected as part of a labeling cluster.
MS/MS Identified (ISO)		The total number of identified tandem MS spectra, where the precursor ion was detected as an isotopic pattern.
MS/MS Identified (PEAK)		The total number of identified tandem MS spectra, where the precursor ion was detected as a single peak.
MS/MS Identified [%]		The percentage of identified tandem MS spectra.
MS/MS Identified (SIL) [%]		The percentage of identified tandem MS spectra, where the precursor ion was detected as part of a labeling cluster.

MS/MS Identified (ISO) [%]	The percentage of identified tandem MS spectra, where the precursor ion was detected as an isotopic pattern.
MS/MS Identified (PEAK) [%]	The percentage of identified tandem MS spectra, where the precursor ion was detected as a single peak.
Peptide Sequences Identified	The total number of unique peptide amino acid sequences identified from the recorded tandem mass spectra.
Peaks	The total number of peaks detected in the full scans.
Peaks Sequenced	The total number of peaks sequenced by tandem MS.
Peaks Sequenced [%]	The percentage of peaks sequenced by tandem MS.
Peaks Repeatedly Sequenced	The total number of peaks repeatedly sequenced (i.e. 1 or more times) by tandem MS.
Peaks Repeatedly Sequenced [%]	The percentage of peaks repeatedly sequenced (i.e. 1 or more times) by tandem MS.
Isotope Patterns	The total number of detected isotope patterns.
Isotope Patterns Sequenced	The total number of isotope patterns sequenced by tandem MS.
Isotope Patterns Sequenced (z>1)	The total number of isotope patterns sequenced by tandem MS with a charge state of 2 or more.
Isotope Patterns Sequenced [%]	The percentage of isotope patterns sequenced by tandem MS.
Isotope Patterns Sequenced (z>1) [%]	The percentage of isotope patterns sequenced by tandem MS with a charge state of 2 or more.
Isotope Patterns Repeatedly Sequenced	The total number of isotope patterns repeatedly sequenced (i.e. 1 or more times) by tandem MS.
Isotope Patterns Repeatedly Sequenced [%]	The percentage of isotope patterns repeatedly sequenced (i.e. 1 or more times) by tandem MS.
Multiplets	The total number of detected labeling triplets.
Multiplets z=1	The total number of detected labeling triplets with a charge state of 1.
Multiplets z=2	The total number of detected labeling triplets with a charge state of 2.
Multiplets z=3	The total number of detected labeling triplets with a charge state of 3.
Multiplets z=4	The total number of detected labeling triplets with a charge state of 4.
Multiplets z=5	The total number of detected labeling triplets with a charge state of 5.
Multiplets z=6	The total number of detected labeling triplets with a charge state of 6.
Multiplets z=7	The total number of detected labeling triplets with a charge state of 7.
Multiplets Sequenced	The total number of labeling triplets sequenced by tandem MS.
Multiplets Sequenced [%]	The percentage of labeling triplets sequenced by tandem MS.
Multiplets Repeatedly Sequenced	The total number of labeling triplets repeatedly sequenced (i.e. 1 or more times) by tandem MS.
Multiplets Repeatedly Sequenced [%]	The percentage of labeling triplets repeatedly sequenced (i.e. 1 or more times) by tandem MS.
Multiplets Identified	The total number of labeling triplets identified.
Multiplets Identified [%]	The percentage of labeling triplets identified.
Recalibrated	When marked with '+', the masses taken from the raw file were recalibrated.
Av. Absolute Mass Deviation [ppm]	The average absolute mass deviation found comparing to the identification mass in parts per million.
Mass Standard Deviation [ppm]	The standard deviation of the mass deviation found comparing to the identification mass in parts per million.
Av. Absolute Mass Deviation [mDa]	The average absolute mass deviation found comparing to the identification mass in milli-Dalton.
Mass Standard Deviation [mDa]	The standard deviation of the mass deviation found comparing to the identification mass in milli-Dalton.

### Evidence

The evidence file combines all the information about the identified peptides and normally is the only file required for processing the results. Additional information about the peptides, modifications, proteins, etc. can be found in the other files by unique identifier linkage.

Name	Separator	Description
Sequence		The identified AA sequence of the peptide.
Length		The length of the sequence stored in the column 'Sequence'.
K Count		The number of instances of K contained within the sequence. The value for this can reliably be determined in the case of labeling partners based on the distance between the partners. These counts are used to solidify the peptide identification process.
R Count		The number of instances of R contained within the sequence. The value for this can reliably be determined in the case of labeling partners based on the distance between the partners. These counts are used to solidify the peptide identification process.
Modifications		Post-translational modifications contained within the identified peptide sequence.
Modified sequence		Sequence representation including the post-translational modifications (abbreviation of the modification in brackets before the modified AA). The sequence is always surrounded by underscore characters ('_').
Deamidation (NQ) Probabilities		Sequence representation of the peptide including PTM positioning probabilities ([01], where 1 is best match) for 'Deamidation (NQ)'.
Oxidation (M) Probabilities		Sequence representation of the peptide including PTM positioning probabilities ([01], where 1 is best match) for 'Oxidation (M)'.
Phospho (STY) Probabilities		Sequence representation of the peptide including PTM positioning probabilities ([01], where 1 is best match) for 'Phospho (STY)'.
Deamidation (NQ) Score Diffs		Sequence representation for each of the possible PTM positions in each possible configuration, the difference is calculated between the identification score with the PTM added to that position and the best scoring identification where no PTM is added to that position. When this value is negative, it is unlikely that the particular modification is located at this position.
Oxidation (M) Score Diffs		Sequence representation for each of the possible PTM positions in each possible configuration, the difference is calculated between the identification score with the PTM added to that position and the best scoring identification where no PTM is added to that position. When this value is negative, it is unlikely that the particular modification is located at this position.
Phospho (STY) Score Diffs		Sequence representation for each of the possible PTM positions in each possible configuration, the difference is calculated between the identification score with the PTM added to that position and the best scoring identification where no PTM is added to that position. When this value is negative, it is unlikely that the particular modification is located at this position.
Acetyl (Protein N-term)		The number of occurrences of the modification 'Acetyl (Protein N-term)'.
Deamidation (NQ)		The number of occurrences of the modification 'Deamidation (NQ)'.
Oxidation (M)		The number of occurrences of the modification 'Oxidation (M)'.
Phospho (STY)		The number of occurrences of the modification 'Phospho (STY)'.
Missed cleavages		Number of missed enzymatic cleavages.
Proteins		The identifiers of the proteins this particular peptide is associated with.
Leading proteins		The identifiers of the proteins in the proteinGroups file, with this protein as best match, this particular peptide is associated with. When multiple matches are found here, the best scoring protein can be found in the 'Leading Razor Protein' column.
Leading razor protein		The identifier of the best scoring protein, from the proteinGroups file this, this peptide is associated to.

Туре	The type of the feature. 'MSMS' for an MS/MS spectrum without an MS1 isotope pattern assigned. 'ISO-MSMS' MS1 isotope cluster identified by MS/MS. 'MULTI-MSMS' MS1 labeling cluster identified by MS/MS. 'MULTI-SECPEP' MS1 labeling cluster identified by MS/MS as second peptide. 'MULTI-MATCH' MS1 labeling cluster identified by matching between runs. In case of label-free data there is no difference between 'MULTI' and 'ISO'.
Labeling State	Labeling state of the precursor isotope pattern used to identify the peptide.
Raw file	The name of the RAW-file the mass spectral data was derived from.
Fraction	The fraction in which this peptide was detected.
Experiment	
MS/MS m/z	The m/z used for fragmentation (not necessarily the mono- isotopic m/z).
Charge	The charge-state of the precursor ion.
m/z	The recalibrated mass-over-charge value of the precursor ion.
Mass	The predicted monoisotopic mass of the identified peptide sequence.
Resolution	The resolution of precursor ion measured in Full Width at Half Maximum (FWHM).
Uncalibrated - Calibrated m/z [ppm]	The difference between the uncalibrated and recalibrated mass-over-charge value of the precursor ion measured in parts-per-million. This gives an indication of the mass drift in the original data, which was automatically corrected by MaxQuant.
Uncalibrated - Calibrated m/z [Da]	The difference between the uncalibrated and recalibrated mass-over-charge value of the precursor ion measured in parts-per-million. This gives an indication of the mass drift in the original data, which was automatically corrected by MaxQuant.
Mass error [ppm]	Mass error of the recalibrated mass-over-charge value of the precursor ion in comparison to the predicted monoisotopic mass of the identified peptide sequence in parts per million.
Mass error [Da]	Mass error of the recalibrated mass-over-charge value of the precursor ion in comparison to the predicted monoisotopic mass of the identified peptide sequence in milli-Dalton.
Uncalibrated mass error [ppm]	Mass error of the uncalibrated mass-over-charge value of the precursor ion in comparison to the predicted monoisotopic mass of the identified peptide sequence.
	Note: This column can contain missing values (denoted as NaN).
Uncalibrated mass error [Da]	Mass error of the uncalibrated mass-over-charge value of the precursor ion in comparison to the predicted monoisotopic mass of the identified peptide sequence.
	Note: This column can contain missing values (denoted as NaN).
Max intensity m/z 0	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Max intensity m/z 1	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Max intensity m/z 2	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Retention time	The uncalibrated retention time in minutes in the elution profile of the precursor ion.
Retention length	The total retention time length of the peak (last time point first time point).
Calibrated retention time	The recalibrated retention time in minutes in the elution profile of the precursor ion.
Calibrated retention time start	The recalibrated retention start in minutes in the elution profile of the precursor ion.
Calibrated retention time finish	The recalibrated retention finish in minutes in the elution profile of the precursor ion.
Retention time calibration	The difference in minutes between the uncalibrated and recalibrated retention time. This gives an indication of the retention time drift in the original data, which was automatically corrected by MaxQuant.
Match time difference	Note: This column can contain missing values (NaN).  When the option match between runs is used in MaxQuant, this value indicates the time difference between the feature from the raw file it was taken from and the feature from the raw file it was matched to.

Match m/z difference	When the option match between runs is used in MaxQuant, this value indicates the m/z difference between the feature
	from the raw file it was taken from and the feature from the raw file it was matched to.
Match q-value	This is the q-value for features that have been identified by matching between runs'.
Match score	The andromeda score of the MS/MS identification that is the source of this identification by 'matching between runs'.
Number of data points	The number of data points (peak centroids) collected for this peptide feature.
Number of scans	The number of MS scans that the 3d peaks of this peptide feature are overlapping with.
Number of isotopic peaks	The number of isotopic peaks contained in this peptide feature.
PIF	Short for Parent Ion Fraction; indicates the fraction the target peak makes up of the total intensity in the inclusion window.
Fraction of total spectrum	The percentage the ion intensity makes up of the total intensity of the whole spectrum.
Base peak fraction	The percentage the parent ion intensity in comparison to the highest peak in the MS spectrum.
PEP	Posterior Error Probability of the identification. This value essentially operates as a p-value, where smaller is more significant.
MS/MS count	The number of sequencing events for this sequence, which matches the number of identifiers stored in the column MS/MS IDs.
MS/MS scan number	The RAW-file derived scan number of the MS/MS with the highest peptide identification score (the highest score is stored in the column 'Score').
Score	Andromeda score for the best associated MS/MS spectrum.
Delta score	Score difference to the second best identified peptide.
Combinatorics	Number of possible distributions of the modifications over the peptide sequence.
Ratio M/L	The ratio between two medium and light label partners.
Ratio M/L normalized	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L shift	
Ratio H/L	The ratio between two heavy and light label partners.
Ratio H/L normalized	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L shift	
Ratio H/M	The ratio between two heavy and medium label partners.
Ratio H/M normalized	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M shift	
Intensity	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Reporter PIF	
Reporter fraction	
Reverse	When marked with '+', this particular peptide was found to be part of a protein derived from the reversed part of the decoy database. These should be removed for further data analysis.
Potential contaminant	When marked with '+', this particular peptide was found to be part of a commonly occurring contaminant. These should be removed for further data analysis.
id	A unique (consecutive) identifier for each row in the evidence table, which is used to cross-link the information in this file with the information stored in the other files.
Protein group IDs	The identifier of the protein-group this redundant peptide sequence is associated with, which can be used to look up the extended protein information in the file 'proteinGroups.txt'. As a single peptide can be linked to multiple proteins (e.g. in the case of razor-proteins), multiple ids can be stored here separated by a semicolon. As a protein can be identified by multiple peptides, the same id can be found in different rows.
Peptide ID  Mod. peptide ID	The identifier of the non-redundant peptide sequence.  Identifier of the associated modification summary stored in the
MO/MO ID-	file 'modificationSpecificPeptides.txt'.
MS/MS IDs	Identifier(s) of the associated MS/MS summary(s) stored in the file 'msms.txt'.

Best MS/MS	Identifier(s) of the best MS/MS associated spectrum stored in the file 'msms.txt'.
Deamidation (NQ) site IDs	Identifier(s) of the modification summary stored in the file 'Deamidation (NQ)Sites.txt'.
Oxidation (M) site IDs	Identifier(s) of the modification summary stored in the file 'Oxidation (M)Sites.txt'.
Phospho (STY) site IDs	Identifier(s) of the modification summary stored in the file 'Phospho (STY)Sites.txt'.

### **Peptides**

The peptides table contains information on the identified peptides in the processed raw-files.

Name	Separator	Description
Sequence		The amino acid sequence of the identified peptide.
N-term cleavage window		Sequence window from -15 to 15 around the N-terminal cleavage site of this peptide.
C-term cleavage window		Sequence window from -15 to 15 around the C-terminal cleavage site of this peptide.
Amino acid before		The amino acid in the protein sequence before the peptide.
First amino acid		The amino acid in the first position of the peptide sequence.
Second amino acid		The amino acid in the first position of the peptide sequence.
Second last amino acid		The amino acid in the last position of the peptide sequence.
Last amino acid		The amino acid in the last position of the peptide sequence.
Amino acid after		The amino acid in the protein sequence after the peptide.
A Count		The number of instances of the 'A' amino acid contained within the sequence.
R Count		The number of instances of the 'R' amino acid contained within the sequence.
N Count		The number of instances of the 'N' amino acid contained within the sequence.
D Count		The number of instances of the 'D' amino acid contained within the sequence.
C Count		The number of instances of the 'C' amino acid contained within the sequence.
Q Count		The number of instances of the 'Q' amino acid contained within the sequence.
E Count		The number of instances of the 'E' amino acid contained within the sequence.
G Count		The number of instances of the 'G' amino acid contained within the sequence.
H Count		The number of instances of the 'H' amino acid contained within the sequence.
I Count		The number of instances of the 'I' amino acid contained within the sequence.
L Count		The number of instances of the 'L' amino acid contained within the sequence.
K Count		The number of instances of the 'K' amino acid contained within the sequence.
M Count		The number of instances of the 'M' amino acid contained within the sequence.
F Count		The number of instances of the 'F' amino acid contained within the sequence.
P Count		The number of instances of the 'P' amino acid contained within the sequence.
S Count		The number of instances of the 'S' amino acid contained within the sequence.
T Count		The number of instances of the 'T' amino acid contained within the sequence.
W Count		The number of instances of the 'W' amino acid contained within the sequence.
Y Count		The number of instances of the 'Y' amino acid contained within the sequence.
V Count		The number of instances of the 'V' amino acid contained within the sequence.
U Count		The number of instances of the 'U' amino acid contained within the sequence.
O Count		The number of instances of the 'O' amino acid contained within the sequence.
Length		The length of the sequence stored in the column "Sequence".
Missed cleavages		Number of missed enzymatic cleavages.
Mass		Monoisotopic mass of the peptide.
Proteins		Identifiers of proteins this peptide is associated with.
Leading razor protein		Identifier of the leading protein in the protein group which uses this peptide for quantification. (Either unique or razor.)
Start position		Position of the first amino acid of this peptide in the protein sequence. (one-based)
End position		Position of the last amino acid of this peptide in the protein sequence. (one-based)

Unique (Proteins)  When marked with ***, this particular peptide is unique to a single protein sequence in the fast afte(s).  Charges  All charge states that have been observed.  Pester For Protability of the identification. This value essentially operates as a p-value, where smaller is more significant.  Score  Highest Andromeda score for the associated MSMNS peptral. Indicates whether this experiment was identified by MSMNS or only by matching between runs.  Identification type IB_30_6  Indicates whether this experiment was identified by MSMNS or only by matching between runs.  Identification type IB_30_6  Indicates whether this experiment was identified by MSMNS or only by matching between runs.  Identification type IB_30_6  Ide	Unique (Groups)	When marked with '+', this particular peptide is unique to a single protein group in the proteinGroups file.
PEP Posterior Error Probability of the identification. This value essentially operates as a p-value, where smaller is more significant.  Score Highest Andromeda score for the associated MS/MS spectra. Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Identification type IB_30_6 only by matching between runs.  Identification type IB_21_24 only by matching between runs.  Identification type IB_21_24 only by matching between runs.  Identification type IB_23_6.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Identification type IB_23_6.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Identification type KR_1_24 only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/	Unique (Proteins)	When marked with '+', this particular peptide is unique to a single protein sequence in the fasta file(s).
essentially operates as a jar value, where smaller is more significant.  Score Highest Andromeda score for the associated MS/MS spectra. Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Identification type IB_30_6 Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Identification type IB_21_24 Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Identification type IB_23_6 Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Identification type KR_1_24 Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Identification type KR_30_6 Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Identification type KR_30_6 Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Identification type KR_30_6 Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Fraction Average  Fraction 3  Fraction 3  Fraction 4  Fraction 5  Fraction 4  Fraction 5  Fraction 6  Fraction 6  Fraction 7  Fraction 8  Fraction 8  Fraction 9  Fraction 10  Fraction 10  Fraction 10  Fraction 11  Fraction 12  Fraction 13  Fraction 14  Fraction 15  Fraction 15  Fraction 16  Fraction 17  Fraction 20  Fraction 31  Fraction 33  Fraction 34  Fraction 35  Fraction 44  Fraction 45  Fraction 47  Fraction 49  Fraction 49  Fraction 49  Fraction 49  Fraction 50  Fraction 50  Fraction 51  Fraction 55  Fraction 56  Fraction 57  Fraction 57  Fraction 57  Fraction 57  Fraction 57  Fraction 57  Fraction 59	Charges	All charge states that have been observed.
Identification type IB_1_24  Identification type IB_30_6  Identification type IB_30_6  Identification type IB_30_6  Identification type IB_21_24  Identification type IB_23_6  Identification type KR_1_24  Identification type KR_1_24  Identification type KR_30_6  Identification type III.  Identification type KR_30_6  Identification type III.  Identificat	PEP	essentially operates as a p-value, where smaller is more
Identification type IB_1_24  Identification type IB_30_6  Identification type IB_30_6  Identification type IB_30_6  Identification type IB_21_24  Identification type IB_23_6  Identification type KR_1_24  Identification type KR_1_24  Identification type KR_30_6  Identification type III.  Identification type KR_30_6  Identification type III.  Identificat	Score	Highest Andromeda score for the associated MS/MS spectra.
only by matching between runs.  Identification type IB2_1_24  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Identification type KR_1_24  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Identification type KR_30_6  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Identification type KR_30_6  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was identified by MS/MS or only by matching between runs.  Indicates whether this experiment was	Identification type IB_1_24	Indicates whether this experiment was identified by MS/MS or
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Only by matching between runs.	Identification type IB2_30_6	
only by matching between runs.  Fraction Std. Dev. Fraction Std. Dev. Fraction 2 Fraction 3 Fraction 5 Fraction 6 Fraction 7 Fraction 8 Fraction 9 Fraction 10 Fraction 10 Fraction 12 Fraction 13 Fraction 14 Fraction 15 Fraction 16 Fraction 17 Fraction 18 Fraction 19 Fraction 19 Fraction 10 Fraction 10 Fraction 11 Fraction 12 Fraction 13 Fraction 14 Fraction 15 Fraction 16 Fraction 17 Fraction 18 Fraction 20 Fraction 31 Fraction 31 Fraction 32 Fraction 34 Fraction 34 Fraction 44 Fraction 45 Fraction 47 Fraction 48 Fraction 49 Fraction 69 Fraction 50 Fraction 50 Fraction 51 Fraction 52 Fraction 69 Fraction 69 Fraction 55 Fraction 69 Fraction 50 Fraction 50 Fraction 51 Fraction 52 Fraction 55 Fraction 55 Fraction 56 Fraction 57	Identification type KR_1_24	Indicates whether this experiment was identified by MS/MS or only by matching between runs.
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Fraction 41         Fraction 42         Fraction 43         Fraction 44         Fraction 45         Fraction 46         Fraction 47         Fraction 48         Fraction 50         Fraction 51         Fraction 52         Fraction 53         Fraction 54         Fraction 55         Fraction 56         Fraction 57         Fraction 58	Fraction 34	
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Fraction 60	
Fraction 71	
Fraction 72	
Fraction 73	
Fraction 74	
Fraction 75	
Experiment IB_1_24	Number of evidence entries for this 'Experiment'.
Experiment IB_30_6	Number of evidence entries for this 'Experiment'.
Experiment IB2_1_24	Number of evidence entries for this 'Experiment'.
Experiment IB2_30_6	Number of evidence entries for this 'Experiment'.
Experiment KR_1_24	Number of evidence entries for this 'Experiment'.
Experiment KR_30_6	Number of evidence entries for this 'Experiment'.
Ratio M/L	The ratio between two medium and light label partners.
Ratio M/L normalized	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L variability [%]	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type	
Ratio H/L	The ratio between two heavy and light label partners.
Ratio H/L normalized	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L variability [%]	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type	
Ratio H/M	The ratio between two heavy and medium label partners.
Ratio H/M normalized	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M variability [%]	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type	
Ratio M/L IB_1_24	The ratio between two medium and light label partners.
Ratio M/L normalized IB_1_24	The ratio between two medium and light label partners.
Ratio M/L variability [%] IB_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB_1_24	
Ratio H/L IB_1_24	The ratio between two heavy and light label partners.
Ratio H/L normalized IB_1_24	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L variability [%] IB_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB_1_24	
Ratio H/M IB_1_24	The ratio between two heavy and medium label partners.
Ratio H/M normalized IB_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M variability [%] IB_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.

