MolDiscovery: Learning Mass Spectrometry Fragmentation of Small Molecules

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MolDiscovery: Learning Mass Spectrometry Fragmentation of Small Molecules

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Abstract

Identification of small molecules is a critical task in various areas of life science. Recent advances in mass spectrometry have enabled the collection of tandem mass spectra of small molecules from hundreds of thousands of environments. To identify which molecules are present in a sample, one can search mass spectra collected from the sample against millions of molecular structures in small molecule databases. This is a challenging task as currently it is not clear how small molecules are fragmented in mass spectrometry. The existing approaches use the domain knowledge from chemistry to predict fragmentation of molecules. However, these rule-based methods fail to explain many of the peaks in mass spectra of small molecules. Recently, spectral libraries with tens of thousands of labelled mass spectra of small molecules have emerged, paving the path for learning more accurate fragmentation models for mass spectral database search. We present molDiscovery, a mass spectral database search method that improves both efficiency and accuracy of small molecule identification by (i) utilizing an efficient algorithm to generate mass spectrometry fragmentations, and (ii) learning a probabilistic model to match small molecules with their mass spectra. We show our database search is an order of magnitude more efficient than the state-of-the-art methods, which enables searching against databases with millions of molecules. A search of over 8 million spectra from the Global Natural Product Social molecular networking infrastructure shows that our probabilistic model can correctly identify nearly six times more unique small molecules than previous methods. Moreover, by applying molDiscovery on microbial datasets with both mass spectral and genomics data we successfully discovered the novel biosynthetic gene clusters of three families of small molecules.

Availability: The command-line version of molDiscovery and its online web service through the GNPS infrastructure are available at https://github.com/mohimanilab/molDiscovery.

Introduction

A crucial problem in various areas of life science is to determine which known small molecules are present/absent in a specific sample. For example, physicians are devoted to discovering small molecule biomarkers in plasma/oral/urinal/fecal/tissue samples from a patient for disease diagnosis [1] and prognosis [2]. Epidemiologists are interested in identifying small molecule disease risk factors from diet [3] and environment [4]. Ecologists are interested in characterizing the molecules produced by microbes in various microbial communities [5]. Natural product scientists need to identify all

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the known molecules in their sample, clearing the path towards the discovery of novel antimicrobial or antitumor molecules [6, 7].

Recent advances in high-throughput mass spectrometry have enabled collection of billions of mass spectra from hundreds of thousands of host-oriented/environmental samples [8, 9, 10, 11]. A mass spectrum is the fingerprint of a small molecule, which can be represented by a set of mass peaks (Figure 1AB). In order to identify small molecules in a sample with tens of thousands of spectra, one can either (i) de novo predict small molecule structure corresponding to mass spectra, (ii) search these mass spectra against tens of thousands of reference spectra in spectral libraries, or (iii) search these mass spectra against millions of molecular structures in small molecule databases. De novo prediction can potentially identify both known and novel small molecules. However, it is rarely used in practice due to the intrinsic complexity of small molecule structure and the low signal-to-noise ratio of mass spectral data. Spectral library search is recognized as the most reliable mass spectral annotation method. Nevertheless, current reference spectral libraries are limited to tens of thousands of molecules, and the majority of known small molecules are not represented in any reference spectral library. Furthermore, collecting mass spectra of all known small molecules individually would be expensive and time-consuming. The most frequently used strategy for small molecule identification is in-silico search of small molecule structure databases. This approach enables small molecule identification in known databases, such as PubChem [12], dictionary of natural products (DNP) [13], and AntiMarin [14]. Moreover, in-silico database search also applies to discovery of novel small molecules through genome mining [15].

The majority of in silico database search methods are rule-based models that incorporate domain knowledge such as bond types, hydrogen rearrangement, and dissociation energy [16, 17, 18, 19, 20, 21] to predict fragmentations of small molecules and score small molecule-spectrum pairs. However, due to the complex rules involved in small molecule fragmentation, these methods are computationally inefficient for heavy small molecules. Moreover, these predictions, which are based on expert knowledge, fail to explain many of the peaks in mass spectra.

Recently, spectral libraries with tens of thousands of labelled mass spectra of small molecules (small molecule-spectrum matches) have emerged, paving the path for learning more accurate fragmentation models in order to improve sensitivity and specificity of mass spectral database search [10]. In this paper, we improve the efficiency and accuracy of small molecule identification by (i) designing an efficient algorithm to generate mass spectrometry fragmentations and (ii) developing a probabilistic model to identify small molecules from their mass spectra. Our results show that molDiscovery greatly increases the accuracy of small molecule identification, while making the search an order of magnitude more efficient. After searching 8 million tandem mass spectra from the Global Natural Product Social molecular networking infrastructure (GNPS) [10], molDiscovery identified 56,971 small molecules at 0% false discovery rate (FDR), an eight times increase compared to existing methods. On a subset of the GNPS repository with known genomes, molDiscovery correctly linked 20 known and three novel biosynthetic gene clusters to their molecular products.

