DeepHalo Documentation

DeepHalo: A Hierarchically Optimized Workflow for High-Confidence and Highthroughput Detection of Halogenated Compounds in Complex HRMS Data Version 1.0.0

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Introduction

DeepHalo is a hierarchically optimized workflow designed for the detection and dereplication of halogenated compounds in **high-resolution** mass spectrometry (HRMS) data. It integrates cutting-edge deep learning models, robust isotope validation, comprehensive scoring, and dual dereplication strategies to deliver high accuracy and efficiency. Key applications include natural product discovery and halogenated metabolite annotation.

Data Support

DeepHalo supports **high-resolution** mass spectrometry (HRMS) data, with or without tandem mass spectrometry (MS/MS) information. Note that the MS² extraction function currently supports only data-dependent acquisition data (DDA). Additionally, DeepHalo exclusively supports the **.mzML** file format for analysis.

Core Features

- 1. Halogen Prediction
 - Element Prediction Model (EPM)
 - o Dual-branch Isotope Neural Network (IsoNN) architecture

- o High accuracy Cl/Br detection (>99.9% precision based on benchmark results)
- o Wide mass range coverage (50-2000 Da)
- o Robust interference resistance to B/Se/Fe/dehydro isomers

2. Isotope Pattern Validation

- Dual Validation System
 - o Mass Dimension: Statistical rule-based correction.
 - o Intensity Dimension: Autoencoder-based Anomaly Detection Model (ADM).

3. Multi-Level Halogen Confidence Scoring (H-score)

- Dual levels
 - o Prediction based on centroid-level isotope patterns
 - Prediction based on Scan-level isotope patterns
 - o H-score integration for comprehensive assessment on the above both levels

4. Dereplication

- Dual Strategy Approach
 - MS1-based dereplication using Custom Database Matching: Validates exact mass, halogen patterns, and isotope intensity similarity.
 - MS2-Based Dereplication by Integrating GNPS: MS2 molecular networking, halogenated compound annotation, graphML file enhancement.

Technical Advantages

- **High Throughput**: automatically process unlimited samples in several to dozens of seconds each on standard hardware (Core i9, 16GB RAM).
- Accuracy: >98.3% precision in halogen detection across simulated and experimental LC-MS datasets.
- **Integration with GNPS**: enhance molecular network annotation in the element dimension by embedding DeepHalo results into GNPS output graphML file
- **Efficient Dereplication**: significantly higher dereplication rate compared to molecular networking alone in GNPS

Installation

Prerequisites

• Python 3.10 (Verify with **python** --version).

Installation Methods

From PyPI

pip install DeepHalo

From Local Wheel

pip install path/to/DeepHalo-xxx.whl

From Source

git clone https://github.com/xieyying/DeepHalo.git cd DeepHalo pip install -e .

Quickstart

1. Detect Halogenated Compounds in mzML Files

halo detect -i /path/to/mzml_files -o /path/to/output_directory -ms2

2. Dereplication with GNPS and/or Custom Database

halo dereplicate -o /path/to/output_directory -g /path/to/GNPS_results -ud /path/to/custom_database.csv

Command-Line Usage

General Help

halo --help # List all commands

halo [command] --help # Show options for a specific command (e.g., 'halo detect --help')

Commands

1. Analyze mzML Files

halo detect -i <input path> -o <project path> [-c <config file>] [-b <blank samples dir>] [-ms2]

2. Dereplication

 $halo\ dereplicate\ -o\ <project_path>\ [-g\ <GNPS_folder>]\ [-ud\ <user_database.csv>]$

3. Create Training Dataset

halo create-ds project path > [-c <config file >]

4. Train Model

halo train <project path> [-c <config file>] [-m search]

Output Directory Structure

Instruction

The whole DeepHalo analysis process including 4 steps

Step 1: Halogenate Mining

Run the command:

```
halo detect -i <input_path> -o <project_path> [-c <config_file>] [-b <blank_samples_dir>] [-ob] [-ms2]
```

Parameters:

- -i INPUT_PATH (Required)
 - o Input .mzML file or directory
 - o Example (Linux): -i /home/data/ms files
 - o Example (Windows):-i D:\data\ms files
- -o OUTPUT DIR (Required)
 - Output directory path
 - o Example (Linux): -o /home/analysis/output
 - o Example (Windows): -o D:\analysis\output
- -c <config file> (Optional)
 - o Parameters for LCMS data analysis
 - o Example(Linux): -c /home/deephalo/config.toml
 - o Example (Windows): -c D:\deephalo\config.toml

- -b <black_samples_dir> (Optional)
 - Blank .mzML file or directory
 - o Example (Linux): -b /home/analysis/blank
 - o Example (Windows): -b D:\analysis\blank
 - o Run blank exclusion during analysis
 - Default: disabled
- -ob (Optional)
 - o Force regenerate blank feature detection results for blank exclusion
 - Default: disabled
- -ms2 (Optional)
 - o Enable MS2 data extraction
 - o Optional parameter (required for steps 2-4)
 - Default: disabled