Ratio H/M count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB_1_24	
Ratio M/L IB_30_6	The ratio between two medium and light label partners.
Ratio M/L normalized IB_30_6	The ratio between two medium and light label partners.
Ratio M/L variability [%] IB_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB_30_6	
Ratio H/L IB_30_6	The ratio between two heavy and light label partners.
Ratio H/L normalized IB_30_6	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L variability [%] IB_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB_30_6	
Ratio H/M IB_30_6	The ratio between two heavy and medium label partners.
Ratio H/M normalized IB_30_6	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M variability [%] IB_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB_30_6	
Ratio M/L IB2_1_24	The ratio between two medium and light label partners.
Ratio M/L normalized IB2_1_24	The ratio between two medium and light label partners.
Ratio M/L variability [%] IB2_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB2_1_24	
Ratio H/L IB2_1_24	The ratio between two heavy and light label partners.
Ratio H/L normalized IB2_1_24	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L variability [%] IB2_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB2_1_24	
Ratio H/M IB2_1_24	The ratio between two heavy and medium label partners.
Ratio H/M normalized IB2_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M variability [%] IB2_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB2_1_24	
Ratio M/L IB2_30_6	The ratio between two medium and light label partners.
Ratio M/L normalized IB2_30_6	The ratio between two medium and light label partners.
Ratio M/L variability [%] IB2_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.

Ratio M/L count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB2_30_6	
Ratio H/L IB2_30_6	The ratio between two heavy and light label partners.
Ratio H/L normalized IB2_30_6	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L variability [%] IB2_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB2_30_6	
Ratio H/M IB2_30_6	The ratio between two heavy and medium label partners.
Ratio H/M normalized IB2_30_6	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M variability [%] IB2_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB2_30_6	
Ratio M/L KR_1_24	The ratio between two medium and light label partners.
Ratio M/L normalized KR_1_24	The ratio between two medium and light label partners.
Ratio M/L variability [%] KR_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type KR_1_24	
Ratio H/L KR_1_24	The ratio between two heavy and light label partners.
Ratio H/L normalized KR_1_24	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L variability [%] KR_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type KR_1_24	
Ratio H/M KR_1_24	The ratio between two heavy and medium label partners.
Ratio H/M normalized KR_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M variability [%] KR_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type KR_1_24	
Ratio M/L KR_30_6	The ratio between two medium and light label partners.
Ratio M/L normalized KR_30_6	The ratio between two medium and light label partners.
Ratio M/L variability [%] KR_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type KR_30_6	
Ratio H/L KR_30_6	The ratio between two heavy and light label partners.
Ratio H/L normalized KR_30_6	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L variability [%] KR_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.

Ratio H/L count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type KR_30_6	
Ratio H/M KR_30_6	The ratio between two heavy and medium label partners.
Ratio H/M normalized KR_30_6	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M variability [%] KR_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type KR_30_6	
Intensity	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Intensity IB_1_24	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Intensity IB_30_6	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Intensity IB2_1_24	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB2_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB2_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB2_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Intensity IB2_30_6	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB2_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB2_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB2_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Intensity KR_1_24	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L KR_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M KR_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H KR_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.

Intensity KR_30_6	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L KR_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M KR_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H KR_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Reverse	When marked with '+', this particular peptide was found to be part of a protein derived from the reversed part of the decoy database. These should be removed for further data analysis.
Potential contaminant	When marked with '+', this particular peptide was found to be part of a commonly occurring contaminant. These should be removed for further data analysis.
id	A unique (consecutive) identifier for each row in the peptides table, which is used to cross-link the information in this table with the information stored in the other tables.
Protein group IDs	The identifiers of the protein groups this peptide was linked to, referenced against the proteinGroups table.
Mod. peptide IDs	Identifier(s) for peptide sequence(s), associated with the peptide, referenced against the corresponding modified peptides table.
Evidence IDs	Identifier(s) for analyzed peptide evidence associated with the protein group referenced against the evidence table.
MS/MS IDs	The identifiers of the MS/MS scans identifying this peptide, referenced against the msms table.
Best MS/MS	The identifier of the best (in terms of quality) MS/MS scan identifying this peptide, referenced against the msms table.
Deamidation (NQ) site IDs	Identifier(s) for site(s) associated with the protein group, which show(s) evidence of the modification, referenced against the appropriate modification site file.
Oxidation (M) site IDs	Identifier(s) for site(s) associated with the protein group, which show(s) evidence of the modification, referenced against the appropriate modification site file.
Phospho (STY) site IDs	Identifier(s) for site(s) associated with the protein group, which show(s) evidence of the modification, referenced against the appropriate modification site file.
MS/MS Count	

# Modification-specific peptides

Name	Separator	Description
Sequence		The identified AA sequence of the peptide.
K Count		The number of instances of the 'K' AA contained within the sequence. The value for this can reliably be determined in the case of SILAC partners based on the distance between the partners. These counts are used to solidify the peptide identification process.
R Count		The number of instances of the 'R' AA contained within the sequence. The value for this can reliably be determined in the case of SILAC partners based on the distance between the partners. These counts are used to solidify the peptide identification process.
Modifications		Post-translational modifications contained within the sequence. When no modifications exist, this is set to 'unmodified'.
Mass		Charge corrected mass of the precursor ion.
Mass Fractional Part		The values after the decimal point (ie value - floor(value)).
Protein Groups		IDs of the protein groups to which this peptide belongs.
Proteins		The identifiers of the proteins this particular peptide is associated with.
Unique (Groups)		When marked with '+', this particular peptide is unique to a single protein group in the proteinGroups file.
Unique (Proteins)		When marked with '+', this particular peptide is unique to a single protein sequence in the fasta file(s).
Acetyl (Protein N-term)		Number of Acetyl (Protein N-term) on this peptide.
Deamidation (NQ)		Number of Deamidation (NQ) on this peptide.
Oxidation (M)		Number of Oxidation (M) on this peptide.
Phospho (STY)		Number of Phospho (STY) on this peptide.
Missed cleavages		Number of missed enzymatic cleavages.
Identification type IB_1_24		Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Identification type IB_30_6		Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Identification type IB2_1_24		Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Identification type IB2_30_6		Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Identification type KR_1_24		Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Identification type KR_30_6		Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Fraction Average		
Fraction Std. Dev.		
Fraction 1		
Fraction 2		
Fraction 3		
Fraction 4		
Fraction 5		
Fraction 6		
Fraction 7		
Fraction 8		
Fraction 9		
Fraction 10		
Fraction 11		
Fraction 12		
Fraction 13		
Fraction 14		
Fraction 15		
Fraction 16		
Fraction 17		
Fraction 18		
Fraction 19		
Fraction 20		
Fraction 31		
Fraction 32		
		•

Fraction 33 Fraction 34 Fraction 35 Fraction 41 Fraction 42 Fraction 43	
Fraction 35 Fraction 41 Fraction 42	
Fraction 41 Fraction 42	
Fraction 42	•
Fraction 43	
Fraction 44	
Fraction 45	
Fraction 46	
Fraction 47	
Fraction 48	
Fraction 49	
Fraction 50	
Fraction 51	
Fraction 52	
Fraction 53	
Fraction 54	
Fraction 55	
Fraction 56	
Fraction 57	
Fraction 58	
Fraction 59	
Fraction 60	
Fraction 71	
Fraction 72	
Fraction 73	
Fraction 74	
Fraction 75	
Experiment IB_1_24	Number of evidence entries for this 'Experiment'.
Experiment IB_30_6	Number of evidence entries for this 'Experiment'.
Experiment IB2_1_24	Number of evidence entries for this 'Experiment'.
Experiment IB2_30_6	Number of evidence entries for this 'Experiment'.
Experiment KR_1_24	Number of evidence entries for this 'Experiment'.
Experiment KR_30_6	Number of evidence entries for this 'Experiment'.
Retention time	Retention time in minutes averaged over the evidence entries belonging to this modification-specific peptide.
Calibrated retention time	Calibrated retention time averaged over the evidence entries belonging to this modification-specific peptide. Obviously this only makes sense if retention time recalibration has been performed which is the case when matching between run is selected.
Charges	All charge states that have been observed.
PEP	Posterior Error Probability of the identification. This value essentially operates as a p-value, where smaller is more significant.
MS/MS scan number	The RAW-file derived scan number of the MS/MS with the highest peptide identification score (the highest score is stored in the column 'Score').
Raw file	The name of the RAW-file the mass spectral data was derived from.
Score	Andromeda score for the best identified among the associated MS/MS spectra.
Delta score	Score difference to the second best identified peptide.
Reverse	When marked with '+', this particular peptide was found to be part of a protein derived from the reversed part of the decoy database. These should be removed for further data analysis.
Potential contaminant	When marked with '+', this particular peptide was found to be part of a commonly occurring contaminant. These should be removed for further data analysis.
Ratio M/L	The ratio between two medium and light label partners.
Ratio M/L normalized	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L variability [%]	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.

Ratio H/L	The ratio between two heavy and light label partners.
Ratio H/L normalized	Normalized ratio between two medium and light label partners.
Ratio H/L variability [%]	The median of ratio sub-populations was shifted to 1.  Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type	quantitation that are quantities with the re-quantity method.
Ratio H/M	The ratio between two heavy and medium label partners.
Ratio H/M normalized	Normalized ratio between two heavy and medium label
	partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M variability [%]	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type	
Ratio M/L IB_1_24	The ratio between two medium and light label partners.
Ratio M/L normalized IB_1_24	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L variability [%] IB_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB_1_24	
Ratio H/L IB_1_24	The ratio between two heavy and light label partners.
Ratio H/L normalized IB_1_24	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L variability [%] IB_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB_1_24	
Ratio H/M IB 1 24	The ratio between two heavy and medium label partners.
Ratio H/M normalized IB_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M variability [%] IB_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB_1_24	
Ratio M/L IB_30_6	The ratio between two medium and light label partners.
Ratio M/L normalized IB_30_6	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L variability [%] IB_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB_30_6	
Ratio H/L IB_30_6	The ratio between two heavy and light label partners.
Ratio H/L normalized IB_30_6	Normalized ratio between two medium and light label partners.  The median of ratio sub-populations was shifted to 1.
Ratio H/L variability [%] IB_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB_30_6	Number of redundant peptides (MS1 features) used for

Ratio H/L iso-count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB_30_6	
Ratio H/M IB_30_6	The ratio between two heavy and medium label partners.
Ratio H/M normalized IB_30_6	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M variability [%] IB_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB_30_6	
Ratio M/L IB2_1_24	The ratio between two medium and light label partners.
Ratio M/L normalized IB2_1_24	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L variability [%] IB2_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB2_1_24	
Ratio H/L IB2_1_24	The ratio between two heavy and light label partners.
Ratio H/L normalized IB2_1_24	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L variability [%] IB2_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB2_1_24	
Ratio H/M IB2_1_24	The ratio between two heavy and medium label partners.
Ratio H/M normalized IB2_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M variability [%] IB2_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB2_1_24	
Ratio M/L IB2_30_6	The ratio between two medium and light label partners.
Ratio M/L normalized IB2_30_6	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L variability [%] IB2_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB2_30_6	
Ratio H/L IB2_30_6	The ratio between two heavy and light label partners.
Ratio H/L normalized IB2_30_6	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L variability [%] IB2_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB2_30_6	
Ratio H/M IB2_30_6	The ratio between two heavy and medium label partners.
Ratio H/M normalized IB2_30_6	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M variability [%] IB2_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.

Ratio H/M count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB2_30_6	
Ratio M/L KR_1_24	The ratio between two medium and light label partners.
Ratio M/L normalized KR_1_24	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L variability [%] KR_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type KR_1_24	
Ratio H/L KR_1_24	The ratio between two heavy and light label partners.
Ratio H/L normalized KR_1_24	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L variability [%] KR_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type KR_1_24	
Ratio H/M KR_1_24	The ratio between two heavy and medium label partners.
Ratio H/M normalized KR_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M variability [%] KR_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type KR_1_24	
Ratio M/L KR_30_6	The ratio between two medium and light label partners.
Ratio M/L normalized KR_30_6	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L variability [%] KR_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type KR_30_6	
Ratio H/L KR_30_6	The ratio between two heavy and light label partners.
Ratio H/L normalized KR_30_6	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L variability [%] KR_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type KR_30_6	
Ratio H/M KR_30_6	The ratio between two heavy and medium label partners.
Ratio H/M normalized KR_30_6	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M variability [%] KR_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type KR_30_6	
Intensity	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.

Intensity L	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Intensity IB_1_24	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Intensity IB_30_6	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Intensity IB2_1_24	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB2_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB2_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB2_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Intensity IB2_30_6	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB2_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB2_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB2_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Intensity KR_1_24	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L KR_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M KR_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H KR_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Intensity KR_30_6	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L KR_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M KR_30_6	Summed up extracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H KR_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
id	A unique (consecutive) identifier for each row in the peptides table, which is used to cross-link the information in this table with the information stored in the other tables.
Protein group IDs	The identifiers of the protein groups this peptide was linked to, referenced against the proteinGroups table.
Peptide ID	Identifier of the associated peptide sequence summary, which can be found in the file 'peptides.txt'.
Evidence IDs	Identifier(s) for analyzed peptide evidence associated with the protein group referenced against the evidence table.
MS/MS IDs	The identifiers of the MS/MS scans identifying this peptide, referenced against the msms table.
Best MS/MS	The identifier of the best (in terms of quality) MS/MS scan identifying this peptide, referenced against the msms table.

Deamidation (NQ) site IDs	Identifier(s) for site(s) associated with this peptide, which show(s) evidence of the modification, referenced against the appropriate modification site file.
Oxidation (M) site IDs	Identifier(s) for site(s) associated with this peptide, which show(s) evidence of the modification, referenced against the appropriate modification site file.
Phospho (STY) site IDs	Identifier(s) for site(s) associated with this peptide, which show(s) evidence of the modification, referenced against the appropriate modification site file.
MS/MS Count	

# Deamidation (NQ)Sites

Name	Separator	Description
Proteins		Identifiers of proteins this site is associated with.
Positions within proteins		For each protein identifier in the 'Proteins' column you find here the position of the site in the respective protein sequence. The index of the first amino acid in the sequence is 1.
Leading proteins		
Protein		Identifier of the protein this peptide is associated with.
Fasta headers		Descriptions of proteins this peptide is associated with.
Localization prob		
Score diff		
PEP		The posterior error probability (PEP) of the best identified modified peptide containing this site.
Score		The Andromeda score of the best identified modified peptide containing this site.
Delta score		The Andromeda delta score of the best identified modified peptide containing this site.
Score for localization		The Andromeda score of the MS/MS spectrum used for calculating the localization score for this site.
Localization prob IB_1_24		
Score diff IB_1_24		
PEP IB_1_24		
Score IB_1_24		
Localization prob IB_30_6		
Score diff IB_30_6		
PEP IB_30_6		
Score IB_30_6		
Localization prob IB2_1_24		
Score diff IB2_1_24		
PEP IB2_1_24		
Score IB2_1_24		
Localization prob IB2_30_6		
Score diff IB2_30_6		
PEP IB2_30_6		
Score IB2_30_6		
Localization prob KR_1_24		
Score diff KR_1_24		
PEP KR_1_24		
Score KR_1_24		
Localization prob KR_30_6		
Score diff KR 30 6		
PEP KR_30_6		
Score KR_30_6		
Diagnostic peak		
Number of Deamidation (NQ)		Different numbers of Deamidation (NQ) on peptides that this site is involved in.
Amino acid		
Sequence window		
Modification window		
Peptide window coverage		
Deamidation (NQ) Probabilities		
Deamidation (NQ) Score diffs		
Position in peptide		
Charge		Charge state of the precursor ion.
Mass error [ppm]		Mass error of the recalibrated mass-over-charge value of the precursor ion in comparison to the predicted monoisotopic mass of the identified peptide sequence.
Identification type IB_1_24		Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Identification type IB_30_6		Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Identification type IB2_1_24		Indicates whether this experiment was identified by MS/MS or only by matching between runs.