Results

Outline of molDiscovery pipeline. The molDiscovery pipeline starts by (i) constructing metabolite graphs and (ii) generating fragmentation graphs. For the latter, molDiscovery uses a new efficient algorithm for finding bridges and 2-cuts in the metabolite graphs. Afterwards, molDiscovery proceeds with (iii) learning a probabilistic model for matching fragmentation graphs and mass spec-
molDiscovery training

\[ \text{Score} = S_1 + S_2 + S_3 + S_4 + S_5 \]

molDiscovery scoring

**Figure 1:** MolDiscovery Framework. Panels (A)-(F) show the training procedures of molDiscovery, while panels (G)-(K) are the scoring procedures based on the pretrained probabilistic model. (A) A reference molecule \( R \) in spectral library. (B) The reference spectrum of \( R \). (C) The fragmentation graph of \( R \). The root node represents the whole molecule, while the other nodes are fragments of it. The edges are annotated with the type of bonds where the fragmentation occurs. Here, a fragment will be annotated with red, blue, green or purple if it corresponds to a mass peak in reference spectrum with logRank = 1, 2, 3, 4 respectively (see the Methods section). (D) Annotation of the reference spectrum of \( R \). A mass peak will be annotated with a fragment if its mass is identical to the mass of the fragment plus charge, within a tolerance. (E) A table counting the number of fragments observed in training data with specific bondType and logRank, referred to as count matrix. In this example, since the number of peaks is small, all the present peaks have logRanks between 1 and 6, and the absent ones are shown with the lowest possible logRank = 7. Note that we allow logRanks between 1 and 64. The null column corresponds to the experimental peaks which cannot be explained by the fragmentation graph. (F) The probabilistic model \( P(\text{logRank} | \text{bondType}) \), which is computed by normalizing the count matrix. (G) A molecule \( Q \) in a chemical structure database. (H) A query spectrum. (I) The fragmentation graph of \( Q \). (J) Annotation of the query spectrum with the fragmentation graph of \( Q \). (K) Computation of the query spectrum match score, which is the sum of scores of all the fragments in the annotated fragmentation graph. Here we represent \( P(\text{logRank} = i | \text{bondType}) \) by \( P(i | \text{bondType}) \). For simplicity, logRank\textsubscript{pca}, 2-cuts columns, and rows of logRank = 5, 6 are not shown.
tra (Figure 1A-E), (iv) scoring small molecule-spectrum pairs (Figure 1F-K), and (v) computing FDR.

In the past we introduced Dereplicator+ [22], a database search method for identification of small molecules from their mass spectra, that follows a similar series of steps to search mass spectra against chemical structures. However, Dereplicator+ uses a brute-force method for fragmentation graph construction, and the naive shared-peak-count for scoring.

Datasets. The GNPS spectral library contains 8,964 small molecule-spectrum matches [10]. After removing spectra corresponding to duplicate molecules, we obtain 4,781 non-redundant spectra, which are used as training and testing data for molDiscovery. The MassBank of North America (MoNA) contains 86,209 positive mode LC-MS/MS spectra. After filtering for unique spectra using SPLASH [23] and isolating singly-charged molecule-spectrum matches, spectra from 3,278 unique molecules are extracted (determined using RDKit [24] canonicalized SMILES). These two spectral libraries are searched against the combined natural product structure database (CombinedDB) of DNP (77,057 non-redundant molecules) [13] and AntiMarin (60,908 molecules) [14].

Furthermore, we benchmark molDiscovery against Dereplicator+ [22] using 46 high-resolution GNPS spectral datasets (Supplementary Table S1) collected from various environments (about 8 million mass spectra in total). These spectra are searched against a combined structure database (AllDB) of 719,958 compounds from DNP, AntiMarin, UNPD [25], HMDB [26], LMSD [27], FooDB [28], NPAtlas [29], KEGG [30], DrugBank [31], StreptomeDB [32], GNPS spectral library, MIBiG [33], PhenolDB [34] and Quorumpeps [35] databases.