Example Usage:

halo analyze-mzml -i D:\data\ms files -o D:\analysis\output -ms2

Output Files

- config.toml: Analysis parameters
- error files.txt: Failed mzML files
- halo\: MS1 information
- ms2 output\: MS2 data

Notes

- Results are filtered by H-score
- To disable filtering: Set H-score to 0 in config.toml file
- H-score visible in Cytoscape after dereplication
- For failed files:
 - 1. Retry analysis
 - 2. If error persists, reconvert raw data to mzML

Step 2: Molecular Networking analysis employing GNPS

- 1. Submit MS2 data in ms2 output to GNPS platform
- 2. Analyze molecular similarity using GNPS platform
- 3. Download and unzip results

Example GNPS output structure:

```
☐—Demo_GNPS_output

| METABOLOMICS-SNETS-V2-xxx-main.graphml
| params.xml
| ☐—clusterinfo
| ☐—clusterinfosummarygroup_attributes_withIDs
| ☐—gnps_molecular_network_graphml
| ☐—result_specnets_DB
```

Step 3: Dereplication

Run the command:

halo dereplicate -o PROJECT PATH [-g GNPS DIR] [-ud DATABASE FILE]

- -o PROJECT PATH (Required)
 - o Output directory (same as used in function of 'detect')
 - o Example: -o D:\analysis\output
- -g GNPS DIR (Optional)
 - o GNPS results directory containing .GraphML file
 - o Example: -g D:\analysis\Demo GNPS output
- -ud DATABASE FILE (Optional)
 - o Custom database file (CSV/JSON)
 - Example: -ud D:\databases\compounds.csv

Output Details

Upon completion of the analysis, the following files and directories will be generated:

1. 'dereplication' Directory

- o Contains processed MS1 data in CSV format.
- If a database was provided, compound match information will also be included in the csv file.

2. GraphML File (generated when GNPS results are provided)

- o The file name will have the suffix adding DeepHalo results.
- This file contains the mean H-score and classification predicted from centroid isotope patterns, along with any database match information if a dereplication database was used.

3. Processed Database File (if a database is provided)

- o The file name will include the suffix DeepHalo dereplication ready database.
- This file is formatted for future analyses, significantly reducing processing time in subsequent runs.

Example DATABASE FILE:

compound_name	formula	Smiles
Compound1	C6H5Cl	ClC1=CC=CC=C1
Compound2	C6H4Cl2	ClC1=CC=C(Cl)C=C1

Important Notes

1. File Format

o The input file must be in CSV format.

2. Required Columns

- o compound name
- o formula

3. Optional (but Recommended) Column

o Smiles (These aid in structure visualization).

4. Additional Requirements

- o Ensure that the formula column contains valid molecular formulas.
- o The file must be encoded in UTF-8.

Step 4: Visualization in Cytoscape

To visualize the results, open the GraphML file (with the suffix *_adding_DeepHalo_results.graphml) in Cytoscape. The file includes the following annotated variables:

• H scoreMean:

- o **Range:** [0, 1]
- O **Default Threshold:** 0.4, above which the presence of halogen is inferred.
- o **Interpretation:** Higher values indicate a greater likelihood of halogen presence.
- **Other Factors Affecting the Score:**
 - Sample complexity
 - Mass spectrometer resolution

• classification:

- Isotope pattern classifications (0-7):
 - 0: X-type (mixed/polyhalogenated)
 - 1: Br-type (Br/Cl₃)
 - 2: Cl-type (Cl/Cl₂)
 - 3: Se-type (Selenium compounds)
 - 4: B-type (Boron compounds)
 - 5: Fe-type (Iron compounds)

- 6: C-type (compounds containing only C, H, O, N, F, P, S, Na)
- 7: artifact-type (artifact isotope patterns)

• Inty_cosine_score:

Cosine similarity between experimental and theoretical isotope peak intensities. Higher values indicate a greater likelihood of a known compound

• Compound_names and Adducts:

These fields provide the known molecular names and adduct information sourced from a user-provided database, offering essential metadata for compound identification.

Smiles

Provides the known molecular structure, sourced from either the GNPS library and a user-provided database.

error_ppm

Represents the mass error between the measured m/z value of the test molecule and that of its corresponding known compound match.

Dependencies,

- pandas == 2.0.3
- numpy == 1.22.0
- tensorflow == 2.10.1
- scikit-learn == 1.3.1
- pyopenms == 3.1.0
- Full list: See <u>README.md</u>

License

Distributed under the MIT License.

For methodology details and benchmarks, refer to the <u>GitHub repository</u>.