Identification type IB2_30_6	Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Identification type KR_1_24	Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Identification type KR_30_6	Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Ratio M/L	The ratio between two medium and light label partners.
Ratio M/L1	The ratio between two medium and light label partners.
Ratio M/L 2	The ratio between two medium and light label partners.
Ratio M/L 3	The ratio between two medium and light label partners.
Ratio M/L normalized	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized1	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized2	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized3	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep.	
Ratio M/L localized	
Ratio M/L variability [%]	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type	
Ratio H/L	The ratio between two heavy and light label partners.
Ratio H/L1	The ratio between two heavy and light label partners.
Ratio H/L2	The ratio between two heavy and light label partners.
Ratio H/L 3	The ratio between two heavy and light label partners.
Ratio H/L normalized	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized1	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized2	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized3	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep.	
Ratio H/L localized	
Ratio H/L variability [%]	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L type	
Ratio H/L count	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M	The ratio between two heavy and medium label partners.
Ratio H/M1	The ratio between two heavy and medium label partners.
Ratio H/M2	The ratio between two heavy and medium label partners.
Ratio H/M3	The ratio between two heavy and medium label partners.
Ratio H/M normalized	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized1	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized2	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized3	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep.	
Ratio H/M localized	
Ratio H/M variability [%]	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type	

Occupancy M	
Occupancy M Occupancy H	
Ratio M/L IB 1 24	The ratio between two medium and light label partners.
Ratio M/L IB 1 24 1	The ratio between two medium and light label partners.
Ratio M/L IB_1_242	The ratio between two medium and light label partners.
Ratio M/L IB_1_243	The ratio between two medium and light label partners.
Ratio M/L normalized IB_1_24	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB_1_241	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB_1_242	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB_1_243	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep. IB_1_24	
Ratio M/L localized IB_1_24	
Ratio M/L variability [%] IB_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB_1_24	
Ratio H/L IB_1_24	The ratio between two heavy and light label partners.
Ratio H/L IB_1_241	The ratio between two heavy and light label partners.
Ratio H/L IB_1_242	The ratio between two heavy and light label partners.
Ratio H/L IB_1_243	The ratio between two heavy and light label partners.
Ratio H/L normalized IB_1_24  Ratio H/L normalized IB_1_241	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.  Normalized ratio between two medium and light label partners.
	The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB_1_242	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.  Normalized ratio between two medium and light label partners.
Ratio H/L normalized IB_1_243	The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep. IB_1_24 Ratio H/L localized IB 1 24	
Ratio H/L variability [%] IB_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB_1_24	
Ratio H/M IB_1_24	The ratio between two heavy and medium label partners.
Ratio H/M IB_1_241	The ratio between two heavy and medium label partners.
Ratio H/M IB_1_242	The ratio between two heavy and medium label partners.
Ratio H/M IB_1_243	The ratio between two heavy and medium label partners.
Ratio H/M normalized IB_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_1_241	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_1_242	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_1_243	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep. IB_1_24 Ratio H/M localized IB 1 24	
Ratio H/M variability [%] IB_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the
Ratio H/M count IB_1_24	natural logarithm of ratios times 100.  Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB_1_24	
Occupancy L IB_1_24	
Occupancy M IB_1_24	
Occupancy H IB_1_24	
Ratio M/L IB_30_6	The ratio between two medium and light label partners.

Ratio M/L IB 30 6 1	The ratio between two medium and light label partners.
Ratio M/L IB 30 6 2	The ratio between two medium and light label partners.
Ratio M/L IB 30 6 3	The ratio between two medium and light label partners.
Ratio M/L normalized IB_30_6	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB_30_61	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB_30_62	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB_30_63	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep. IB_30_6	The inicial of falle out populatione was stilled to 1.
Ratio M/L localized IB 30 6	
Ratio M/L variability [%] IB_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB_30_6	
Ratio H/L IB_30_6	The ratio between two heavy and light label partners.
Ratio H/L IB_30_61	The ratio between two heavy and light label partners.
Ratio H/L IB_30_62	The ratio between two heavy and light label partners.
Ratio H/L IB_30_63	The ratio between two heavy and light label partners.
Ratio H/L normalized IB_30_6	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB_30_61	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB_30_62	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB_30_63	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep. IB_30_6	
Ratio H/L localized IB_30_6	
Ratio H/L variability [%] IB_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB_30_6	
Ratio H/M IB_30_6	The ratio between two heavy and medium label partners.
Ratio H/M IB_30_61	The ratio between two heavy and medium label partners.
Ratio H/M IB_30_62	The ratio between two heavy and medium label partners.
Ratio H/M IB_30_63	The ratio between two heavy and medium label partners.
Ratio H/M normalized IB_30_6	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_30_61	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_30_62	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_30_63	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep. IB_30_6	
Ratio H/M localized IB_30_6	
Ratio H/M variability [%] IB_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB_30_6	
Occupancy L IB_30_6	
Occupancy M IB_30_6	
Occupancy H IB_30_6	
Ratio M/L IB2_1_24	The ratio between two medium and light label partners.
Ratio M/L IB2_1_241	The ratio between two medium and light label partners.
Ratio M/L IB2_1_242	The ratio between two medium and light label partners.
Ratio M/L IB2_1_243	The ratio between two medium and light label partners.

Ratio M/L normalized IB2_1_24	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB2_1_241	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB2_1_242	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB2_1_243	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep. IB2_1_24	
Ratio M/L localized IB2_1_24	
Ratio M/L variability [%] IB2_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB2_1_24	
Ratio H/L IB2_1_24	The ratio between two heavy and light label partners.
Ratio H/L IB2 1 24 1	The ratio between two heavy and light label partners.
Ratio H/L IB2 1 24 2	The ratio between two heavy and light label partners.
Ratio H/L IB2 1 24 3	The ratio between two heavy and light label partners.
	Normalized ratio between two medium and light label partners.
Ratio H/L normalized IB2_1_24	The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB2_1_241	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB2_1_242	Normalized ratio between two medium and light label partners.  The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB2_1_243	Normalized ratio between two medium and light label partners.  The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep. IB2_1_24	
Ratio H/L localized IB2_1_24	
Ratio H/L variability [%] IB2_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB2_1_24	
Ratio H/M IB2_1_24	The ratio between two heavy and medium label partners.
Ratio H/M IB2_1_241	The ratio between two heavy and medium label partners.
Ratio H/M IB2_1_242	The ratio between two heavy and medium label partners.
Ratio H/M IB2 1 24 3	The ratio between two heavy and medium label partners.
	, , ,
Ratio H/M normalized IB2_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB2_1_241	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB2_1_242	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB2_1_243  Retio H/M normal per IB2_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep. IB2_1_24	
Ratio H/M localized IB2_1_24	
Ratio H/M variability [%] IB2_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB2_1_24	
Occupancy L IB2_1_24	
Occupancy M IB2_1_24	
Occupancy H IB2_1_24	The well-between the Property of the Control of the
Ratio M/L IB2_30_6	The ratio between two medium and light label partners.
Ratio M/L IB2_30_61	The ratio between two medium and light label partners.
Ratio M/L IB2_30_62	The ratio between two medium and light label partners.
Ratio M/L IB2_30_63	The ratio between two medium and light label partners.
Ratio M/L normalized IB2_30_6	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.

Ratio M/L normalized IB2_30_61	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB2_30_62	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB2 30 6 3	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep. IB2_30_6	The median of raile say paperations had office to the
Ratio M/L localized IB2_30_6	
Ratio M/L variability [%] IB2_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB2_30_6	
Ratio H/L IB2_30_6	The ratio between two heavy and light label partners.
Ratio H/L IB2_30_61	The ratio between two heavy and light label partners.
Ratio H/L IB2_30_62	The ratio between two heavy and light label partners.
Ratio H/L IB2_30_63	The ratio between two heavy and light label partners.
Ratio H/L normalized IB2_30_6	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB2_30_61	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB2_30_62	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB2_30_63	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep. IB2_30_6	
Ratio H/L localized IB2_30_6	
Ratio H/L variability [%] IB2_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB2_30_6	Number of redundant peptides (MS1 features) used for guantitation.
Ratio H/L iso-count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB2_30_6	
Ratio H/M IB2_30_6	The ratio between two heavy and medium label partners.
Ratio H/M IB2_30_61	The ratio between two heavy and medium label partners.
Ratio H/M IB2_30_62	The ratio between two heavy and medium label partners.
Ratio H/M IB2_30_63	The ratio between two heavy and medium label partners.
Ratio H/M normalized IB2_30_6	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB2_30_61	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB2_30_62	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB2_30_63	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep. IB2_30_6	
Ratio H/M localized IB2_30_6	
Ratio H/M variability [%] IB2_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB2_30_6	
Occupancy L IB2_30_6	
Occupancy M IB2_30_6	
Occupancy H IB2_30_6	
Ratio M/L KR_1_24	The ratio between two medium and light label partners.
Ratio M/L KR_1_241	The ratio between two medium and light label partners.
Ratio M/L KR_1_242	The ratio between two medium and light label partners.
Ratio M/L KR_1_243	The ratio between two medium and light label partners.
Ratio M/L normalized KR_1_24	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized KR_1_241	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.

Ratio M/L normalized KR_1_242	Normalized ratio between two heavy and light label partners.
Ratio IVI/E Hormalized RR_1_242	The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized KR_1_243	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep. KR_1_24	
Ratio M/L localized KR_1_24	
Ratio M/L variability [%] KR_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type KR_1_24	
Ratio H/L KR_1_24	The ratio between two heavy and light label partners.
Ratio H/L KR_1_241	The ratio between two heavy and light label partners.
Ratio H/L KR_1_242	The ratio between two heavy and light label partners.
Ratio H/L KR_1_243	The ratio between two heavy and light label partners.
Ratio H/L normalized KR_1_24	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_1_241	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_1_242	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_1_243	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep. KR_1_24	
Ratio H/L localized KR_1_24	
Ratio H/L variability [%] KR_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type KR_1_24	
Ratio H/M KR_1_24	The ratio between two heavy and medium label partners.
Ratio H/M KR_1_241	The ratio between two heavy and medium label partners.
Ratio H/M KR_1_242	The ratio between two heavy and medium label partners.
Ratio H/M KR_1_243	The ratio between two heavy and medium label partners.
Ratio H/M normalized KR_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_1_241	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_1_242	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_1_243	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep. KR_1_24	
Ratio H/M localized KR_1_24	0 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
Ratio H/M variability [%] KR_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type KR_1_24	
Occupancy L KR_1_24	
Occupancy M KR_1_24	
Occupancy H KR_1_24	
Ratio M/L KR_30_6	The ratio between two medium and light label partners.
Ratio M/L KR_30_61	The ratio between two medium and light label partners.
Ratio M/L KR_30_62	The ratio between two medium and light label partners.
Ratio M/L KR_30_63	The ratio between two medium and light label partners.
Ratio M/L normalized KR_30_6	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized KR_30_61	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized KR_30_62	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.

Ratio M/L normalized KR_30_63	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep. KR_30_6	The median of ratio out populations trac office to 1.
Ratio M/L localized KR_30_6	
Ratio M/L variability [%] KR_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the
Ratio M/L count KR_30_6	natural logarithm of ratios times 100.  Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type KR_30_6	
Ratio H/L KR_30_6	The ratio between two heavy and light label partners.
Ratio H/L KR_30_61	The ratio between two heavy and light label partners.
Ratio H/L KR_30_62	The ratio between two heavy and light label partners.
Ratio H/L KR_30_63	The ratio between two heavy and light label partners.
Ratio H/L normalized KR_30_6	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_30_61	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_30_62	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_30_63	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep. KR_30_6	
Ratio H/L localized KR_30_6	
Ratio H/L variability [%] KR_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type KR_30_6	
Ratio H/M KR_30_6	The ratio between two heavy and medium label partners.
Ratio H/M KR_30_61	The ratio between two heavy and medium label partners.
Ratio H/M KR_30_62	The ratio between two heavy and medium label partners.
Ratio H/M KR_30_63	The ratio between two heavy and medium label partners.
Ratio H/M normalized KR_30_6	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_30_61	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_30_62	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_30_63	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep. KR_30_6	
Ratio H/M localized KR_30_6	
Ratio H/M variability [%] KR_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type KR_30_6	
Occupancy L KR_30_6	
Occupancy M KR_30_6	
Occupancy H KR_30_6	
Intensity	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L	
Ratio mod/base M	
Ratio mod/base H	

Intensity IB_1_24	Summed up extracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a label of a label of the identified AB sequence in case of a label of the identified AB sequence.
	labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L IB_1_24	
Ratio mod/base M IB_1_24	
Ratio mod/base H IB_1_24	
Intensity IB_30_6	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L IB_30_6	
Ratio mod/base M IB_30_6	
Ratio mod/base H IB_30_6	
Intensity IB2_1_24	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB2_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB2_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB2_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L IB2_1_24	
Ratio mod/base M IB2_1_24	
Ratio mod/base H IB2_1_24	
Intensity IB2_30_6	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB2_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB2_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB2_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L IB2_30_6	
Ratio mod/base M IB2_30_6	
Ratio mod/base H IB2_30_6	
Intensity KR_1_24	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L KR_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M KR_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H KR_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L KR_1_24	
Ratio mod/base M KR_1_24	
Ratio mod/base H KR_1_24	
Intensity KR_30_6	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L KR_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M KR_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H KR_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L KR_30_6	

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Ratio mod/base H KR_30_6	
Reverse	When marked with '+', this particular peptide was found to be part of a protein derived from the reversed part of the protein sequence database. These should be removed for further data analysis.
otential contaminant	When marked with '+', this particular peptide was found to be part of a commonly occurring contaminant. These should be removed for further data analysis.
1	A unique (consecutive) identifier for each row in the site table, which is used to cross-link the information in this file with the information stored in the other files.
Protein group IDs	The identifier of the protein-group this peptide sequence is associated with, which can be used to look up the extended protein information in the file 'proteinGroups.txt'.  As a single peptide can be linked to multiple proteins (e.g. in the case of razor-proteins), multiple id's can be stored here separated by a semicolon.  As a protein can be identified by multiple peptides, the same id can be found in different rows.
Positions	The positions of the modifications in the protein amino acid sequence.
Position	The position of the modification in the protein amino acid sequence.
Peptide IDs	Identifier(s) of the associated peptide sequence(s) summary, which can be found in the file 'peptides.txt'.
Mod. peptide IDs	Identifier(s) of the associated peptide sequence(s) summary, which can be found in the file 'modificationSpecificPeptides.txt'.
vidence IDs	Identifier(s) for analyzed peptide evidence associated with the protein group referenced against the evidence table.
MS/MS IDs	The identifiers of the MS/MS scans identifying this peptide, referenced against the msms table.
Sest localization evidence ID	
Best localization MS/MS ID	
Sest localization raw file	
Best localization scan number	
Best score evidence ID	
Sest score MS/MS ID	
sest score raw file	
Best score scan number	
Best PEP evidence ID	
Best PEP MS/MS ID	
Best PEP raw file	
Sest PEP scan number	

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# Oxidation (M)Sites

Name	Separator	Description
Proteins		Identifiers of proteins this site is associated with.
Positions within proteins		For each protein identifier in the 'Proteins' column you find here the position of the site in the respective protein sequence. The index of the first amino acid in the sequence is 1.
Leading proteins		
Protein		Identifier of the protein this peptide is associated with.
Fasta headers		Descriptions of proteins this peptide is associated with.
Localization prob		
Score diff		
PEP		The posterior error probability (PEP) of the best identified modified peptide containing this site.
Score		The Andromeda score of the best identified modified peptide containing this site.
Delta score		The Andromeda delta score of the best identified modified peptide containing this site.
Score for localization		The Andromeda score of the MS/MS spectrum used for calculating the localization score for this site.
Localization prob IB_1_24		
Score diff IB_1_24		
PEP IB_1_24		
Score IB_1_24		
Localization prob IB_30_6		
Score diff IB_30_6		
PEP IB_30_6		
Score IB_30_6		
Localization prob IB2_1_24		
Score diff IB2 1 24		
PEP IB2 1 24		
Score IB2 1 24		
Localization prob IB2_30_6		
Score diff IB2_30_6		
PEP IB2 30 6		
Score IB2 30 6		
Localization prob KR_1_24		
Score diff KR_1_24		
PEP KR 1 24		
Score KR_1_24		
Localization prob KR_30_6		
Score diff KR 30 6		
PEP KR 30 6		
Score KR_30_6		
Diagnostic peak		
Number of Oxidation (M)		Different numbers of Oxidation (M) on peptides that this site is involved in.
Amino acid		
Sequence window		
Modification window		
Peptide window coverage		
Oxidation (M) Probabilities		
Oxidation (M) Score diffs		
Position in peptide		
Charge		Charge state of the precursor ion.
Mass error [ppm]		Mass error of the recalibrated mass-over-charge value of the precursor ion in comparison to the predicted monoisotopic mass of the identified peptide sequence.
Identification type IB_1_24		Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Identification type IB_30_6		Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Identification type IB2_1_24		Indicates whether this experiment was identified by MS/MS or only by matching between runs.

Identification type IB2_30_6	Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Identification type KR_1_24	Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Identification type KR_30_6	Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Ratio M/L	The ratio between two medium and light label partners.
Ratio M/L1	The ratio between two medium and light label partners.
Ratio M/L 2	The ratio between two medium and light label partners.
Ratio M/L 3	The ratio between two medium and light label partners.
Ratio M/L normalized	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized1	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized2	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized3	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep.	
Ratio M/L localized	
Ratio M/L variability [%]	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type	
Ratio H/L	The ratio between two heavy and light label partners.
Ratio H/L1	The ratio between two heavy and light label partners.
Ratio H/L2	The ratio between two heavy and light label partners.
Ratio H/L 3	The ratio between two heavy and light label partners.
Ratio H/L normalized	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized1	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized2	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized3	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep.	
Ratio H/L localized	
Ratio H/L variability [%]	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L type	
Ratio H/L count	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M	The ratio between two heavy and medium label partners.
Ratio H/M1	The ratio between two heavy and medium label partners.
Ratio H/M2	The ratio between two heavy and medium label partners.
Ratio H/M3	The ratio between two heavy and medium label partners.
Ratio H/M normalized	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized1	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized2	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized3	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep.	
Ratio H/M localized	
Ratio H/M variability [%]	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type	

Occupancy M	
Occupancy M Occupancy H	
Ratio M/L IB 1 24	The ratio between two medium and light label partners.
Ratio M/L IB 1 24 1	The ratio between two medium and light label partners.
Ratio M/L IB_1_242	The ratio between two medium and light label partners.
Ratio M/L IB_1_243	The ratio between two medium and light label partners.
Ratio M/L normalized IB_1_24	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB_1_241	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB_1_242	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB_1_243	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep. IB_1_24	
Ratio M/L localized IB_1_24	
Ratio M/L variability [%] IB_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB_1_24	
Ratio H/L IB_1_24	The ratio between two heavy and light label partners.
Ratio H/L IB_1_241	The ratio between two heavy and light label partners.
Ratio H/L IB_1_242	The ratio between two heavy and light label partners.
Ratio H/L IB_1_243	The ratio between two heavy and light label partners.
Ratio H/L normalized IB_1_24  Ratio H/L normalized IB_1_241	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.  Normalized ratio between two medium and light label partners.
	The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB_1_242	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.  Normalized ratio between two medium and light label partners.
Ratio H/L normalized IB_1_243	The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep. IB_1_24 Ratio H/L localized IB 1 24	
Ratio H/L variability [%] IB_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB_1_24	
Ratio H/M IB_1_24	The ratio between two heavy and medium label partners.
Ratio H/M IB_1_241	The ratio between two heavy and medium label partners.
Ratio H/M IB_1_242	The ratio between two heavy and medium label partners.
Ratio H/M IB_1_243	The ratio between two heavy and medium label partners.
Ratio H/M normalized IB_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_1_241	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_1_242	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_1_243	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep. IB_1_24 Ratio H/M localized IB 1 24	
Ratio H/M variability [%] IB_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the
Ratio H/M count IB_1_24	natural logarithm of ratios times 100.  Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB_1_24	
Occupancy L IB_1_24	
Occupancy M IB_1_24	
Occupancy H IB_1_24	
Ratio M/L IB_30_6	The ratio between two medium and light label partners.