Figure 2: Heatmap of $P(\log\text{Rank}|\text{bondType, logRank}_{pa})$. Each row represents bond type bondType and parent’s intensity rank logRank_{pa} of a fragment. The row ”null” refers to the null distribution $P(\log\text{Rank}|null)$. Each column represents the logRank of a child fragment. When logRank_{pa} is 0, it means the parent is the root (precursor molecule). (A) logRank distribution over different bondTypes for logRank_{pa} = 3. (B) logRank distribution of OC bond over different logRank_{pa}. Since there is only one fragment in the fragmentation graph that could have logRank = 1, the parent fragment of such fragment can not generate another fragment with logRank = 1, hence $P(1|NC, 1) = 0$. (C) logRank distribution over different bondTypes for logRank_{pa} = 7. Supplementary Figure S1-2 shows the complete heatmaps for charge +1 and +2 respectively.
Preferential fragmentation patterns in mass spectrometry. MolDiscovery learned a probabilistic model (see Methods) that reveals the preferences in mass spectrometry fragmentation. First, mass spectrometry fragmentation has preference for bond type $bondType$. Figure 2A shows that the breakage of $bondTypes$ NC and OC lead to fragments with higher rank (i.e. logRanks closer to 1) than CC bond, indicating that they are more likely to be broken by mass spectrometry. In addition, bridges (NC, OC and CC) tend to generate fragments with higher logRank than 2-cuts (for example, NC_NC, NC_OC, CC_OC), suggesting bridges are more likely to be fragmented than 2-cuts. Note that the logRank distribution of fragments due to the 2-cut CC_CC is similar to the null model distribution (random noise), which implies that in practice the fragmentation of CC_CC 2-cut rarely happens.

In addition, parent fragments with high ranks are more likely to produce high-rank children fragments, while fragments with low ranks are similar to random noise. For example, for fragments produced by the breakage of OC bond (Figure 2B), if their parents have high ranks (1, 2 or 3), they are likely to be higher rank than those with lower-rank parents (6, 7). Moreover, molDiscovery automatically learns to discard fragments with low rank parents which are not informative for predicting mass spectrometry fragmentations. The distribution of logRank for these fragments is similar to noise (null distribution, Figure 2C). Therefore, the log-likelihood scores for such fragments are close to zero.

Benchmarking the efficiency of fragmentation graph construction algorithms. We compared the performance of the fragmentation graph construction algorithms in molDiscovery and Dereplicator+. For both algorithms, we allow for at most two bridges and one 2-cut. The algorithms are benchmarked on CombinedDB. Figure 3A and Supplementary Figure S3A show the

![Figure 3](image-url)
average fragmentation graph construction time for small molecules in different mass ranges. The results show that the average fragmentation graph construction time for Dereplicator+ grows exponentially as the mass of small molecules increase, while the construction time only grows linearly for molDiscovery. When the mass is greater than 600 Da, molDiscovery is two orders of magnitude faster than Dereplicator+.

Figure 3B and Supplementary Figure 3B compare the maximum memory consumption of molDiscovery with Dereplicator+. For both chemical structure databases, Dereplicator+ only works for molecules with masses less than 1000 Da, while molDiscovery can handle molecules with masses up to 2000 Da.

**Benchmarking database search accuracy of molDiscovery.** The accuracy of database search by molDiscovery is compared to four state-of-the-art methods on the non-redundant GNPS spectral library. First we split the spectra into five batches, each with 956 and one with 957 tandem mass spectra. In order to evaluate their performance on larger chemical structure databases, we combined the molecules in the non-redundant GNPS spectral library with non-redundant DNP (77,057 molecules). Then we performed 5-fold cross validation for molDiscovery against this combined database of chemical structures. The database search results show that molDiscovery on average can correctly identify 74.2% small molecules in the test dataset as top ranked identifications in comparison to 63.6% for Dereplicator+, and lower rates for MAGMa+ [37], and CFM-ID [38] (Figure 4A). CSI:FingerID [39] achieves an average top ranked identification rate of 73.5%. However, of the 4,781 molecules tested 3,042 molecules appear in the CSI:FingerID positive mode training dataset. If we only evaluate the 1,739 compounds that are not present in the CSI:FingerID training dataset, molDiscovery achieves 73.9% and 92.2% accuracy for top 1 and top 10 identifications.

