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LC-MS/MS Label-Free Proteomic Data Analysis Parameters V.2

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Protocol status: Working

We use this protocol and it's working

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Keywords: HuBMAP, BIOMIC, MSRC, Vanderbilt, Proteomics, MaxQuant, Data Analysis

Abstract

List of parameters and settings for searching label free proteomic data in MaxQuant version 1.6.7.



- 1 Label-free proteomic samples were searched using MaxQuant version 1.6.7.

- 2 *Group Specific Parameter settings included:*
Standard
Multiplicity = 1
Variable modifications: Oxidation (M); Acetylation (Protein N-term), Carbamidomethyl (C)
Fixed modifications: Carbamidomethyl (C)
Max number of modifications per peptide: 5
Instrument settings: Orbitrap (default settings)
Digestion: Specific, Trypsin /P
Max missed cleavages: 2
LFQ: None

- 3 *Global Parameters included:*
Database: UniProt Human Proteome (UP000005640, 9606) Reviewed, downloaded July 30, 2019
Min peptide length: 7
Max peptide mass: 4600 Da
MS/MS analyzer: Default settings
Identification: Default settings