Ratio M/L IB 30 6 1	The ratio between two medium and light label partners.
Ratio M/L IB 30 6 2	The ratio between two medium and light label partners.
Ratio M/L IB 30 6 3	The ratio between two medium and light label partners.
Ratio M/L normalized IB_30_6	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB_30_61	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB_30_62	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB_30_63	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep. IB_30_6	The inicial of falle out populatione was stilled to 1.
Ratio M/L localized IB 30 6	
Ratio M/L variability [%] IB_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB_30_6	
Ratio H/L IB_30_6	The ratio between two heavy and light label partners.
Ratio H/L IB_30_61	The ratio between two heavy and light label partners.
Ratio H/L IB_30_62	The ratio between two heavy and light label partners.
Ratio H/L IB_30_63	The ratio between two heavy and light label partners.
Ratio H/L normalized IB_30_6	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB_30_61	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB_30_62	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB_30_63	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep. IB_30_6	
Ratio H/L localized IB_30_6	
Ratio H/L variability [%] IB_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB_30_6	
Ratio H/M IB_30_6	The ratio between two heavy and medium label partners.
Ratio H/M IB_30_61	The ratio between two heavy and medium label partners.
Ratio H/M IB_30_62	The ratio between two heavy and medium label partners.
Ratio H/M IB_30_63	The ratio between two heavy and medium label partners.
Ratio H/M normalized IB_30_6	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_30_61	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_30_62	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_30_63	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep. IB_30_6	
Ratio H/M localized IB_30_6	
Ratio H/M variability [%] IB_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB_30_6	
Occupancy L IB_30_6	
Occupancy M IB_30_6	
Occupancy H IB_30_6	
Ratio M/L IB2_1_24	The ratio between two medium and light label partners.
Ratio M/L IB2_1_241	The ratio between two medium and light label partners.
Ratio M/L IB2_1_242	The ratio between two medium and light label partners.
Ratio M/L IB2_1_243	The ratio between two medium and light label partners.

Ratio M/L normalized IB2_1_24	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB2_1_241	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB2_1_242	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB2_1_243	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep. IB2_1_24	
Ratio M/L localized IB2_1_24	
Ratio M/L variability [%] IB2_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB2_1_24	
Ratio H/L IB2_1_24	The ratio between two heavy and light label partners.
Ratio H/L IB2 1 24 1	The ratio between two heavy and light label partners.
Ratio H/L IB2 1 24 2	The ratio between two heavy and light label partners.
Ratio H/L IB2 1 24 3	The ratio between two heavy and light label partners.
	Normalized ratio between two medium and light label partners.
Ratio H/L normalized IB2_1_24	The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB2_1_241	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB2_1_242	Normalized ratio between two medium and light label partners.  The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB2_1_243	Normalized ratio between two medium and light label partners.  The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep. IB2_1_24	
Ratio H/L localized IB2_1_24	
Ratio H/L variability [%] IB2_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB2_1_24	
Ratio H/M IB2_1_24	The ratio between two heavy and medium label partners.
Ratio H/M IB2_1_241	The ratio between two heavy and medium label partners.
Ratio H/M IB2_1_242	The ratio between two heavy and medium label partners.
Ratio H/M IB2 1 24 3	The ratio between two heavy and medium label partners.
	, , ,
Ratio H/M normalized IB2_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB2_1_241	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB2_1_242	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB2_1_243  Retio H/M normal per IB2_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep. IB2_1_24	
Ratio H/M localized IB2_1_24	
Ratio H/M variability [%] IB2_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB2_1_24	
Occupancy L IB2_1_24	
Occupancy M IB2_1_24	
Occupancy H IB2_1_24	The well-between the Property of the Control of the
Ratio M/L IB2_30_6	The ratio between two medium and light label partners.
Ratio M/L IB2_30_61	The ratio between two medium and light label partners.
Ratio M/L IB2_30_62	The ratio between two medium and light label partners.
Ratio M/L IB2_30_63	The ratio between two medium and light label partners.
Ratio M/L normalized IB2_30_6	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.

Normalized ratio between two heavy and light label partners.
The median of ratio sub-populations was shifted to 1.
Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Number of redundant peptides (MS1 features) used for quantitation.
Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
The ratio between two heavy and light label partners.
The ratio between two heavy and light label partners.
The ratio between two heavy and light label partners.
The ratio between two heavy and light label partners.
Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Number of redundant peptides (MS1 features) used for quantitation.
Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
The ratio between two heavy and medium label partners.
The ratio between two heavy and medium label partners.
The ratio between two heavy and medium label partners.
The ratio between two heavy and medium label partners.
Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Number of redundant peptides (MS1 features) used for quantitation.
Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
The ratio between two medium and light label partners.
The ratio between two medium and light label partners.
The ratio between two medium and light label partners.
The ratio between two medium and light label partners.
Normalized ratio between two heavy and light label partners.  The median of ratio sub-populations was shifted to 1.
Normalized ratio between two heavy and light label partners.

Ratio M/L normalized KR_1_242	Normalized ratio between two heavy and light label partners.
Ratio IVI/E Hormalized RR_1_242	The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized KR_1_243	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep. KR_1_24	
Ratio M/L localized KR_1_24	
Ratio M/L variability [%] KR_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type KR_1_24	
Ratio H/L KR_1_24	The ratio between two heavy and light label partners.
Ratio H/L KR_1_241	The ratio between two heavy and light label partners.
Ratio H/L KR_1_242	The ratio between two heavy and light label partners.
Ratio H/L KR_1_243	The ratio between two heavy and light label partners.
Ratio H/L normalized KR_1_24	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_1_241	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_1_242	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_1_243	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep. KR_1_24	
Ratio H/L localized KR_1_24	
Ratio H/L variability [%] KR_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type KR_1_24	
Ratio H/M KR_1_24	The ratio between two heavy and medium label partners.
Ratio H/M KR_1_241	The ratio between two heavy and medium label partners.
Ratio H/M KR_1_242	The ratio between two heavy and medium label partners.
Ratio H/M KR_1_243	The ratio between two heavy and medium label partners.
Ratio H/M normalized KR_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_1_241	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_1_242	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_1_243	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep. KR_1_24	
Ratio H/M localized KR_1_24	0 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
Ratio H/M variability [%] KR_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type KR_1_24	
Occupancy L KR_1_24	
Occupancy M KR_1_24	
Occupancy H KR_1_24	
Ratio M/L KR_30_6	The ratio between two medium and light label partners.
Ratio M/L KR_30_61	The ratio between two medium and light label partners.
Ratio M/L KR_30_62	The ratio between two medium and light label partners.
Ratio M/L KR_30_63	The ratio between two medium and light label partners.
Ratio M/L normalized KR_30_6	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized KR_30_61	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized KR_30_62	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.

Ratio M/L normalized KR_30_63	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep. KR_30_6	The median of falle out populations trac office to 1.
Ratio M/L localized KR_30_6	
Ratio M/L variability [%] KR_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the
Ratio M/L count KR_30_6	natural logarithm of ratios times 100.  Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type KR_30_6	
Ratio H/L KR_30_6	The ratio between two heavy and light label partners.
Ratio H/L KR_30_61	The ratio between two heavy and light label partners.
Ratio H/L KR_30_62	The ratio between two heavy and light label partners.
Ratio H/L KR_30_63	The ratio between two heavy and light label partners.
Ratio H/L normalized KR_30_6	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_30_61	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_30_62	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_30_63	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep. KR_30_6	
Ratio H/L localized KR_30_6	
Ratio H/L variability [%] KR_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type KR_30_6	
Ratio H/M KR_30_6	The ratio between two heavy and medium label partners.
Ratio H/M KR_30_61	The ratio between two heavy and medium label partners.
Ratio H/M KR_30_62	The ratio between two heavy and medium label partners.
Ratio H/M KR_30_63	The ratio between two heavy and medium label partners.
Ratio H/M normalized KR_30_6	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_30_61	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_30_62	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_30_63	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep. KR_30_6	
Ratio H/M localized KR_30_6	
Ratio H/M variability [%] KR_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type KR_30_6	
Occupancy L KR_30_6	
Occupancy M KR_30_6	
Occupancy H KR_30_6	
Intensity	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L	
Ratio mod/base M	
Ratio mod/base H	

Intensity IB_1_24	Summed up extracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a label of a label of the identified AB sequence in case of a label of the identified AB sequence.
	labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L IB_1_24	
Ratio mod/base M IB_1_24	
Ratio mod/base H IB_1_24	
Intensity IB_30_6	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L IB_30_6	
Ratio mod/base M IB_30_6	
Ratio mod/base H IB_30_6	
Intensity IB2_1_24	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB2_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB2_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB2_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L IB2_1_24	
Ratio mod/base M IB2_1_24	
Ratio mod/base H IB2_1_24	
Intensity IB2_30_6	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB2_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB2_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB2_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L IB2_30_6	
Ratio mod/base M IB2_30_6	
Ratio mod/base H IB2_30_6	
Intensity KR_1_24	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L KR_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M KR_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H KR_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L KR_1_24	
Ratio mod/base M KR_1_24	
Ratio mod/base H KR_1_24	
Intensity KR_30_6	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L KR_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M KR_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H KR_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L KR_30_6	

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Ratio mod/base H KR_30_6	
Reverse	When marked with '+', this particular peptide was found to be part of a protein derived from the reversed part of the protein sequence database. These should be removed for further data analysis.
otential contaminant	When marked with '+', this particular peptide was found to be part of a commonly occurring contaminant. These should be removed for further data analysis.
1	A unique (consecutive) identifier for each row in the site table, which is used to cross-link the information in this file with the information stored in the other files.
Protein group IDs	The identifier of the protein-group this peptide sequence is associated with, which can be used to look up the extended protein information in the file 'proteinGroups.txt'.  As a single peptide can be linked to multiple proteins (e.g. in the case of razor-proteins), multiple id's can be stored here separated by a semicolon.  As a protein can be identified by multiple peptides, the same id can be found in different rows.
Positions	The positions of the modifications in the protein amino acid sequence.
Position	The position of the modification in the protein amino acid sequence.
Peptide IDs	Identifier(s) of the associated peptide sequence(s) summary, which can be found in the file 'peptides.txt'.
Mod. peptide IDs	Identifier(s) of the associated peptide sequence(s) summary, which can be found in the file 'modificationSpecificPeptides.txt'.
vidence IDs	Identifier(s) for analyzed peptide evidence associated with the protein group referenced against the evidence table.
MS/MS IDs	The identifiers of the MS/MS scans identifying this peptide, referenced against the msms table.
Sest localization evidence ID	
Best localization MS/MS ID	
Sest localization raw file	
Best localization scan number	
Best score evidence ID	
Sest score MS/MS ID	
sest score raw file	
Best score scan number	
Best PEP evidence ID	
Best PEP MS/MS ID	
Best PEP raw file	
Sest PEP scan number	

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# Phospho (STY)Sites

Name	Separator	Description
Proteins		Identifiers of proteins this site is associated with.
Positions within proteins		For each protein identifier in the 'Proteins' column you find here the position of the site in the respective protein sequence. The index of the first amino acid in the sequence is 1.
Leading proteins		
Protein		Identifier of the protein this peptide is associated with.
Fasta headers		Descriptions of proteins this peptide is associated with.
Localization prob		
Score diff		
PEP		The posterior error probability (PEP) of the best identified
		modified peptide containing this site.
Score		The Andromeda score of the best identified modified peptide containing this site.
Delta score		The Andromeda delta score of the best identified modified peptide containing this site.
Score for localization		The Andromeda score of the MS/MS spectrum used for calculating the localization score for this site.
Localization prob IB_1_24		
Score diff IB_1_24		
PEP IB_1_24		
Score IB_1_24		
Localization prob IB_30_6		
Score diff IB_30_6		
PEP IB_30_6		
Score IB_30_6		
Localization prob IB2_1_24		
Score diff IB2 1 24		
PEP IB2 1 24		
Score IB2 1 24		
Localization prob IB2_30_6		
Score diff IB2_30_6		
PEP IB2 30 6		
Score IB2 30 6		
Localization prob KR_1_24		
Score diff KR_1_24		
PEP KR 1 24		
Score KR_1_24		
Localization prob KR_30_6		
Score diff KR_30_6		
PEP KR_30_6		
Score KR_30_6		
Diagnostic peak		
Number of Phospho (STY)		Different numbers of Phospho (STY) on peptides that this site is involved in.
Amino acid		
Sequence window		
Modification window		
Peptide window coverage		
Phospho (STY) Probabilities		
Phospho (STY) Score diffs		
Position in peptide		
Charge		Charge state of the precursor ion.
Mass error [ppm]		Mass error of the recalibrated mass-over-charge value of the precursor ion in comparison to the predicted monoisotopic mass of the identified peptide sequence.
Identification type IB_1_24		Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Identification type IB_30_6		Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Identification type IB2_1_24		Indicates whether this experiment was identified by MS/MS or only by matching between runs.

Identification type IB2_30_6	Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Identification type KR_1_24	Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Identification type KR_30_6	Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Ratio M/L	The ratio between two medium and light label partners.
Ratio M/L1	The ratio between two medium and light label partners.
Ratio M/L 2	The ratio between two medium and light label partners.
Ratio M/L 3	The ratio between two medium and light label partners.
Ratio M/L normalized	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized1	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized2	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized3	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep.	
Ratio M/L localized	
Ratio M/L variability [%]	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type	
Ratio H/L	The ratio between two heavy and light label partners.
Ratio H/L1	The ratio between two heavy and light label partners.
Ratio H/L2	The ratio between two heavy and light label partners.
Ratio H/L 3	The ratio between two heavy and light label partners.
Ratio H/L normalized	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized1	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized2	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized3	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep.	
Ratio H/L localized	
Ratio H/L variability [%]	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L type	
Ratio H/L count	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M	The ratio between two heavy and medium label partners.
Ratio H/M1	The ratio between two heavy and medium label partners.
Ratio H/M2	The ratio between two heavy and medium label partners.
Ratio H/M3	The ratio between two heavy and medium label partners.
Ratio H/M normalized	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized1	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized2	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized3	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep.	
Ratio H/M localized	
Ratio H/M variability [%]	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type	

Occurred by M	
Occupancy M	
Occupancy H Ratio M/L IB 1 24	The ratio between two medium and light label partners.
Ratio M/L IB 1 24 1	The ratio between two medium and light label partners.
Ratio M/L IB 1 24 2	The ratio between two medium and light label partners.
Ratio M/L IB_1_243	The ratio between two medium and light label partners.
Ratio M/L normalized IB_1_24	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB_1_241	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB_1_242	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB_1_243	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep. IB_1_24	
Ratio M/L localized IB_1_24	
Ratio M/L variability [%] IB_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB_1_24	
Ratio H/L IB_1_24	The ratio between two heavy and light label partners.
Ratio H/L IB_1_241	The ratio between two heavy and light label partners.
Ratio H/L IB_1_242	The ratio between two heavy and light label partners.
Ratio H/L IB_1_243	The ratio between two heavy and light label partners.
Ratio H/L normalized IB_1_24	Normalized ratio between two medium and light label partners.  The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB_1_241	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB_1_242	Normalized ratio between two medium and light label partners.  The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB_1_243	Normalized ratio between two medium and light label partners.  The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep. IB_1_24 Ratio H/L localized IB 1 24	
Ratio H/L variability [%] IB_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB_1_24	
Ratio H/M IB_1_24	The ratio between two heavy and medium label partners.
Ratio H/M IB_1_241	The ratio between two heavy and medium label partners.
Ratio H/M IB_1_242	The ratio between two heavy and medium label partners.
Ratio H/M IB_1_243	The ratio between two heavy and medium label partners.
Ratio H/M normalized IB_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_1_241	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_1_242	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_1_243	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep. IB_1_24	
Ratio H/M localized IB_1_24	Confficient of contact the contact of the contact o
Ratio H/M variability [%] IB_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB_1_24	
Occupancy L IB_1_24	
Occupancy M IB_1_24	
Occupancy H IB_1_24	
Ratio M/L IB_30_6	The ratio between two medium and light label partners.

Ratio M/L IB 30 6 1	The ratio between two medium and light label partners.
Ratio M/L IB 30 6 2	The ratio between two medium and light label partners.
Ratio M/L IB 30 6 3	The ratio between two medium and light label partners.
Ratio M/L normalized IB_30_6	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB_30_61	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB_30_62	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB_30_63	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep. IB_30_6	The inicial of falle out populatione was stilled to 1.
Ratio M/L localized IB 30 6	
Ratio M/L variability [%] IB_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB_30_6	
Ratio H/L IB_30_6	The ratio between two heavy and light label partners.
Ratio H/L IB_30_61	The ratio between two heavy and light label partners.
Ratio H/L IB_30_62	The ratio between two heavy and light label partners.
Ratio H/L IB_30_63	The ratio between two heavy and light label partners.
Ratio H/L normalized IB_30_6	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB_30_61	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB_30_62	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB_30_63	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep. IB_30_6	
Ratio H/L localized IB_30_6	
Ratio H/L variability [%] IB_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB_30_6	
Ratio H/M IB_30_6	The ratio between two heavy and medium label partners.
Ratio H/M IB_30_61	The ratio between two heavy and medium label partners.
Ratio H/M IB_30_62	The ratio between two heavy and medium label partners.
Ratio H/M IB_30_63	The ratio between two heavy and medium label partners.
Ratio H/M normalized IB_30_6	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_30_61	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_30_62	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_30_63	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep. IB_30_6	
Ratio H/M localized IB_30_6	
Ratio H/M variability [%] IB_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB_30_6	
Occupancy L IB_30_6	
Occupancy M IB_30_6	
Occupancy H IB_30_6	
Ratio M/L IB2_1_24	The ratio between two medium and light label partners.
Ratio M/L IB2_1_241	The ratio between two medium and light label partners.
Ratio M/L IB2_1_242	The ratio between two medium and light label partners.
Ratio M/L IB2_1_243	The ratio between two medium and light label partners.