![Figure 4](image.png)

**Figure 4:** Benchmarking the database search accuracy on (A) GNPS spectral library (4,781 molecules) and (B) GNPS spectral library excluding CSI:FingerID training data (1,739 molecules). For each spectrum, we rank the matching scores given by molDiscovery, Dereplicator+, MAGMa+ 1.0.1, CFM-ID 2.0, and CSI:FingerID 1.4.3, (from SIRIUS 4.4.2 [36]). If there is a tie, we assign the average rank to all the matches in the tie. Then we check if the correct molecule-spectrum match is among the top $k$ ($k = 1, 3, 5, 10$) matches. The x-axis is the rank $k$, and the y-axis is the mean proportion of correct molecule-spectrum matches in the test datasets that are among the top $k$ matches.
| Software         | Runtime (d:hh:mm) | Memory Usage (GB) |
|------------------|-------------------|-------------------|
| molDiscovery     | 0:00:28           | 0.22              |
| Dereplicator+    | 0:03:23           | 25.46             |
| CFM-ID           | 8:00:44           | 0.041             |
| CSI:FingerID     | 7:07:26           | 3.13              |
| MAGMa+           | 3:14:27           | 1.87              |

Table 1: Runtime and memory usage of various in silico search tools. Reported performance is for a search of 4,781 spectra against DNP+GNPS (81,838 compounds). If supported by the software the chemical database was pre-processed before the search was run.

respectively, while CSI:FingerID achieves 66.1% and 71.0% accuracy (Figure 4B). Supplementary Figure 4 shows the top-k accuracies of the methods in the five cross validation batches.

MolDiscovery was also the most efficient method and used only 217 MB of memory as its peak (Table 1) making it seven times faster and two orders of magnitude more memory efficient than Dereplicator+. MolDiscovery was three orders of magnitude faster than CSI:FingerID, CFM-ID, and MAGMa+.

Evaluating molDiscovery sensitivity to platform variation. In order to determine if the probabilistic scoring method described is robust to changes in mass spectrometry platform, a subset of annotated spectra from MoNA dataset including 1,639 HCD-fragmented and 1,639 CID-fragmented spectra were used for evaluation. Two models were trained using the GNPS spectra augmented with either the HCD or CID fragmented MoNA dataset, and then tested on both fragmentation modes. Searches were run against a combined chemical structure database consisting of 3,278 MoNA, 77,057 non-redundant DNP, and 4,781 GNPS molecules. It was shown that the platform on which training data was collected did not affect performance of molDiscovery (Figure 5).

![Figure 5: Accuracy of molDiscovery on tandem mass spectra of different fragmentation modes. MolDiscovery models were trained on the GNPS spectra augmented with MoNA spectra labeled with either CID or HCD fragmentation modes. The two models are then tested on both CID and HCD data. Matches were ranked according to score. The fraction of spectra for which the correct molecule was in the top k (k = 1, 3, 5, 10) matches are shown along the y-axis.](image-url)
Figure 6: Accuracy of Dereplicator+ and molDiscovery on 163 doubly-charged spectra from MoNA. The molDiscovery model was initially trained on 4,781 singly-charged spectra from GNPS and further trained on 813 spectra identified as doubly-charged spectra by Dereplicator+.

Performance on doubly-charged spectra. Only 163 spectra from MoNA dataset are annotated as doubly-charged. In light of the small amount of data we reserved the MoNA dataset as our test dataset, and used high confidence doubly-charged Dereplicator+ identifications of the GNPS datasets as the training data (813 spectra and 180 unique molecules at cutoff score of 15). Since this training data is still small, we bootstrapped doubly-charged parameters using the singly-charged model we previously trained. The training process updates the model parameters according to the doubly-charged training data. Despite the small training dataset, molDiscovery outperformed Dereplicator+ on search of 163 doubly-charged spectra against non-redundant DNP and GNPS chemical database augmented with MoNA molecules (Figure 6).

MolDiscovery performance on different compound classes and mass ranges. To evaluate how well molDiscovery performs on different compound classes, we used ClassyFire [40] to annotate the non-redundant GNPS spectral library. Figure 7A shows that molDiscovery performs as well as or better than all the other tools on 15 out of 17 superclasses. The two superclasses for which molDiscovery has lower accuracy than other methods, Homogeneous non-metal compounds and Hydrocarbon derivatives, had only five and two annotated compounds respectively (Supplementary Figure S5A). Similarly, molDiscovery performs equally or better than the competing methods on compounds from nearly all mass ranges (Figure 7B). It is worth noting that the accuracy of CSI:FingerID drops a lot on the compounds that are not in its training data (Supplementary Figure S6).

Performance on large-scale spectral datasets. We benchmarked the performance of molDiscovery against Dereplicator+ on 45 GNPS datasets containing a total of 8 million tandem mass spectra (Supplementary Table S1). In contrast to the GNPS spectral library where all the spectra are annotated and their molecules are known, the spectra in the GNPS spectral datasets are not annotated, and they could correspond to novel or known molecules. Figure 8 shows the numbers of small molecule-spectrum matches and unique compounds from the AllDB database identified by molDiscovery and Dereplicator+ at various FDR levels. At the strict 0% FDR level, molDiscov-
ery identified eight times more spectra (56971 versus 7684) and six times more unique compounds (3185 versus 542) than Dereplicator+. MolDiscovery search took 34 days on 10 threads, and CSI-FingerID, CFM-ID and MAGMa+ are not benchmarked, as the searches would have taken hundreds of years. The top 100 identifications by molDiscovery is shown in Supplementary Table S2.

Validating molDiscovery identifications using a literature search. We benchmarked molDiscovery against Dereplicator+ on top 100 identifications from GNPS datasets by performing literature search. For this task, we used the extensively studied GNPS dataset MSV000079450 (∼400,000 spectra from Pseudomonas isolates) [41, 42]. Out of the top 100 small molecule-spectra matches reported by molDiscovery, 78 correspond to compounds having Pseudomonas origin based on taxonomies reported for molecules in AntiMarin database. The second largest genus among the identifications (20 out of 100) is Bacillus. The molecule-spectrum matches from Bacillus are likely to be true positives as the dataset is known to be contaminated with Bacillus species [42]. While the top 100 identifications from Dereplicator+ also contains 20 Bacillus matches, the number of hits related to Pseudomonas species is 62, 25% lower than molDiscovery. 18 identifications of Dereplicator+ are annotated as having fungi origin, which are likely false positives.

MolDiscovery links small molecules to their biosynthetic gene clusters (BGCs). We cross-validated molDiscovery and genome mining by searching 46 microbial datasets (922 unique microbial strains, 8,013,433 spectra) that contained both tandem mass spectrometry data and genomic data. MolDiscovery successfully identified 20 molecules in microbial isolates that contained their known BGCs (Table 2.)
Figure 8: Performance of molDiscovery and Dereplicator+ on GNPS spectral datasets. The curves show the number of (A) small molecule-spectrum matches and (B) unique compounds identified by molDiscovery and Dereplicator+ in the search of 8 million spectra from 46 GNPS spectral datasets against 719,958 compounds of AllDB at different FDR levels.

Table 2: MolDiscovery identified 20 molecules in bacterial isolates that contained their known BGCs. A genome is reported to contain a known BGC from the MiBiG database [33] if the genome has total BLAST hit [43] to the BGC with at least 90% coverage and 70% identity. Taxa similarity means similarity between the taxon of source organism and the organism reported in MiBIG. PK stands for polyketides, MA stands for N-myristoyl-D-asparagine, OT stands for (R)-N1-((S)-5-oxohexan-2-yl)-2-tetradecanamidosuccinamide, and TA stands for cis-7-tetradecenoyl-D-asparagine.

Moreover, molDiscovery successfully discovered novel BGCs for three small molecule families from streptomyces dataset MSV000083738 [44, 45]. MolDiscovery search results for this dataset are available at https://github.com/mohimanilab/molDiscovery. MolDiscovery identified dinghupetins A-D [46] in Streptomyces sp. NRRL B-5680 and several other Streptomyces strains (Supplementary Table S2). AntiSMASH [47] revealed a non-ribosomal peptide (NRP) biosynthetic gene cluster in the genome of this strain, with seven adenylation domains that are highly specific to the amino acid residues of dinghupetin molecules [48] (Supplementary Figure S7).

MolDiscovery detected lipopeptin family and neopeptin family molecules in Streptomyces hygroscopicus NRRL B-1477 and Streptomyces rimosus NRRL WC3874. Genome mining revealed...
Figure 9: (A) Biosynthetic gene cluster of longicatenamycin family. MolDiscovery identified longicatenamycin variants at FDR 0% in *Streptomyces rimosus* NRRL B-8076. After searching its genome using antiSMASH, we detected an NRP BGC with adenylation domains which show high specificity to the amino acid residues of longicatenamycin. (B) Molecular structures of the two longicatenamycin variants identified by molDiscovery at FDR 0%.

an NRP BGC in *S. hygroscopicus* NRRL B-1477 with seven adenylation domains, which are highly specific to the residues of neopeptin and lipopeptin (Supplementary Figure S8). It has been reported that these two families are structurally similar [49]. Thus, this BGC could be responsible for the production of both families.