Ratio M/L normalized IB2_1_24	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB2_1_241	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB2_1_242	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB2_1_243	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep. IB2_1_24	
Ratio M/L localized IB2_1_24	
Ratio M/L variability [%] IB2_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB2_1_24	
Ratio H/L IB2_1_24	The ratio between two heavy and light label partners.
Ratio H/L IB2 1 24 1	The ratio between two heavy and light label partners.
Ratio H/L IB2 1 24 2	The ratio between two heavy and light label partners.
Ratio H/L IB2 1 24 3	The ratio between two heavy and light label partners.
	Normalized ratio between two medium and light label partners.
Ratio H/L normalized IB2_1_24	The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB2_1_241	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB2_1_242	Normalized ratio between two medium and light label partners.  The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB2_1_243	Normalized ratio between two medium and light label partners.  The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep. IB2_1_24	
Ratio H/L localized IB2_1_24	
Ratio H/L variability [%] IB2_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB2_1_24	
Ratio H/M IB2_1_24	The ratio between two heavy and medium label partners.
Ratio H/M IB2_1_241	The ratio between two heavy and medium label partners.
Ratio H/M IB2_1_242	The ratio between two heavy and medium label partners.
Ratio H/M IB2 1 24 3	The ratio between two heavy and medium label partners.
	, , ,
Ratio H/M normalized IB2_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB2_1_241	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB2_1_242	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB2_1_243  Retio H/M normal per IB2_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep. IB2_1_24	
Ratio H/M localized IB2_1_24	
Ratio H/M variability [%] IB2_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB2_1_24	
Occupancy L IB2_1_24	
Occupancy M IB2_1_24	
Occupancy H IB2_1_24	The well-between the Property of the Control of the
Ratio M/L IB2_30_6	The ratio between two medium and light label partners.
Ratio M/L IB2_30_61	The ratio between two medium and light label partners.
Ratio M/L IB2_30_62	The ratio between two medium and light label partners.
Ratio M/L IB2_30_63	The ratio between two medium and light label partners.
Ratio M/L normalized IB2_30_6	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.

Ratio M/L normalized IB2_30_61	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB2_30_62	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized IB2 30 6 3	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep. IB2_30_6	The median of raise say paperations had office to the
Ratio M/L localized IB2_30_6	
Ratio M/L variability [%] IB2_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB2_30_6	
Ratio H/L IB2_30_6	The ratio between two heavy and light label partners.
Ratio H/L IB2_30_61	The ratio between two heavy and light label partners.
Ratio H/L IB2_30_62	The ratio between two heavy and light label partners.
Ratio H/L IB2_30_63	The ratio between two heavy and light label partners.
Ratio H/L normalized IB2_30_6	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB2_30_61	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB2_30_62	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB2_30_63	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep. IB2_30_6	
Ratio H/L localized IB2_30_6	
Ratio H/L variability [%] IB2_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count IB2_30_6	Number of redundant peptides (MS1 features) used for guantitation.
Ratio H/L iso-count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB2_30_6	
Ratio H/M IB2_30_6	The ratio between two heavy and medium label partners.
Ratio H/M IB2_30_61	The ratio between two heavy and medium label partners.
Ratio H/M IB2_30_62	The ratio between two heavy and medium label partners.
Ratio H/M IB2_30_63	The ratio between two heavy and medium label partners.
Ratio H/M normalized IB2_30_6	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB2_30_61	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB2_30_62	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB2_30_63	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep. IB2_30_6	
Ratio H/M localized IB2_30_6	
Ratio H/M variability [%] IB2_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB2_30_6	
Occupancy L IB2_30_6	
Occupancy M IB2_30_6	
Occupancy H IB2_30_6	
Ratio M/L KR_1_24	The ratio between two medium and light label partners.
Ratio M/L KR_1_241	The ratio between two medium and light label partners.
Ratio M/L KR_1_242	The ratio between two medium and light label partners.
Ratio M/L KR_1_243	The ratio between two medium and light label partners.
Ratio M/L normalized KR_1_24	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized KR_1_241	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.

Ratio M/L normalized KR_1_242	Normalized ratio between two heavy and light label partners.
Ratio IVI/E Hormalized RR_1_242	The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized KR_1_243	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep. KR_1_24	
Ratio M/L localized KR_1_24	
Ratio M/L variability [%] KR_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type KR_1_24	
Ratio H/L KR_1_24	The ratio between two heavy and light label partners.
Ratio H/L KR_1_241	The ratio between two heavy and light label partners.
Ratio H/L KR_1_242	The ratio between two heavy and light label partners.
Ratio H/L KR_1_243	The ratio between two heavy and light label partners.
Ratio H/L normalized KR_1_24	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_1_241	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_1_242	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_1_243	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep. KR_1_24	
Ratio H/L localized KR_1_24	
Ratio H/L variability [%] KR_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type KR_1_24	
Ratio H/M KR_1_24	The ratio between two heavy and medium label partners.
Ratio H/M KR_1_241	The ratio between two heavy and medium label partners.
Ratio H/M KR_1_242	The ratio between two heavy and medium label partners.
Ratio H/M KR_1_243	The ratio between two heavy and medium label partners.
Ratio H/M normalized KR_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_1_241	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_1_242	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_1_243	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep. KR_1_24	
Ratio H/M localized KR_1_24	0 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
Ratio H/M variability [%] KR_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type KR_1_24	
Occupancy L KR_1_24	
Occupancy M KR_1_24	
Occupancy H KR_1_24	
Ratio M/L KR_30_6	The ratio between two medium and light label partners.
Ratio M/L KR_30_61	The ratio between two medium and light label partners.
Ratio M/L KR_30_62	The ratio between two medium and light label partners.
Ratio M/L KR_30_63	The ratio between two medium and light label partners.
Ratio M/L normalized KR_30_6	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized KR_30_61	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L normalized KR_30_62	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.

Ratio M/L normalized KR_30_63	Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio M/L unmod. pep. KR_30_6	The median of falle out populations trac office to 1.
Ratio M/L localized KR_30_6	
Ratio M/L variability [%] KR_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the
Ratio M/L count KR_30_6	natural logarithm of ratios times 100.  Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type KR_30_6	
Ratio H/L KR_30_6	The ratio between two heavy and light label partners.
Ratio H/L KR_30_61	The ratio between two heavy and light label partners.
Ratio H/L KR_30_62	The ratio between two heavy and light label partners.
Ratio H/L KR_30_63	The ratio between two heavy and light label partners.
Ratio H/L normalized KR_30_6	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_30_61	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_30_62	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_30_63	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L unmod. pep. KR_30_6	
Ratio H/L localized KR_30_6	
Ratio H/L variability [%] KR_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type KR_30_6	
Ratio H/M KR_30_6	The ratio between two heavy and medium label partners.
Ratio H/M KR_30_61	The ratio between two heavy and medium label partners.
Ratio H/M KR_30_62	The ratio between two heavy and medium label partners.
Ratio H/M KR_30_63	The ratio between two heavy and medium label partners.
Ratio H/M normalized KR_30_6	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_30_61	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_30_62	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_30_63	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M unmod. pep. KR_30_6	
Ratio H/M localized KR_30_6	
Ratio H/M variability [%] KR_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type KR_30_6	
Occupancy L KR_30_6	
Occupancy M KR_30_6	
Occupancy H KR_30_6	
Intensity	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L	
Ratio mod/base M	
Ratio mod/base H	

Intensity IB_1_24	Summed up extracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a label of a label of the identified AB sequence in case of a label of the identified AB sequence.
	labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L IB_1_24	
Ratio mod/base M IB_1_24	
Ratio mod/base H IB_1_24	
Intensity IB_30_6	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L IB_30_6	
Ratio mod/base M IB_30_6	
Ratio mod/base H IB_30_6	
Intensity IB2_1_24	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB2_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB2_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB2_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L IB2_1_24	
Ratio mod/base M IB2_1_24	
Ratio mod/base H IB2_1_24	
Intensity IB2_30_6	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB2_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB2_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB2_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L IB2_30_6	
Ratio mod/base M IB2_30_6	
Ratio mod/base H IB2_30_6	
Intensity KR_1_24	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L KR_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M KR_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H KR_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L KR_1_24	
Ratio mod/base M KR_1_24	
Ratio mod/base H KR_1_24	
Intensity KR_30_6	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L KR_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M KR_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H KR_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Ratio mod/base L KR_30_6	

<b>-</b>	T
Ratio mod/base M KR_30_6	
Ratio mod/base H KR_30_6	
Occupancy IB_1_24	
Occupancy ratioIB_1_24	
Occupancy error scale IB_1_24	
Occupancy IB_30_6	
Occupancy ratioIB_30_6	
Occupancy error scale IB_30_6	
Occupancy IB2_1_24	
Occupancy ratioIB2_1_24	
Occupancy error scale IB2_1_24	
Occupancy IB2_30_6	
Occupancy ratioIB2_30_6	
Occupancy error scale IB2_30_6	
Occupancy KR_1_24	
Occupancy ratioKR_1_24	
Occupancy error scale KR_1_24	
Occupancy KR_30_6	
Occupancy ratioKR_30_6	
Occupancy error scale KR_30_6	
Reverse	When marked with '+', this particular peptide was found to be part of a protein derived from the reversed part of the protein sequence database. These should be removed for further data analysis.
Potential contaminant	When marked with '+', this particular peptide was found to be part of a commonly occurring contaminant. These should be removed for further data analysis.
id	A unique (consecutive) identifier for each row in the site table, which is used to cross-link the information in this file with the information stored in the other files.
Protein group IDs	The identifier of the protein-group this peptide sequence is associated with, which can be used to look up the extended protein information in the file 'proteinGroups.txt'.  As a single peptide can be linked to multiple proteins (e.g. in the case of razor-proteins), multiple id's can be stored here separated by a semicolon.  As a protein can be identified by multiple peptides, the same id can be found in different rows.
Positions	The positions of the modifications in the protein amino acid sequence.
Position	The position of the modification in the protein amino acid sequence.
Peptide IDs	Identifier(s) of the associated peptide sequence(s) summary, which can be found in the file 'peptides.txt'.
Mod. peptide IDs	Identifier(s) of the associated peptide sequence(s) summary, which can be found in the file 'modificationSpecificPeptides.txt'.
Evidence IDs	Identifier(s) for analyzed peptide evidence associated with the protein group referenced against the evidence table.
MS/MS IDs	The identifiers of the MS/MS scans identifying this peptide, referenced against the msms table.
Best localization evidence ID	
Best localization MS/MS ID	
Best localization raw file	
Best localization scan number	
Best localization scan number	
Best localization scan number Best score evidence ID	
Best localization scan number Best score evidence ID Best score MS/MS ID	
Best localization scan number Best score evidence ID Best score MS/MS ID Best score raw file	
Best localization scan number Best score evidence ID Best score MS/MS ID Best score raw file Best score scan number	
Best localization scan number  Best score evidence ID  Best score MS/MS ID  Best score raw file  Best score scan number  Best PEP evidence ID	

### Protein groups

The Protein Groups table contains information on the identified proteins in the processed raw-files. Each single row contains the group of proteins that could be reconstructed from a set of peptides.

Name	Separator	Description
Protein IDs		Identifiers of proteins contained in the protein group. They are sorted by number of identified peptides in descending order.
Majority protein IDs		These are the IDs of those proteins that have at least half of the peptides that the leading protein has.
Peptide counts (all)		Number of peptides associated with each protein in protein group, occurring in the order as the protein IDs occur in the 'Protein IDs' column. Here distinct peptide sequences are counted. Modified forms or different charges are counted as one peptide.
Peptide counts (razor+unique)		Number of peptides associated with each protein in protein group, occurring in the order as the protein IDs occur in the 'Protein IDs' column. Here distinct peptide sequences are counted. Modified forms or different charges are counted as one peptide.
Peptide counts (unique)		Number of peptides associated with each protein in protein group, occurring in the order as the protein IDs occur in the 'Protein IDs' column. Here distinct peptide sequences are counted. Modified forms or different charges are counted as one peptide.
Fasta headers		Fasta headers(s) of protein(s) contained within the group.
Number of proteins		Number of proteins contained within the group. This corresponds to the number of entries in the column 'Protein IDs'.
Peptides		The total number of peptide sequences associated with the protein group (i.e. for all the proteins in the group).
Razor + unique peptides		The total number of razor + unique peptides associated with the protein group (i.e. these peptides are shared with another protein group).
Unique peptides		The total number of unique peptides associated with the protein group (i.e. these peptides are not shared with another protein group).
Peptides IB_1_24		Number of peptides (distinct peptide sequences) in experiment IB_1_24
Peptides IB_30_6		Number of peptides (distinct peptide sequences) in experiment IB_30_6
Peptides IB2_1_24		Number of peptides (distinct peptide sequences) in experiment IB2_1_24
Peptides IB2_30_6		Number of peptides (distinct peptide sequences) in experiment IB2_30_6
Peptides KR_1_24		Number of peptides (distinct peptide sequences) in experiment KR_1_24
Peptides KR_30_6		Number of peptides (distinct peptide sequences) in experiment KR_30_6
Razor + unique peptides IB_1_24		Number of razor + unique peptides (distinct peptide sequences) in experiment IB_1_24
Razor + unique peptides IB_30_6		Number of razor + unique peptides (distinct peptide sequences) in experiment IB_30_6
Razor + unique peptides IB2_1_24		Number of razor + unique peptides (distinct peptide sequences) in experiment IB2_1_24
Razor + unique peptides IB2_30_6		Number of razor + unique peptides (distinct peptide sequences) in experiment IB2_30_6
Razor + unique peptides KR_1_24		Number of razor + unique peptides (distinct peptide sequences) in experiment KR_1_24
Razor + unique peptides KR_30_6		Number of razor + unique peptides (distinct peptide sequences) in experiment KR_30_6
Unique peptides IB_1_24		Number of unique peptides (distinct peptide sequences) in experiment IB_1_24
Unique peptides IB_30_6		Number of unique peptides (distinct peptide sequences) in experiment IB_30_6
Unique peptides IB2_1_24		Number of unique peptides (distinct peptide sequences) in experiment IB2_1_24
Unique peptides IB2_30_6		Number of unique peptides (distinct peptide sequences) in experiment IB2_30_6
Unique peptides KR_1_24		Number of unique peptides (distinct peptide sequences) in experiment KR_1_24
Unique peptides KR_30_6		Number of unique peptides (distinct peptide sequences) in experiment KR_30_6

Sequence coverage [%]	Percentage of the sequence that is covered by the identified
Unique i razer coguence coverage	peptides of the best protein sequence contained in the group.  Percentage of the sequence that is covered by the identified
Unique + razor sequence coverage [%]	unique and razor peptides of the best protein sequence contained in the group.
Unique sequence coverage [%]	Percentage of the sequence that is covered by the identified unique peptides of the best protein sequence contained in the group.
Mol. weight [kDa]	Molecular weight of the leading protein sequence contained in the protein group.
Sequence length	The length of the leading protein sequence contained in the group.
Sequence lengths	The length of all sequences of the proteins contained in the group.
Fraction average	3 - 1
Fraction 1	
Fraction 2	
Fraction 3	
Fraction 4	
Fraction 5	
Fraction 6	
Fraction 7	
Fraction 8	
Fraction 9	
Fraction 10	
Fraction 11	
Fraction 12	
Fraction 13	
Fraction 14	
Fraction 15	
Fraction 16	
Fraction 17	
Fraction 18	
Fraction 19	
Fraction 20	
Fraction 31	
Fraction 32	
Fraction 33	
Fraction 34	
Fraction 35	
Fraction 41	
Fraction 42	
Fraction 43	
Fraction 44	
Fraction 45	
Fraction 46	
Fraction 47	
Fraction 48	
Fraction 49	
Fraction 50	
Fraction 51	
Fraction 52	
Fraction 53	
Fraction 54	
Fraction 55	
Fraction 56	
Fraction 57	
Fraction 58	
Fraction 59	
Fraction 60	
Fraction 71	
Fraction 72	
Fraction 73	
Fraction 74	
Fraction 75	
Q-value	This is the ratio of reverse to forward protein groups.
	, is the rate of toroide to formal a protein groups.

Protein score which is derived from peptide posterior error probabilities.
Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Indicates whether this experiment was identified by MS/MS or only by matching between runs.
Indicates whether this experiment was identified by MS/MS or only by matching between runs.
The ratio between two medium and light label partners.
Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Number of redundant peptides (MS1 features) used for quantitation.
Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
The ratio between two heavy and light label partners.
Normalized ratio between two medium and light label partners.  The median of ratio sub-populations was shifted to 1.
Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Number of redundant peptides (MS1 features) used for quantitation.
Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
The ratio between two heavy and medium label partners.
Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Number of redundant peptides (MS1 features) used for quantitation.
Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
The ratio between two medium and light label partners.
The ratio between two medium and light label partners.
The ratio between two medium and light label partners.
The ratio between two medium and light label partners.
The ratio between two medium and light label partners.
The ratio between two medium and light label partners.
Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Normalized ratio between two heavy and light label partners.  The median of ratio sub-populations was shifted to 1.
Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Normalized ratio between two heavy and light label partners. The median of ratio sub-populations was shifted to 1.
Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the
natural logarithm of ratios times 100.
Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.

Ratio M/L variability [%] IB2_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L variability [%] KR_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L variability [%] KR_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio M/L count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio M/L iso-count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L iso-count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L iso-count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L iso-count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L iso-count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L iso-count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio M/L type IB_1_24	quantitation that are quantified with the re-quantity method.
Ratio M/L type IB_30_6	
,,	
Ratio M/L type IB2_1_24	
Ratio M/L type IB2_30_6	
Ratio M/L type KR_1_24	
Ratio M/L type KR_30_6	
Ratio H/L IB_1_24	The ratio between two heavy and light label partners.
Ratio H/L IB_30_6	The ratio between two heavy and light label partners.
Ratio H/L IB2_1_24	The ratio between two heavy and light label partners.
Ratio H/L IB2_30_6	The ratio between two heavy and light label partners.
Ratio H/L KR_1_24	The ratio between two heavy and light label partners.
Ratio H/L KR_30_6	The ratio between two heavy and light label partners.
Ratio H/L normalized IB_1_24	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB_30_6	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB2_1_24	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized IB2_30_6	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_1_24	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L normalized KR_30_6	Normalized ratio between two medium and light label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/L variability [%] IB_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L variability [%] IB_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L variability [%] IB2_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L variability [%] IB2_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L variability [%] KR_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/L variability [%] KR_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.

Ratio H/L count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/L iso-count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L iso-count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L iso-count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L iso-count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L iso-count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L iso-count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/L type IB_1_24	
Ratio H/L type IB_30_6	
Ratio H/L type IB2_1_24	
Ratio H/L type IB2_30_6	
Ratio H/L type KR_1_24	
Ratio H/L type KR_30_6	
Ratio H/M IB 1 24	The ratio between two heavy and medium label partners.
Ratio H/M IB 30 6	The ratio between two heavy and medium label partners.
Ratio H/M IB2_1_24	The ratio between two heavy and medium label partners.
Ratio H/M IB2_30_6	The ratio between two heavy and medium label partners.
Ratio H/M KR_1_24	The ratio between two heavy and medium label partners.
Ratio H/M KR_30_6	The ratio between two heavy and medium label partners.
Ratio H/M normalized IB_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB_30_6	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB2_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized IB2_30_6	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_1_24	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M normalized KR_30_6	Normalized ratio between two heavy and medium label partners. The median of ratio sub-populations was shifted to 1.
Ratio H/M variability [%] IB_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M variability [%] IB_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M variability [%] IB2_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M variability [%] IB2_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M variability [%] KR_1_24	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M variability [%] KR_30_6	Coefficient of variability over all redundant quantifiable peptides. It is calculated as the standard deviation of the natural logarithm of ratios times 100.
Ratio H/M count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation.