In addition, molDiscovery identified longicatenamycin family of cyclic hexapeptides in *Streptomyces rimosus* NRRL B-8076 and several other *Streptomyces* strains. Genome mining of the NRRL B-8076 strain revealed a BGC highly specific to longicatenamycin (Figure 9). MolDiscovery also identified cyclic hexapeptide families microphorusrumamides (A and B) and noursamycin (A, D and E) in related *Streptomyces* strains (Supplementary Table S2), suggesting the same BGC might be responsible for their production. Dereplicator+ and CSI-FingerID fail to discover longicatenamycin at 1% FDR threshold.
Discussion

With the advent of high-throughput mass spectrometry, large-scale tandem mass spectral datasets from various environmental/clinical sources have become available. One of the approaches for annotating these datasets is in silico database search. Currently the existing methods for in silico database search of tandem mass spectra are rule-based. These methods can not scale to searching molecules heavier than 1000 Da and fail to explain many of the fragment peaks in tandem mass spectra.

MolDiscovery introduces an efficient algorithm to construct fragmentation graphs of small molecules, enabling in silico database search for molecules up to 2000 Da. MolDiscovery is more efficient than the state-of-the-art methods in terms of run-time and memory consumption. Furthermore, by training on the reference spectra from the GNPS spectral library, MolDiscovery learned a probabilistic model that significantly improves the accuracy of database search. MolDiscovery is shown to be robust against dissociation technique, and it performs well on singly and doubly charged spectra.

Small molecule fragmentation is a complex process that depends not only on the type of fragmented bonds, but also on local/global features of small molecules such as moiety. Currently, molDiscovery ignores all these features. Recent advances in graph-based machine learning has enabled representing complex small molecule structures with continuous vectors, making it feasible to incorporate local/global structural information into the prediction of fragmentation, potentially leading to more accurate fragmentation models. MolDiscovery paves the way for these more sophisticated approaches by collecting larger training data of small molecule-spectrum matches through the search of millions of spectra.

Methods

Constructing fragmentation graphs of small molecules. MolDiscovery first constructs a metabolite graph for a small molecule structure and then generates a fragmentation graph from the metabolite graph (Figure 10). To simplify the modelling of small molecule fragmentation, we assume that mass spectrometry can only break N-C, O-C and C-C bonds. This is a reasonable assumption, as among top ten most frequent bonds in AntiMarin, these three bonds are the only single bonds that do not contain a hydrogen atom (Supplementary Table S3).

To construct a metabolite graph, molDiscovery first disconnects N-C, O-C and C-C bonds. The resulting connected components form the nodes of the metabolite graph. Edges in the metabolite graph correspond to bonds between the connected components (Figure 10AB).

The fragmentation graph of a molecule is a directed acyclic graph with a single source (the metabolite graph) where nodes are fragments of the molecule and directed edges between the nodes represent bridge or 2-cut fragmentations. To construct a fragmentation graph, molDiscovery first searches for all the bridges and 2-cuts of the metabolite graph to obtain depth one fragments of the molecule using Hopcroft and Tarjan’s algorithm for graph manipulation [50]. Each fragment of the molecule can be represented by a fixed length binary vector that indicates the presence or absence of metabolite graph nodes in the fragment (Figure 10D). We observe that depth two fragments can be formed by a bitwise AND operation between their parent fragments and depth one fragments (Figure 10E). This generalizes to computing fragments at depth \( n > 1 \) by intersecting their parents (fragments at depth \( n - 1 \)) with fragments at depth one. The final fragmentation graph is constructed
Learn a probabilistic model to match mass spectra and small molecules. Dereplicator+ uses a naïve scoring scheme which ignores that (i) fragmentation probability of different bonds are different, (ii) bridges have a higher chance of fragmentation than 2-cuts, (iii) peaks with higher intensity have higher chances of corresponding to a fragmentation than lower intensity peaks, (iv) matches are biased towards molecules with large fragmentation graph size, and (v) matches are biased towards spectra with a large number of peaks.

In order to solve the above shortcomings we develop a probabilistic model for matching spectra with small molecules. Given a molecule-spectrum pair in the training data (Figure 1AB), we first construct the fragmentation graph of the molecule using our fragmentation graph construction algorithm (Figure 1C). Each fragment in the fragmentation graph is assigned a \((bondType, logRank)\) label. \(bondType\) represents the bond(s) disconnected in the parent fragment to produce the current fragment. It can either be one bond (bridge) or two bonds (2-cut). Bridges can be OC, NC, CC, while 2-cuts can be their pairwise combinations.