Ratio H/M count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation.
Ratio H/M iso-count IB_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M iso-count IB_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M iso-count IB2_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M iso-count IB2_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M iso-count KR_1_24	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M iso-count KR_30_6	Number of redundant peptides (MS1 features) used for quantitation that are quantified with the re-quantify method.
Ratio H/M type IB_1_24	
Ratio H/M type IB_30_6	
Ratio H/M type IB2_1_24	
Ratio H/M type IB2_30_6	
Ratio H/M type KR_1_24	
Ratio H/M type KR_30_6	
Sequence coverage IB_1_24 [%]	Percentage of the sequence that is covered by the identified peptides in this sample of the longest protein sequence contained within the group.
Sequence coverage IB_30_6 [%]	Percentage of the sequence that is covered by the identified peptides in this sample of the longest protein sequence contained within the group.
Sequence coverage IB2_1_24 [%]	Percentage of the sequence that is covered by the identified peptides in this sample of the longest protein sequence contained within the group.
Sequence coverage IB2_30_6 [%]	Percentage of the sequence that is covered by the identified peptides in this sample of the longest protein sequence contained within the group.
Sequence coverage KR_1_24 [%]	Percentage of the sequence that is covered by the identified peptides in this sample of the longest protein sequence contained within the group.
Sequence coverage KR_30_6 [%]	Percentage of the sequence that is covered by the identified peptides in this sample of the longest protein sequence contained within the group.
Intensity	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Intensity IB_1_24	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Intensity IB_30_6	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Intensity IB2_1_24	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB2_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB2_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.

Intensity H IB2_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Intensity IB2_30_6	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L IB2_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M IB2_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H IB2_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Intensity KR_1_24	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L KR_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M KR_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H KR_1_24	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Intensity KR_30_6	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L KR_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M KR_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H KR_30_6	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Only identified by site	When marked with '+', this particular protein group was identified only by a modification site.
Reverse	When marked with '+', this particular protein group contains no protein, made up of at least 50% of the peptides of the leading protein, with a peptide derived from the reversed part of the decoy database. These should be removed for further data analysis. The 50% rule is in place to prevent spurious protein hits to erroneously flag the protein group as reverse.
Potential contaminant	When marked with '+', this particular protein group was found to be a commonly occurring contaminant. These should be removed for further data analysis.
id	A unique (consecutive) identifier for each row in the proteinGroups table, which is used to cross-link the information in this file with the information stored in the other files.
Peptide IDs	Identifier(s) of the associated peptide sequence(s) summary, which can be found in the file 'peptides.txt'.
Peptide is razor	Indicates for each peptide ID if it is a razor or group unique peptide (true) or a non unique non razor peptide (false).
Mod. peptide IDs	
Evidence IDs	
MS/MS IDs	
Best MS/MS	The identifier of the best (in terms of quality) MS/MS scans identifying the peptides of this protein, referenced against the msms table.
Deamidation (NQ) site IDs	Identifier(s) for site(s) associated with the protein group, which show(s) evidence of the modification, referenced against the appropriate modification site file.
Oxidation (M) site IDs	Identifier(s) for site(s) associated with the protein group, which show(s) evidence of the modification, referenced against the appropriate modification site file.
Phospho (STY) site IDs	Identifier(s) for site(s) associated with the protein group, which show(s) evidence of the modification, referenced against the appropriate modification site file.
Deamidation (NQ) site positions	Positions of the sites in the leading protein of this group.
Oxidation (M) site positions	Positions of the sites in the leading protein of this group.
Phospho (STY) site positions	Positions of the sites in the leading protein of this group.

### All peptides

The type of detection for the peptide. MULTI – A labeling multiple traval detected.  So – An isotope pattern was detected.  The charge state of the peptide.  The charge state of the peptide.  The mass of the neutral peptide ((m²z-proton) " charge).  Mass — The mass of the neutral peptide ((m²z-proton) " charge).  Mass — The mass of the neutral peptide ((m²z-proton) " charge).  Mass — The mass of the neutral peptide ((m²z-proton) " charge).  Mass — The mass of the neutral peptide ((m²z-proton) " charge).  Mass — The mass of the neutral peptide ((m²z-proton) " charge).  Mumber of data points on the peptide measured in Full Width at Half Maximum (FWHM).  Number of data points peptide feature are overlapping with.  Number of scans — The number of Ms scans that the 3d peaks of this peptide feature are overlapping with.  Number of isotopic peaks — The number of isotopic peaks contained in this peptide feature are overlapping with.  Number of isotopic peaks contained in this peptide feature are overlapping with.  Number of isotopic peaks contained in this peptide feature are overlapping with.  Number of isotopic peaks contained in this peptide feature are overlapping with.  Number of isotopic peaks contained in this peptide feature are overlapping with.  Number of sotopic peaks contained in this peptide feature.  Short for Parent lon Fraction; indicates the fraction the target peak makes up of the total intensity in the inclusion window.  Mass fractional part — The values after the radix point (ie value - floor(value)).  Mass fractional part — The values after the radix point (ie value - floor(value)).  Mass precision (ppm) — The precision of the peptide in parts-per million.  Max intensity m/2 0 — Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence, in case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Max intensity m/2 2 — Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the i	Name	Separator	Description
multiplet was detected. ISO — An isotope pattern was detected. Incalibrated m/2  The charge state of the peptide. Incalibrated m/2  Mass  The mass of the neutral peptide ((m/2-proton) * charge). Incalibrated m/2  Mass of the neutral peptide ((m/2-proton) * charge). Incalibrated m/2  Mumber of data points (peak centroids) collected for this peptide feature.  Number of data points (peak centroids) collected for this peptide feature.  Number of scans  The number of MS scans that the 3d peaks of this peptide feature.  Number of isotopic peaks  The number of isotopic peaks contained in this peptide feature are overlapping with.  Number of isotopic peaks  Short for Parent Ion Fraction: indicates the fraction the target peak makes up of the total intensity in the inclusion window.  Mass Iractional part  The values after the radix point (ie value - floor/value).  Mass deficit  Empirically derived deviation measure to the next nearest integer scaled to center around 0. Can be used to visually detect containmants in a plot setting Mass against this value.  Mass precision (ppm)  The precision of the mass detection of the peptide in parts-per million.  Max intensity m/2 0  Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AX sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Max intensity m/2 1  Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AX sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Max intensity m/2 2  Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AX sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Retention length  The retention limp of the peak d	Raw file		Name of the raw file the spectral data was extracted from.
ISO _ An isotope pattern was detected.	Туре		
The mass divided by the charge of the charged peptide. Mass Incalibrated m/2 m/2 before re-calibrations have been applied.  The mass of the neutral peptide ((m/2-proton) " charge).  Incalibrated m/2 m/2 before re-calibrations have been applied.  The resolution of the peak detected for the peptide measured in Full Width at Half Maximum (FWHM).  The number of data points (peak centroids) collected for this peptide feature.  Sumber of scans The number of MS scans that the 3d peaks of this peptide feature re-coveraging with the status are overlapping with the status of the status are overlapping with the status of the st			
The mass of the neutral peptide ((m/z-proton) * charge).  Incalibrated m/z  m/z before re-calibrations have been applied.  Resolution  The resolution of the peak detected for the peptide measured in Full Width at Half Maximum (FWHM).  The number of data points peak centroids) collected for this peptide feature.  Number of data points peaks on the peptide feature of the peptide feature are overlapping with.  Number of isotopic peaks  The number of isotopic peaks contained in this peptide feature.  Short for Parent Ion Fraction; indicates the fraction the target peak makes up of the total intensity in the inclusion window. The values after the radius point (fe value — Ionof value).  Mass deficit  Empirically derived deviation measure to the next nearest integer scaled to center around 0. Can be used to visually detect contaminants in a plot setting Mass against this value.  M*abs precision (ppm)  The precision of the mass detection of the peptide in parts-per million.  Max intensity m/z 0  Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Max intensity m/z 1  Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Setention length  The retention time of the peak detected for the peptide measured in minumber.  The retention time width of the peak (last time point) in seconds.  The retention time width of the peak was encountered. The last scan number at which the peak was encountered. When marked with ** in this particular MS/MS scan was identified as a peptide; when marked	Charge		The charge state of the peptide.
Juncalibrated m/z  Resolution  The resolution of the peak detected for the peptide measured in Full Witch at Half Maximum (FWHM).  The number of data points  The number of data points (peak centroids) collected for this peptide feature.  The number of scans  The number of scans that the 3d peaks of this peptide feature are overlapping with.  Number of isotopic peaks  The number of isotopic peaks ontained in this peptide feature are overlapping with.  Short for Parent Ion Fraction; indicates the fraction that page the peak makes up of the total intensity in the inclusion window.  The values after the radix point (ie value - floor(value)).  Mass fractional part  The values after the radix point (ie value - floor(value)).  Empirically derived deviation measure to the next nearest integer scaled to center around 0. Can be used to visually detect contaminants in a plot setting Mass against this value.  M"a+b - round(m"a+b) m: the peptide mass a: 0,99854  b: -0.04  Mass precision (ppm)  The precision of the mass detection of the peptide in parts-per million.  Max intensity m/z 0  Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AX sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Max intensity m/z 2  Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AX sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Retention time  Retention time  The retention time of the peak detected for the peptide measured in minutes.  The first scan number at which the peak was encountered. The first scan number at which the peak was encountered. The first scan number at which the peak was encountered. The last scan number at which the peak was encountered. The last scan number at which the peak was encountered. The last scan number at which the peak was encountered. When marked with "+ in particular MS/MS scan wa	m/z		The mass divided by the charge of the charged peptide.
Resolution  The resolution of the peak detected for the peptide measured in Full With at Half Maximum (FWHM).  The number of data points (peak centroids) collected for this peptide feature.  The number of MS scans that the 3d peaks of this peptide feature are overlapping with.  The number of isotopic peaks ontained in this peptide feature are overlapping with.  The number of isotopic peaks ontained in this peptide feature are overlapping with.  The number of isotopic peaks ontained in this peptide feature are overlapping with.  The values after the radix point (ie value - floor(value)).  The values after the radix point (ie value - floor(value)).  Empirically derived deviation measure to the next nearest integer scaled to center around 0. Can be used to visually detect contaminants in a plot setting Mass against this value.  m'a+b - round(m'a+b)  m' the peptide mass  a: 0.99854  b: -0.04  Max intensity m/z 0  Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment his is the total intensity of all the isotopic patterns in the label cluster.  Max intensity m/z 1  Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Max intensity m/z 2  Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Retention length  The retention time of the peak detected for the peptide measured in minutes.  Retention length  The first scan number at which the peak was encountered. When marked with by identified AA sequence in case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  The last scan number at which the peak was encountered. When marked with by ithis particular MS/MS	Mass		The mass of the neutral peptide ((m/z-proton) * charge).
In Full Width at Half Maximum (FWHM).  The number of data points peak centroids) collected for this peptide feature.  Number of scans  The number of Mis cans that the 3d peaks of this peptide feature are overlapping with.  The number of isotopic peaks  The number of Isotopic peaks contained in this peptide feature are overlapping with.  The number of Isotopic peaks contained in this peptide feature are overlapping with.  The number of Isotopic peaks contained in this peptide feature are overlapping with.  Short for Parent Ion Fraction; indicates the fraction the target peak makes up of the total intensity in the inclusion window.  In the values after the radik point ((e value - Indor(value)).  In the values after the radik point ((e value - Indor(value)).  Empirically derived deviation measure to the next nearest integer scaled to center around 0. Can be used to visually detect contaminants in a plot setting Mass against this value.  In the peptide mass  2. 0.98954  2. 0.09854  2. 0.09854  3. 0.989554  3. 0.98956  3. 0.989	Uncalibrated m/z		m/z before re-calibrations have been applied.
peptide feature.  The number of Scans that the 3d peaks of this peptide feature are overlapping with.  The number of isotopic peaks contained in this peptide feature are overlapping with.  The number of isotopic peaks contained in this peptide feature.  Short for Parent lon Fraction; indicates the fraction the target peak makes up of the total intensity in the inclusion window. Mass fractional part  Mass deficit  Empirically derived deviation measure to the next nearest integer scaled to center around 0. Can be used to visually detect contaminants in a plot setting Mass against this value.  m°a+b – round(m°a+b)  m; the peptide mass a: 0.99954  b: -0.04  Mass precision [ppm]  The precision of the mass detection of the peptide in parts-per million.  Max intensity m/z 0  Summed up eXtracted ton Current (XIC) of all isotopic clusters associated with the identified As sequence, in case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Summed up eXtracted ton Current (XIC) of all isotopic clusters associated with the identified As sequence, in case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Summed up eXtracted ton Current (XIC) of all isotopic clusters associated with the identified As sequence, in case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Summed up eXtracted ton Current (XIC) of all isotopic clusters associated with the identified As expendence, in case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Retention time  The retention time of the peak detected for the peptide measured in minutes.  Retention length  The fold retention time width of the peak (last time point – first time point) in seconds.  The first scan number at which the peak was encountered.  When marked with '+ this particular MS/MS scan was assigned to a decoy sequence.  When marked with '+ this particular MS/MS sca	Resolution		
teature are overlapping with.  The number of isotopic peaks on the peptide feature  Short for Parent Ion Fraction; indicates the fraction the target peak makes up of the total intensity in the inclusion window.  Mass fractional part  The values after the radix point (ie value - floor(value)).  Empirically derived deviation measure to the next nearest integer scaled to cental part of visually detect contaminants in a plot setting Mass against this value.  m°a+b – round(m°a+b) m; the peptide mass a: 0.99954 b: -0.04  Mass precision [ppm]  The precision of the mass detection of the peptide in parts-per million.  Max intensity m/z 0  Summed up extracted fon Current (XIC) of all isotopic clusters associated with the identified AS expence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Summed up extracted fon Current (XIC) of all isotopic clusters associated with the identified AS expence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Summed up extracted fon Current (XIC) of all isotopic clusters associated with the identified AS expence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Summed up extracted fon Current (XIC) of all isotopic clusters associated with the identified AS expence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Retention time  The total retention time width of the peak detected for the peptide measured in minutes.  Retention length  The total retention time width of the peak (last time point – first impoint) in seconds.  The first scan number at which the peak was encountered.  When marked with "this particular MS/MS scan was assigned to a decoy sequence.  When marked with "this particular MS/MS scan was assigned to a decoy sequence.  When marked with "this particular MS/MS scan was assigned to a decoy sequence.  The identified As aseque	Number of data points		
Short for Parent Ion Fraction; indicates the fraction the target peak makes up of the total intensity in the inclusion window.  Mass fractional part  The values after the radix point (ie value - floor(value)).  Empirically derived deviation measure to the next nearest integer scaled to center around 0. Can be used visually detect contaminants in a plot setting Mass against this value.  m*a+b - round(m*a+b) m* the peptide mass a: 0.99594 b: -0.04  Mass precision [ppm]  The precision of the mass detection of the peptide in parts-per million.  Max intensity m/z 0  Summed up extracted lon Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Max intensity m/z 1  Summed up extracted lon Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Summed up extracted lon Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Summed up extracted lon Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Retention time  Retention time  The retention time of the peak detected for the peptide measured in minutes.  Retention length  The retention time width of the peak (last time point) – first time point) in seconds.  Retention length (FWHM)  The full width at half maximum value retention time width of the peak in seconds.  The first scan number at which the peak was encountered.  When marked with '+' this particular MS/MS can was assigned to a decoy sequence.  When marked with '+' this particular MS/MS scan was assigned to a decoy sequence.  The intensity of the sequence of t	Number of scans		
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Empirically derived deviation measure to the next nearest integer scaled to center around 0. Can be used to visually detect contaminants in a plot setting Mass against this value.  m*a+b - round(m*a+b) m: the peptide mass a: 0.99354 b: -0.04  Mass precision [ppm]  The precision of the mass detection of the peptide in parts-per million.  Max intensity m/z 0  Summed up extracted lon Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Summed up extracted lon Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Summed up extracted lon Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Summed up extracted lon Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Summed up extracted lon Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Retention time  Retention length  The retention time of the peak detected for the peptide measured in minutes.  Retention length  The total retention time width of the peak (last time point) in seconds.  Retention length (FWHM)  The full width at half maximum value retention time width of the peak in seconds.  The issue can number at which the peak was encountered.  When marked with '-' this particular MS/MS scan was identified as a peptide, when marked with '-' no identification was made.  Unique identified AA sequence of the peptide.  The length of the sequence of the peptide.  Prot	PIF		
integer scaled to center around 0. Can be used to visually detect contaminants in a plot setting Mass against this value.  m°a+b – round(m°a+b) m: the peptide mass a: 0.99954 b: 0.04  Mass precision (ppm)  The precision of the mass detection of the peptide in parts-per million.  Max intensity m/z 0  Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Max intensity m/z 1  Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Max intensity m/z 2  Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Max intensity m/z 2  Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Retention time  The retention time of the peak detected for the peptide measured in minutes.  Retention length  The total retention time width of the peak (last time point – first time point) in seconds.  Retention length (FWHM)  The full width at half maximum value retention time width of the peak in seconds.  The last scan number at which the peak was encountered.  When marked with '+' this particular MS/MS scan was assigned to a decoy sequence.  When marked with '+' this particular MS/MS scan was assigned to a decoy sequence.  When marked with '+' this particular MS/MS scan was assigned to a decoy sequence.  The length of the sequence stored in the column "Sequence".  Note: This column only set when this MS/MS spectrum has been identified.  Note: This column only set when this MS/MS spectrum	Mass fractional part		The values after the radix point (ie value - floor(value)).
a: 0.99954   b: -0.04	Mass deficit		integer scaled to center around 0. Can be used to visually detect contaminants in a plot setting Mass against this value.  m*a+b - round(m*a+b)
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associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Max intensity m/z 2  Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Retention time  The retention time of the peak detected for the peptide measured in minutes.  Retention length  The total retention time width of the peak (last time point – first time point) in seconds.  Retention length (FWHM)  The full width at half maximum value retention time width of the peak in seconds.  The first scan number at which the peak was encountered.  Max scan number  The last scan number at which the peak was encountered.  When marked with '-' this particular MS/MS scan was identified as a peptide; when marked with '-' no identification was made.  Reverse  When marked with '+' this particular MS/MS scan was assigned to a decoy sequence.  MS/MS IDS  Unique identifier linking this identification to the MS/MS scans.  Sequence  The identified AA sequence of the peptide.  The length of the sequence stored in the column "Sequence".  Note: This column only set when this MS/MS spectrum has been identified.  Post-translational modifications contained within the sequence of modified AAs.  Note: This column only set when this MS/MS spectrum has been identified.	Mass precision [ppm]		The precision of the mass detection of the peptide in parts-per- million.
associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  Retention time  The retention time of the peak detected for the peptide measured in minutes.  Retention length  The total retention time width of the peak (last time point – first time point) in seconds.  Retention length (FWHM)  The full width at half maximum value retention time width of the peak in seconds.  The first scan number at which the peak was encountered.  Max scan number  The last scan number at which the peak was encountered.  When marked with '+' this particular MS/MS scan was identified as a peptide; when marked with '-' no identification was made.  Reverse  When marked with '+' this particular MS/MS scan was assigned to a decoy sequence.  Wish IDs  Unique identifier linking this identification to the MS/MS scans.  Sequence  The identified AA sequence of the peptide.  Post-translational modifications contained within the sequence when no modifications exist, this is set to 'unmodified'.  Note: This column only set when this MS/MS spectrum has been identifier of proteins this peptide is associated with.  Note: This column only set when this MS/MS spectrum has been identifier.	Max intensity m/z 0		labeled experiment this is the total intensity of all the isotopic
associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.  The retention time of the peak detected for the peptide measured in minutes.  The total retention time width of the peak (last time point – first time point) in seconds.  Retention length (FWHM)  The full width at half maximum value retention time width of the peak in seconds.  The full width at half maximum value retention time width of the peak in seconds.  The first scan number at which the peak was encountered.  Max scan number  The last scan number at which the peak was encountered.  When marked with '+' this particular MS/MS scan was identified as a peptide; when marked with '-' no identification was made.  Reverse  When marked with '+' this particular MS/MS scan was assigned to a decoy sequence.  MS/MS IDs  Unique identifier linking this identification to the MS/MS scans.  The identified AA sequence of the peptide.  The length of the sequence stored in the column "Sequence".  Modifications  Post-translational modifications contained within the sequence When no modifications exist, this is set to 'unmodified'.  Note: This column only set when this MS/MS spectrum has been identified.  Proteins  Identifiers of proteins this peptide is associated with.  Note: This column only set when this MS/MS spectrum has been identified.	Max intensity m/z 1		labeled experiment this is the total intensity of all the isotopic
Retention length  The total retention time width of the peak (last time point – first time point) in seconds.  Retention length (FWHM)  The full width at half maximum value retention time width of the peak in seconds.  The first scan number at which the peak was encountered.  Max scan number  The last scan number at which the peak was encountered.  When marked with '+' this particular MS/MS scan was identified as a peptide; when marked with '-' no identification was made.  Reverse  When marked with '+' this particular MS/MS scan was assigned to a decoy sequence.  MS/MS IDs  Unique identifier linking this identification to the MS/MS scans.  Sequence  The identified AA sequence of the peptide.  In length of the sequence stored in the column "Sequence".  Modifications  Post-translational modifications contained within the sequence When no modifications exist, this is set to 'unmodified'.  Note: This column only set when this MS/MS spectrum has been identified.  Proteins  Identifiers of proteins this peptide is associated with.  Note: This column only set when this MS/MS spectrum has been identified.	Max intensity m/z 2		labeled experiment this is the total intensity of all the isotopic
time point) in seconds.  Retention length (FWHM)  The full width at half maximum value retention time width of the peak in seconds.  The first scan number at which the peak was encountered.  The last scan number at which the peak was encountered.  When marked with '+' this particular MS/MS scan was identified as a peptide; when marked with '-' no identification was made.  Reverse  When marked with '+' this particular MS/MS scan was assigned to a decoy sequence.  Unique identifier linking this identification to the MS/MS scans.  Sequence  In identified AA sequence of the peptide.  The length of the sequence stored in the column "Sequence".  Wodifications  Post-translational modifications contained within the sequence When no modifications exist, this is set to 'unmodified'.  Note: This column only set when this MS/MS spectrum has been identified.  Proteins  Identifiers of proteins this peptide is associated with.  Note: This column only set when this MS/MS spectrum has been identified.	Retention time		
peak in seconds.  The first scan number at which the peak was encountered.  The last scan number at which the peak was encountered.  When marked with '+' this particular MS/MS scan was identified was a peptide; when marked with '-' no identification was made.  Reverse When marked with '+' this particular MS/MS scan was assigned to a decoy sequence.  MS/MS IDs Unique identifier linking this identification to the MS/MS scans.  Gequence The identified AA sequence of the peptide.  Length The length of the sequence stored in the column "Sequence".  Modifications Post-translational modifications contained within the sequence When no modifications exist, this is set to 'unmodified'.  Note: This column only set when this MS/MS spectrum has been identified.  Sequence representation of the peptide including location(s) or modified AAs.  Note: This column only set when this MS/MS spectrum has been identified.  Proteins Identifiers of proteins this peptide is associated with.  Note: This column only set when this MS/MS spectrum has been identified.	Retention length		The total retention time width of the peak (last time point – first time point) in seconds.
Max scan number  The last scan number at which the peak was encountered.  When marked with '+' this particular MS/MS scan was identified as a peptide; when marked with '-' no identification was made.  Reverse  When marked with '+' this particular MS/MS scan was assigned to a decoy sequence.  WS/MS IDs  Unique identifier linking this identification to the MS/MS scans.  Sequence  The identified AA sequence of the peptide.  The length of the sequence stored in the column "Sequence".  Wodifications  Post-translational modifications contained within the sequence When no modifications exist, this is set to 'unmodified'.  Note: This column only set when this MS/MS spectrum has been identified.  Proteins  Proteins  Identifiers of proteins this peptide is associated with.  Note: This column only set when this MS/MS spectrum has	Retention length (FWHM)		The full width at half maximum value retention time width of the peak in seconds.
When marked with '+' this particular MS/MS scan was identified as a peptide; when marked with '-' no identification was made.  Reverse  When marked with '+' this particular MS/MS scan was assigned to a decoy sequence.  MS/MS IDs  Unique identifier linking this identification to the MS/MS scans.  Gequence  The identified AA sequence of the peptide.  The length of the sequence stored in the column "Sequence".  Modifications  Post-translational modifications contained within the sequence When no modifications exist, this is set to 'unmodified'.  Note: This column only set when this MS/MS spectrum has been identified.  Sequence representation of the peptide including location(s) or modified AAs.  Note: This column only set when this MS/MS spectrum has been identified.  Proteins  Identifiers of proteins this peptide is associated with.  Note: This column only set when this MS/MS spectrum has	Min scan number		The first scan number at which the peak was encountered.
identified as a peptide; when marked with '-' no identification was made.  When marked with '+' this particular MS/MS scan was assigned to a decoy sequence.  MS/MS IDs  Unique identifier linking this identification to the MS/MS scans.  Sequence  In identified AA sequence of the peptide.  In length of the sequence stored in the column "Sequence".  Modifications  Post-translational modifications contained within the sequence When no modifications exist, this is set to 'unmodified'.  Note: This column only set when this MS/MS spectrum has been identified.  Sequence representation of the peptide including location(s) or modified AAs.  Note: This column only set when this MS/MS spectrum has been identified.  Proteins  Identifiers of proteins this peptide is associated with.  Note: This column only set when this MS/MS spectrum has	Max scan number		The last scan number at which the peak was encountered.
assigned to a decoy sequence.  MS/MS IDs  Unique identifier linking this identification to the MS/MS scans.  The identified AA sequence of the peptide.  The length of the sequence stored in the column "Sequence".  Modifications  Post-translational modifications contained within the sequence When no modifications exist, this is set to 'unmodified'.  Note: This column only set when this MS/MS spectrum has been identified.  Sequence representation of the peptide including location(s) o modified AAs.  Note: This column only set when this MS/MS spectrum has been identified.  Proteins  Identifiers of proteins this peptide is associated with.  Note: This column only set when this MS/MS spectrum has	Identified		identified as a peptide; when marked with '-' no identification
The identified AA sequence of the peptide.  The length of the sequence stored in the column "Sequence".  Modifications  Post-translational modifications contained within the sequence When no modifications exist, this is set to 'unmodified'.  Note: This column only set when this MS/MS spectrum has been identified.  Sequence representation of the peptide including location(s) or modified AAs.  Note: This column only set when this MS/MS spectrum has been identified.  Proteins  Identifiers of proteins this peptide is associated with.  Note: This column only set when this MS/MS spectrum has	Reverse		
The length of the sequence stored in the column "Sequence".  Post-translational modifications contained within the sequence When no modifications exist, this is set to 'unmodified'.  Note: This column only set when this MS/MS spectrum has been identified.  Sequence representation of the peptide including location(s) or modified AAs.  Note: This column only set when this MS/MS spectrum has been identified.  Proteins  Identifiers of proteins this peptide is associated with.  Note: This column only set when this MS/MS spectrum has	MS/MS IDs		Unique identifier linking this identification to the MS/MS scans.
Post-translational modifications contained within the sequence When no modifications exist, this is set to 'unmodified'.  Note: This column only set when this MS/MS spectrum has been identified.  Sequence representation of the peptide including location(s) or modified AAs.  Note: This column only set when this MS/MS spectrum has been identified.  Proteins  Identifiers of proteins this peptide is associated with.  Note: This column only set when this MS/MS spectrum has	Sequence		The identified AA sequence of the peptide.
When no modifications exist, this is set to 'unmodified'.  Note: This column only set when this MS/MS spectrum has been identified.  Sequence representation of the peptide including location(s) o modified AAs.  Note: This column only set when this MS/MS spectrum has been identified.  Proteins  Identifiers of proteins this peptide is associated with.  Note: This column only set when this MS/MS spectrum has	Length		The length of the sequence stored in the column "Sequence".
been identified.   Sequence representation of the peptide including location(s) of modified AAs.   Note: This column only set when this MS/MS spectrum has been identified.   Proteins   Identifiers of proteins this peptide is associated with.	Modifications		Post-translational modifications contained within the sequence. When no modifications exist, this is set to 'unmodified'.
modified AAs.  Note: This column only set when this MS/MS spectrum has been identified.  Proteins  Identifiers of proteins this peptide is associated with.  Note: This column only set when this MS/MS spectrum has			
been identified.  Proteins  Identifiers of proteins this peptide is associated with.  Note: This column only set when this MS/MS spectrum has	Modified sequence		Sequence representation of the peptide including location(s) of modified AAs.
Note: This column only set when this MS/MS spectrum has			
	Proteins		Identifiers of proteins this peptide is associated with.
Score The score of the identification (higher is better).	Score		The score of the identification (higher is better).

PEP	The posterior error probability of the identification (smaller is better).
Lys Count	The number of instances of Lys contained within the sequence. The value for this can reliably be determined in the case of label partners, based on the distance between the partners. These counts are used to solidify the peptide identification process.
Arg Count	The number of instances of Arg contained within the sequence. The value for this can reliably be determined in the case of label partners, based on the distance between the partners. These counts are used to solidify the peptide identification process.
Ratio M/L	The ratio between two medium and light multiplet members.
Ratio M/L normalized	Normalized ratio between two medium and light multiplet members. The median of the total ratio population was shifted to 1.
Ratio H/L	The ratio between two heavy and light multiplet members.
Ratio H/L normalized	Normalized ratio between two heavy and light multiplet members. The median of the total ratio population was shifted to 1.
Ratio H/M	The ratio between two heavy and medium multiplet members.
Ratio H/M normalized	Normalized ratio between two heavy and medium multiplet members. The median of the total ratio population was shifted to 1.
Intensity	Summed up eXtracted Ion Current (XIC) of all isotopic clusters associated with the identified AA sequence. In case of a labeled experiment this is the total intensity of all the isotopic patterns in the label cluster.
Intensity L	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the light label partner.
Intensity M	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the medium label partner.
Intensity H	Summed up eXtracted Ion Current (XIC) of the isotopic cluster belonging to the heavy label partner.
Intensities L	Elution profile of the light peptide.
Intensities M	Elution profile of the medium peptide.
Intensities H	Elution profile of the heavy peptide.
Isotope pattern L	Isotope pattern of the light peptide.
Isotope pattern M	Isotope pattern of the medium peptide.
Isotope pattern H	Isotope pattern of the heavy peptide.
MS/MS Count	The number of MS/MS spectra recorded for the peptide.
MSMS Scan Numbers	The scan numbers where the MS/MS spectra were recorded.
MSMS label States	The label partner detected for the peptide. The value 0 is always the light partner. In the case of double label labeling 1 is the heavy partner. In the case of triple label labeling 1 is the medium and 2 the heavy partner.
MSMS Isotope Indices	Indices of the isotopic peaks that the MS/MS spectra reside on. A value of 0 corresponds to the monoisotopic peak.
DP Mass Difference	Dependent peptide search: Mass Difference
DP Time Difference	Dependent peptide search: Time Difference
DP Score	Dependent peptide search: Score
DP PEP	Dependent peptide search: PEP
DP Positional Probability	Dependent peptide search: Positional Probability
DP Base Sequence	Dependent peptide search: Base Sequence
DP Probabilities DP AA	Dependent peptide search: Probabilities  Dependent peptide search: AA
DP Base Raw File	Dependent peptide search: AA  Dependent peptide search: Base Scan Number
DP Base Raw File  DP Base Scan Number	Dependent peptide search: Base Scan Number  Dependent peptide search: Base Scan Number
D. Dago Coall Nathbel	popondoni popilao scarcii. Dase scari Nullisel
DP Mod Scan Number	Dependent peptide search: Mod Scan Number
DP Mod Scan Number DP Decov	Dependent peptide search: Mod Scan Number  Dependent peptide search: Decov
DP Mod Scan Number DP Decoy DP Proteins	Dependent peptide search: Mod Scan Number  Dependent peptide search: Decoy  Dependent peptide search: Proteins
DP Decoy	Dependent peptide search: Decoy
DP Decoy DP Proteins	Dependent peptide search: Decoy Dependent peptide search: Proteins
DP Decoy DP Proteins DP Cluster Index	Dependent peptide search: Decoy Dependent peptide search: Proteins Dependent peptide search: Cluster Index
DP Decoy DP Proteins DP Cluster Index DP Cluster Mass	Dependent peptide search: Decoy Dependent peptide search: Proteins Dependent peptide search: Cluster Index Dependent peptide search: Cluster Mass
DP Decoy DP Proteins DP Cluster Index DP Cluster Mass DP Cluster Mass SD	Dependent peptide search: Decoy Dependent peptide search: Proteins Dependent peptide search: Cluster Index Dependent peptide search: Cluster Mass Dependent peptide search: Cluster Mass SD
DP Decoy DP Proteins DP Cluster Index DP Cluster Mass DP Cluster Mass SD DP Cluster Size Total	Dependent peptide search: Decoy Dependent peptide search: Proteins Dependent peptide search: Cluster Index Dependent peptide search: Cluster Mass Dependent peptide search: Cluster Mass SD Dependent peptide search: Cluster Size Total
DP Decoy DP Proteins DP Cluster Index DP Cluster Mass DP Cluster Mass SD DP Cluster Size Total DP Cluster Size Forward	Dependent peptide search: Decoy Dependent peptide search: Proteins Dependent peptide search: Cluster Index Dependent peptide search: Cluster Mass Dependent peptide search: Cluster Mass SD Dependent peptide search: Cluster Size Total Dependent peptide search: Cluster Size Forward
DP Decoy DP Proteins DP Cluster Index DP Cluster Mass DP Cluster Mass SD DP Cluster Size Total DP Cluster Size Forward DP Cluster Size Reverse	Dependent peptide search: Decoy Dependent peptide search: Proteins Dependent peptide search: Cluster Index Dependent peptide search: Cluster Mass Dependent peptide search: Cluster Mass SD Dependent peptide search: Cluster Size Total Dependent peptide search: Cluster Size Forward Dependent peptide search: Cluster Size Reverse
DP Decoy DP Proteins DP Cluster Index DP Cluster Mass DP Cluster Mass SD DP Cluster Size Total DP Cluster Size Forward DP Cluster Size Reverse DP Modification	Dependent peptide search: Decoy Dependent peptide search: Proteins Dependent peptide search: Cluster Index Dependent peptide search: Cluster Mass Dependent peptide search: Cluster Mass SD Dependent peptide search: Cluster Size Total Dependent peptide search: Cluster Size Forward Dependent peptide search: Cluster Size Reverse Dependent peptide search: Modification

DP Base Ratio H/L	Dependent peptide search: Base Ratio H/L
DP Base Ratio H/L Normalized	Dependent peptide search: Base Ratio H/L Normalized
DP Base Ratio H/M	Dependent peptide search: Base Ratio H/M
DP Base Ratio H/M Normalized	Dependent peptide search: Base Ratio H/M Normalized
DP Occupancy L	Dependent peptide search: Occupancy L
DP Occupancy M	Dependent peptide search: Occupancy M
DP Occupancy H	Dependent peptide search: Occupancy H

#### MS scans

The msScans table contains information about the full scans, which can be used to verify data quality and generated useful statistics about the interaction between the samples and LC.

Name	Separator	Description
Raw file		The name of the RAW-file the mass spectral data originates from.
Scan number		The scan number (defined in the raw-file) at which the full scan was made.
Scan index		The consecutive index of this full scan.
Retention time		The retention time at which the full scan was made.
Cycle time		The total time (full scan including the tandem MS scans) this full scan has taken up.
Ion injection time		The total injection time that was required to capture the specified amount of ions. This value is limited by a maximum, which can be used to determine whether the time has maxed out (indicative of a bad acquisition).
Base peak intensity		The intensity of the most intense ion in the spectrum.
Total ion current		The total intensity acquired in the full scan.
MS/MS count		The number of tandem MS scans that were made based on this full scan (e.g. a top 10 method selects the top 10 most intense ions in the scan and fragments those).
Mass calibration		The applied mass correction in Th to the full scan.
Fraction		The fraction measured with this full scan.
Experiment		
Peak length		The average time between the start and the end of the peaks detected in the full scan.
Isotope pattern length		The average time between the start and the end of the isotope patterns detected in the full scan.
Multiplet length		The average time between the start and the end of the isotope patterns of the labeling multiplets detected in the full scan.
Peaks / s		The average number of peaks detected per second of chromatography.
Single peaks / s		The average number of single peaks detected per second of chromatography.
Isotope patterns / s		The average number of isotope patterns detected per second of chromatography.
Single isotope patterns / s		The average number of single isotope patterns detected per of second chromatography.
Multiplets / s		The average number of labeling multiplets detected per of second chromatography.
Identified multiplets / s		The percentage of labeling multiplets actually identified.
Multiplet identification rate [%]		The percentage of the detected labeling multiplets that were identified.
MS/MS / s		The average number of MS/MS events per second of chromatography.
Identified MS/MS / s		The average number of identified MS/MS events per second of chromatography.
MS/MS identification rate [%]		The percentage of tandem MS scans that were identified.
Intens Comp Factor		Taken from the Thermo RAW file.
CTCD Comp		Taken from the Thermo RAW file.
RawOvFtT		For Thermo Fisher only. TIC estimation done with the orbitrap cell.
AGC Fill		Taken from the Thermo RAW file.

### MZ range

Name	Separator	Description
Raw file		The name of the RAW-file the mass spectral data was derived from.
m/z		The mass-over-charge value.
Peaks / Da		The average number of peaks detected per Dalton.
Single peaks / Da		The average number of single peaks detected per Dalton.
Isotope patterns / Da		The average number of isotope patterns detected per Dalton.
Single isotope patterns / Da		The average number of single isotope patterns detected per Dalton.
SILAC pairs / Da		The average number of SILAC pairs detected per Dalton.
Identified SILAC pairs / Da		The percentage of SILAC pairs actually identified.
SILAC identification rate [%]		The percentage of the detected SILAC pairs that were identified.
MS/MS / Da		The average number of MS/MS events per Dalton.
Identified MS/MS / Da		The average number of identified MS/MS events per Dalton.
Identification rate [%]		The percentage of tandem MS scans that were identified.