\(logRank\) represents the intensity of the mass peak corresponding to the fragment (Figure 1CD). The mass spectrometry community has used intensity rank as an abundance measure of mass peaks in a spectrum [51, 52]. The higher the intensity rank is (closer to rank 1), the more abundant is
the corresponding fragment. To reduce the number of parameters and avoid overfitting, we group peaks according to their logRank instead of directly using intensity rank (Supplementary Note S1). A fragment will be annotated with a logRank between 1 – 6 if there is a peak with rank between 1 – 64 in the spectrum within 0.01 Da of the mass of the fragment (Figure 1CD). If there is no such mass peak, the fragment will be annotated with logRank = 7.

In the annotated fragmentation graph (Figure 1C) we assume that (i) the logRank of each fragment depends only on its bondType and the logRank of its parent. Here, we only consider direct parent, as considering grandparents of the fragments increases the number of parameters by an order of magnitude, resulting in overfitting. We further assume (ii) logRank of each mass peak is independent from the logRank of other peaks. While this assumption is sometimes wrong, e.g. only one peak can have logRank 1, we use this assumption to simplify our probabilistic model. Finally, we assume (iii) the root node has logRank 0. Given a small molecule structure Molecule and its fragmentation graph FG, the probability of generating a spectrum Spectrum is as follows:

\[
P(Spectrum|Molecule) = \prod_{peak\in Spectrum} P(peak|FG) = \prod_{peak\in Spectrum_{FG}} P(peak|FG) \prod_{peak\in Spectrum_{null}} P(peak|FG)
\]

where Spectrum_{FG} represents all the peaks within 0.01 Da of some fragment in FG, and Spectrum_{null} represents the rest of the peaks. If multiple fragments match the same peak, we randomly pick one with lowest depth. Since we use logRank as a measure of abundance and by definition all peaks in Spectrum_{FG} correspond to a fragment in FG, we can rewrite the equation as follows:

\[
P(Spectrum|Molecule) = \prod_{frag\in FG} P(logRank_{frag}|FG) \prod_{peak\in Spectrum_{null}} P(peak|FG)
\]

where \(frag\) is a fragment in the fragmentation graph. Then, by assumption (i):

\[
P(Spectrum|Molecule) = \prod_{frag\in FG} P(logRank_{frag}|bondType_{frag}, logRank_{pa(frag)}) \prod_{peak\in Spectrum_{null}} P(peak|FG)
\]

where \(pa(frag)\) is the parent of \(frag\). Similarly, we can obtain the probability of generating a random spectrum (null model):

\[
P(Spectrum|null) = \prod_{peak\in Spectrum} P(peak|null)
\]

In order to learn \(P(logRank_{frag}|bondType, logRank_{pa(frag)})\), we can directly count the number of the fragments with a particular logRank for each (bondType = \(b\), logRank_{pa} = \(p\)) combination in the training data as follows:

\[
P(logRank = l|bondType = b, logRank_{pa} = p) = \frac{\sum_i |\{frag \in FG_i|logRank_{frag} = l, bondType_{frag} = b, logRank_{pa(frag)} = p\}|}{\sum_i |\{frag \in FG_i|bondType_{frag} = b, logRank_{pa(frag)} = p\}|}
\]

where \(FG_i\) is the fragmentation graph of the \(i^{th}\) molecule-spectrum pair in the training data.
Similarly, for the null model, we can compute the logRank distribution of all the mass peaks as follows:

\[
P(\text{logRank} = l | \text{null}) = \frac{\sum_i |\{\text{peak} \in \text{Spectrum}_i | \text{logRank}_{\text{peak}} = l\}|}{\sum_i |\{\text{peak} \in \text{Spectrum}_i\}|}
\]

where \(\text{Spectrum}_i\) is the spectrum of the \(i^{th}\) molecule-spectrum pair.

**Scoring a spectrum against a small molecule.** Given a query tandem mass spectrum (Figure 1H) and a small molecule in a chemical structure database (Figure 1G), if the spectrum precursor mass is within 0.02 Dalton of the mass of the small molecule and the user-specified adduct, they will become a candidate pair for scoring. We then construct the fragmentation graph of the molecule and annotate the query spectrum against the fragmentation graph (Figure 1IJ). Based on the described probabilistic model, we use a log-likelihood ratio model (Figure 1K) to score the spectrum against the small molecule:

\[
\log \frac{P(\text{Spectrum}|\text{Molecule})}{P(\text{Spectrum}|\text{null})} = \log \prod_{\text{peak} \in \text{Spectrum}_{\text{FG}}} \frac{P(\text{peak}|\text{FG})}{P(\text{peak}|\text{null})} \prod_{\text{peak} \in \text{Spectrum}_{\text{null}}} \frac{P(\text{peak}|\text{FG})}{P(\text{peak}|\text{null})}
\]