#### MS/MS scans

Name	Separator	Description
Raw file		Name of the RAW file the spectral MS/MS data was extracted from.
Scan number		RAW file derived scan number for the MS/MS spectrum.
Retention time		Time point along the elution profile at which the MS/MS data was recorded.
Ion injection time		The ion inject time for the MS/MS scan. This can be used to determine if this time equals to the maximum ion inject time, general indicative of a lower quality spectrum.
Total ion current		The total ion current of the MS/MS scan. For Thermo data this value is calculated by summing all the intensity values found in the mass spectral data, which is different from the Xcalibur reported TIC (Xcalibur TIC is about 25% of the value reported here).
Collision energy		The collision energy used for the fragmentation that resulted in this MS/MS scan.
Summations		For time of flight instruments only.
Base peak intensity		The intensity of the most intense ion in the spectrum.
Elapsed time		The time the MS/MS scan took to complete.
Identified		When marked with '+' this particular MS/MS scan was identified as a peptide; when marked with '-' no identification was made.
Matched		When marked with '+' this particular MS/MS scan was retrieved by matching between runs.
Reverse		When marked with '+' this particular MS/MS scan was assigned to a decoy sequence.
MS/MS IDs		Unique identifier linking this identification to the MS/MS scans.
Sequence		The identified AA sequence of the peptide.
Length		The length of the sequence stored in the column "Sequence".
Filtered peaks		Number of peaks after the 'top X per 100 Da' filtering.
m/z		Recalibrated m/z of the precursor ion.
Mass		Charge corrected mass of the precursor ion.
Charge		Charge state of the precursor ion.
Туре		The type of precursor ion as identified by MaxQuant. ISO – isotopic cluster. PEAK – single peak. MULTI – labeling cluster.
Fragmentation		The type of fragmentation used to create the MS/MS spectrum. CID – Collision Induced Dissociation. HCD – High energy Collision induced Dissociation. ETD – Electron Transfer Dissociation.
Mass analyzer		The mass analyzer used to record the MS/MS spectrum. ITMS – Ion trap. FTMS – Fourier transform ICR or orbitrap cell. TOF – Time of flight.
Parent intensity fraction		The percentage the parent ion intensity makes up of the total intensity in the selection window.
Fraction of total spectrum		The percentage the parent ion intensity makes up of the total intensity of the whole MS spectrum.
Base peak fraction		The percentage the parent ion intensity in comparison to the highest peak in he MS spectrum.
Precursor full scan number		The full scan number where the precursor ion was selected for fragmentation.
Precursor intensity		The intensity of the precursor ion at the scan number it was selected.
Precursor apex fraction		The fraction the intensity of the precursor ion makes up of the peak (apex) intensity.
Precursor apex offset		How many full scans the precursor ion is offset from the peak (apex) position.
Precursor apex offset time		How much time the precursor ion is offset from the peak (apex) position.
Scan event number		This number indicates which MS/MS scan this one is in the consecutive order of the MS/MS scans that are acquired after an MS scan.
Modifications		Post-translational modifications contained within the sequence. When no modifications exist, this is set to 'unmodified'.
		Note: This column only set when this MS/MS spectrum has been identified.

Modified sequence	Sequence representation of the peptide including location(s) of modified AAs.
	Note: This column only set when this MS/MS spectrum has been identified.
Proteins	Identifiers of proteins this peptide is associated with.
	Note: This column only set when this MS/MS spectrum has been identified.
Score	The score of the identification (higher is better).
PEP	The posterior error probability of the identification (smaller is better).
Fraction	The identifier of the fraction the sample was taken from.
Experiment	
Reporter PIF	
Reporter fraction	
DP mass difference	This dependent peptide's mass difference to the associated identified peptide.
DP time difference	This dependent peptide's time difference to the associated identified peptide.
DP score	The andromeda identification score.
DP PEP	Posterior Error Probability of the identification. This value essentially operates as a p-value, where smaller is more significant.
DP positional probability	
DP base sequence	
DP probabilities	
DP AA	
DP base raw file	
DP base scan number	
DP mod scan number	
DP decoy	
DP proteins	
DP cluster index	
DP cluster mass	
DP cluster mass SD	
DP cluster size total	
DP cluster size forward	
DP cluster size reverse	
DP modification	Post-translational modifications contained within the sequence. When no modifications exist, this is set to 'unmodified'.
DP peptide length difference	
Intens Comp Factor	Taken from the Thermo RAW file.
CTCD Comp	Taken from the Thermo RAW file.
RawOvFtT	For Thermo Fisher only. TIC estimation done with the orbitrap cell.
AGC Fill	Taken from the Thermo RAW file.
Scan index	Consecutive index of the MS/MS spectrum.
MS scan index	Consecutive index of the MS spectrum prior to this MS/MS spectrum.
MS scan number	Scan number of the MS spectrum prior to this MS/MS spectrum.

#### MS/MS

Name	Separator	Description
Raw file		The name of the RAW file the mass spectral data was read from.
Scan number		The RAW-file derived scan number of the MS/MS spectrum.
Scan index		The consecutive index of the MS/MS spectrum.
Sequence		The identified AA sequence of the peptide.
Length		The length of the sequence stored in the column "Sequence".
Missed cleavages		Number of missed enzymatic cleavages.
Modifications		Post-translational modifications contained within the identified peptide sequence.
Modified sequence		Sequence representation including the post-translational modifications (abbreviation of the modification in brackets before the modified AA). The sequence is always surrounded by underscore characters ('_').
Deamidation (NQ) Probabilities		Sequence representation of the peptide including PTM positioning probabilities ([01], where 1 is best match) for 'Deamidation (NQ)'.
Oxidation (M) Probabilities		Sequence representation of the peptide including PTM positioning probabilities ([01], where 1 is best match) for 'Oxidation (M)'.
Phospho (STY) Probabilities		Sequence representation of the peptide including PTM positioning probabilities ([01], where 1 is best match) for 'Phospho (STY)'.
Deamidation (NQ) Score diffs		
Oxidation (M) Score diffs		
Phospho (STY) Score diffs		
Acetyl (Protein N-term)		
Deamidation (NQ)		
Oxidation (M)		
Phospho (STY)		
Proteins		The identifiers of the proteins the identified peptide is associated with.
Charge		The charge state of the precursor ion.
Fragmentation		The type of fragmentation used to create the MS/MS spectrum. CID – Collision Induced Dissociation. HCD – High energy Collision induced Dissociation. ETD – Electron Transfer Dissociation.
Mass analyzer		The mass analyzer used to record the MS/MS spectrum. ITMS – Ion trap.  FTMS – Fourier transform ICR or orbitrap cell.  TOF – Time of flight.
Туре		The type of precursor ion as identified by MaxQuant. ISO – isotopic cluster.  PEAK – single peak.  MULTI – labeling cluster.
Scan event number		
Isotope index		
m/z		The mass-over-charge of the precursor ion.
Mass		The charge corrected mass of the precursor ion.
Mass error [ppm]		Mass error of the recalibrated mass-over-charge value of the precursor ion in comparison to the predicted monoisotopic mass of the identified peptide sequence expressed in parts per million.
Mass error [Da]		Mass error of the recalibrated mass-over-charge value of the precursor ion in comparison to the predicted monoisotopic mass of the identified peptide sequence expressed in atomic mass units.
Simple mass error [ppm]		
Retention time		The uncalibrated retention time in minutes where the MS/MS spectrum has been acquired.
PEP		Posterior Error Probability of the identification. This value essentially operates as a p-value, where smaller is more significant.
Score		Andromeda score for the best associated MS/MS spectrum.
Delta score		Score difference to the second best identified peptide with a different amino acid sequence.
Score diff		Score difference to the second best positioning of modifications identified peptide with the same amino acid sequence.
Localization prob		

Combinatorics	Number of possible distributions of the modifications over the peptide sequence.
Labeling state	Labeling state of the precursor isotope pattern used to identify the peptide.
PIF	Short for Parent Ion Fraction; indicates the fraction the target peak makes up of the total intensity in the inclusion window.
Fraction of total spectrum	The percentage the parent ion intensity makes up of the total intensity of the whole spectrum.
Base peak fraction	The percentage the parent ion intensity in comparison to the highest peak in he MS spectrum.
Precursor full scan number	The full scan number where the precursor ion was selected for fragmentation.
Precursor Intensity	The intensity of the precursor ion at the scan number it was selected.
Precursor apex fraction	The fraction the intensity of the precursor ion makes up of the peak (apex) intensity.
Precursor apex offset	How many full scans the precursor ion is offset from the peak (apex) position.
Precursor apex offset time	How much time the precursor ion is offset from the peak (apex) position.
Diagnostic peak Phospho (STY) Y	
Matches	The species of the peaks in the fragmentation spectrum after TooN filtering.
Intensities	The intensities of the peaks in the fragmentation spectrum after TopN filtering.
Mass deviations [Da]	The mass deviation of each peak in the fragmentation spectrum in absolute mass units.
Mass deviations [ppm]	The mass deviation of each peak in the fragmentation spectrum in parts per million.
Masses	The masses-over-charge of the peaks in the fragmentation spectrum.
Number of matches	The number of peaks matching to the predicted fragmentation spectrum.
Intensity coverage	The fraction of intensity in the MS/MS spectrum that is annotated.
Peak coverage	The fraction of peaks in the MS/MS spectrum that are annotated.
Neutral loss level	How many neutral losses were applied to each fragment in the Andromeda scoring.
ETD identification type	For ETD spectra several different combinations of ion series are scored. Here the highest scoring combination is indicated
Reverse	When marked with '+', this particular peptide was found to be part of a protein derived from the reversed part of the decoy database. These should be removed for further data analysis.
All scores	
All sequences	
All modified sequences	
Reporter PIF	
Reporter fraction	
id	A unique (consecutive) identifier for each row in the msms table, which is used to cross-link the information in this file with the information stored in the other files.
Protein group IDs	The identifier of the protein-group this redundant peptide sequence is associated with, which can be used to look up the extended protein information in the file 'proteinGroups.txt'. As a single peptide can be linked to multiple proteins (e.g. in the case of razor-proteins), multiple id's can be stored here separated by a semicolon.  As a protein can be identified by multiple peptides, the same id can be found in different rows.
Peptide ID	The identifier of the non-redundant peptide sequence.
Mod. peptide ID	Identifier of the associated modification summary stored in the file 'modificationSpecificPeptides.txt'.
Evidence ID	Identifier of the associated evidence stored in the file 'evidence.txt'.
Deamidation (NQ) site IDs	Identifier of the associated entry stored in the file 'Deamidation (NQ)Sites.txt'.
Oxidation (M) site IDs	Identifier of the associated entry stored in the file 'Oxidation (M)Sites.txt'.
Phospho (STY) site IDs	Identifier of the associated entry stored in the file 'Phospho (STY)Sites.txt'.

## Dependent peptides

Name	Separator	Description
Base peptide sequence		Base peptide sequence. Entries for the dependent peptides table get aggregated if they have the same base peptide sequence, modification and cluster index!
Modification		Modification. Entries for the dependent peptides table get aggregated if they have the same base peptide sequence, modification and cluster index!
Cluster index		Cluster index Entries for the dependent peptides table get aggregated if they have the same base peptide sequence, modification and cluster index!
Cluster mass		Cluster mass
Cluster mass SD		Cluster Mass SD
Cluster size total		Cluster Size Total
Cluster size forward		Cluster Size Forward
Cluster size reverse		Cluster Size Reverse
Base peptide raw file		This column contains the names of raw files that contained the original unmodified base peptide spectrum which were used to search for additional dependent peptides.
Matches in Raw files		This column represents all the raw files where the dependent Peptide could be found.
Туре		The type of detection for the dependent peptide. Multi – A labeling multiplet was detected. Iso – An isotope pattern was detected.
Localization probability		Base peptide sequence combined with the localization probability of possible canidates for the modification
Max Probability		The column contains the highest probability form an aminoacid canidate.
Posterior error probability		Posterior error probability
Aminoacid localizations		This contains all the possible amino acid candidates for the modification as well as information if they could be n or c - Terminal.
Feature Ids		The Feature Id can be used with the according raw file to trace a dependent peptide entry back in other tables. E.g. AllPeptides or MatchedFeatures - tables.
Found by		Information about how Identification and Quantification was acquired. Regular dependent peptide search: base raw file and matched raw file are equal. Dependent base peptide aggregation search: an other raw file was utilized to identify then the one the dep. peptide is in. Match unidentified features: an unidentified feature was detected by match unidentified features but not dependent peptide search.(dependent peptides are unidentified features)
Amount found by regular search		Information about how often specific Identification and Quantification was acquired.
Amount found by base peptide aggregation		Information about how often specific Identification and Quantification was acquired.
Amount found by match unidentified features		Information about how often specific Identification and Quantification was acquired.
AllPeptides Ids		This represents the index in the AllPeptides table of a dependent peptide entry. It's an easier way of finding the dependent peptide entry within the table than with the FeatureId!
Score		
Mass difference		Mass Difference
Time difference		Time Difference
Peptide Length Difference		Peptide Length Difference
Protein IDs		
Modification intensities		T 100 100 100 100 100 100 100 100 100 10
Modification scan numbers		The modification scan numbers can be used to search a dependent peptide entry in msmsScans table.
Base intensity Exp: IB_1_24		This column contains the base intensity for the experiment. If there are multiple dependent peptides found in this raw file than their intensities will get aggregated.
Base intensity Exp: IB_30_6		This column contains the base intensity for the experiment. If there are multiple dependent peptides found in this raw file than their intensities will get aggregated.
Base intensity Exp: IB2_1_24		This column contains the base intensity for the experiment. If there are multiple dependent peptides found in this raw file than their intensities will get aggregated.

Base intensity Exp: IB2_30_6	This column contains the base intensity for the experiment. If
, ,	there are multiple dependent peptides found in this raw file than their intensities will get aggregated.
Base intensity Exp: KR_1_24	This column contains the base intensity for the experiment. If there are multiple dependent peptides found in this raw file than their intensities will get aggregated.
Base intensity Exp: KR_30_6	This column contains the base intensity for the experiment. If there are multiple dependent peptides found in this raw file than their intensities will get aggregated.
Modification intensity Exp: IB_1_24	This column contains the modification intensity for the experiment. If there are multiple dependent peptides found in this raw file than their intensities will get aggregated.
Modification intensity Exp: IB_30_6	This column contains the modification intensity for the experiment. If there are multiple dependent peptides found in this raw file than their intensities will get aggregated.
Modification intensity Exp: IB2_1_24	This column contains the modification intensity for the experiment. If there are multiple dependent peptides found in this raw file than their intensities will get aggregated.
Modification intensity Exp: IB2_30_6	This column contains the modification intensity for the experiment. If there are multiple dependent peptides found in this raw file than their intensities will get aggregated.
Modification intensity Exp: KR_1_24	This column contains the modification intensity for the experiment. If there are multiple dependent peptides found in this raw file than their intensities will get aggregated.
Modification intensity Exp: KR_30_6	This column contains the modification intensity for the experiment. If there are multiple dependent peptides found in this raw file than their intensities will get aggregated.
Ratio mod/base intensity Exp: IB_1_24	This column contains the ratio between modification and base intensity for the specific experiment.
Ratio mod/base intensity Exp: IB_30_6	This column contains the ratio between modification and base intensity for the specific experiment.
Ratio mod/base intensity Exp:	This column contains the ratio between modification and base intensity for the specific experiment.
Ratio mod/base intensity Exp:	This column contains the ratio between modification and base intensity for the specific experiment.
Ratio mod/base intensity Exp: KR 1 24	This column contains the ratio between modification and base intensity for the specific experiment.
Ratio mod/base intensity Exp: KR 30 6	This column contains the ratio between modification and base intensity for the specific experiment.
Base Ratio H/L Exp: IB_1_24	Dependent peptide search: Base Ratio H/L
Base Ratio H/L Exp: IB_30_6	Dependent peptide search: Base Ratio H/L
Base Ratio H/L Exp: IB2_1_24	Dependent peptide search: Base Ratio H/L
Base Ratio H/L Exp: IB2_30_6	Dependent peptide search: Base Ratio H/L
Base Ratio H/L Exp: KR_1_24	Dependent peptide search: Base Ratio H/L
Base Ratio H/L Exp: KR_30_6	Dependent peptide search: Base Ratio H/L
Base Ratio H/L normalized Exp: IB_1_24	Base Ratio of heavy to light isotopes normalized
Base Ratio H/L normalized Exp: IB_30_6	Base Ratio of heavy to light isotopes normalized
Base Ratio H/L normalized Exp: IB2_1_24	Base Ratio of heavy to light isotopes normalized
Base Ratio H/L normalized Exp: IB2_30_6	Base Ratio of heavy to light isotopes normalized
Base Ratio H/L normalized Exp: KR_1_24	Base Ratio of heavy to light isotopes normalized
Base Ratio H/L normalized Exp: KR_30_6	Base Ratio of heavy to light isotopes normalized
Base Ratio M/L Exp: IB_1_24	Dependent peptide search: Base Ratio M/L
Base Ratio M/L Exp: IB_30_6	Dependent peptide search: Base Ratio M/L
Base Ratio M/L Exp: IB2_1_24	Dependent peptide search: Base Ratio M/L
Base Ratio M/L Exp: IB2_30_6	Dependent peptide search: Base Ratio M/L
Base Ratio M/L Exp: KR_1_24	Dependent peptide search: Base Ratio M/L
Base Ratio M/L Exp: KR_30_6	Dependent peptide search: Base Ratio M/L
Base Ratio M/L normalized Exp: IB_1_24	Dependent peptide search: Base Ratio M/L
Base Ratio M/L normalized Exp: IB_30_6	Dependent peptide search: Base Ratio M/L
Base Ratio M/L normalized Exp:	Dependent peptide search: Base Ratio M/L
Base Ratio M/L normalized Exp: IB2_30_6	Dependent peptide search: Base Ratio M/L
Base Ratio M/L normalized Exp: KR_1_24	Dependent peptide search: Base Ratio M/L
Base Ratio M/L normalized Exp: KR_30_6	Dependent peptide search: Base Ratio M/L
Base Ratio H/M Exp: IB_1_24	Dependent peptide search: Base Ratio M/L

Base Ratio H/M Exp: IB_30_6	Dependent peptide search: Base Ratio M/L
Base Ratio H/M Exp: IB2_1_24	Dependent peptide search: Base Ratio M/L
Base Ratio H/M Exp: IB2_30_6	Dependent peptide search: Base Ratio M/L
Base Ratio H/M Exp: KR_1_24	Dependent peptide search: Base Ratio M/L
Base Ratio H/M Exp: KR_30_6	Dependent peptide search: Base Ratio M/L
Base Ratio H/M normalized Exp: IB_1_24	Dependent peptide search: Base Ratio M/L
Base Ratio H/M normalized Exp: IB_30_6	Dependent peptide search: Base Ratio M/L
Base Ratio H/M normalized Exp: IB2_1_24	Dependent peptide search: Base Ratio M/L
Base Ratio H/M normalized Exp: IB2_30_6	Dependent peptide search: Base Ratio M/L
Base Ratio H/M normalized Exp: KR_1_24	Dependent peptide search: Base Ratio M/L
Base Ratio H/M normalized Exp: KR_30_6	Dependent peptide search: Base Ratio M/L