\[
= \log \prod_{\text{frag} \in \text{FG}} \frac{P(\text{logRank}_{\text{frag}}|\text{FG})}{P(\text{logRank}_{\text{frag}}|\text{null})}
\]

\[
= \log \prod_{\text{frag} \in \text{FG}} \frac{P(\text{logRank}_{\text{frag}}|\text{bondType}_{\text{frag}}, \text{logRank}_{\text{pa(\text{frag})}})}{P(\text{logRank}_{\text{frag}}|\text{null})}
\]

\[
= \sum_{\text{frag} \in \text{FG}} \log \frac{P(\text{logRank}_{\text{frag}}|\text{bondType}_{\text{frag}}, \text{logRank}_{\text{pa(\text{frag})}})}{P(\text{logRank}_{\text{frag}}|\text{null})}
\]

Note that in (9), we only need to consider the mass peaks corresponding to fragments in the fragmentation graph since we assume all the other peaks are generated by random chance, hence \(\frac{P(\text{peak}|\text{FG})}{P(\text{peak}|\text{null})} = 1\).

**Computing FDR.** We use target-decoy analysis to estimate FDR [53]. First, we randomly shuffle the fragmentation graph of target molecules to create decoy fragmentation graphs. Then, tandem mass spectra are searched against the target and the decoy fragmentation graph databases respectively. At each score cutoff, there are \(N_{\text{target}}\) matches to the target database and \(N_{\text{decoy}}\) matches to the decoy database, and the FDR is estimated as

\[
FDR = \frac{N_{\text{decoy}}}{N_{\text{target}}}
\]

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Figures

molDiscovery training

(A) A reference molecule R in spectral library. (B) The reference spectrum of R. (C) The fragmentation graph of R. The root node

molDiscovery scoring

(G) A reference molecule R in spectral library. (B) The reference spectrum of R. (C) The fragmentation graph of R. The root node

Score = \( S_1 + S_2 + S_3 + S_4 + S_5 \)

Figure 1

MolDiscovery Framework. Panels (A)-(F) show the training procedures of molDiscovery, while panels (G)-(K) are the scoring procedures based on the pretrained probabilistic model. (A) A reference molecule R in spectral library. (B) The reference spectrum of R. (C) The fragmentation graph of R. The root node
represents the whole molecule, while the other nodes are fragments of it. The edges are annotated with the type of bonds where the fragmentation occur. Here, a fragment will be annotated with red, blue, green or purple if it corresponds to a mass peak in reference spectrum with logRank = 1, 2, 3, 4 respectively (see the Methods section). (D) Annotation of the reference spectrum of R. A mass peak will be annotated with a fragment if its mass is identical to the mass of the fragment plus charge, within a tolerance. (E) A table counting the number of fragments observed in training data with specific bondType and logRank, referred to as count matrix. In this example, since the number of peaks is small, all the present peaks have logRanks 1 to 4, and the absent ones are shown with the lowest possible logRank = 7. Note that we allow logRanks between 1 – 6, corresponding to peaks with ranks between 1 – 64. The null column corresponds to the experimental peaks which cannot be explained by the fragmentation graph. (F) The probabilistic model P(logRank|bondType), which is computed by normalizing the count matrix. (G) A molecule Q in a chemical structure database. (H) A query spectrum. (I) The fragmentation graph of Q. (J) Annotation of the query spectrum with the fragmentation graph of Q. (K) Computation of the query spectrum match score, which is the sum of scores of all the fragments in the annotated fragmentation graph. Here we represent P(logRank = i|bondType) by P(i|bondType). For simplicity, logRankpa, 2-cuts columns, and rows of logRank = 5, 6 are not shown.
**Figure 2**

Heatmap of $P(\text{logRank}|\text{bondType}, \text{logRankpa})$. Each row represents bond type bondType and parent's intensity rank logRankpa of a fragment. The row "null" refers to the null distribution $P(\text{logRank}|\text{null})$. Each column represents the logRank of a child fragment. When logRankpa is 0, it means the parent is the root (precursor molecule). (A) logRank distribution over different bondTypes for logRankpa = 3. (B) logRank distribution of OC bond over different logRankpa. Since there is only one fragment in the fragmentation graph that could have logRank = 1, the parent fragment of such fragment can not generate another fragment with logRank = 1, hence $P(1|\text{NC, 1}) = 0$. (C) logRank distribution over different bondTypes for logRankpa = 7. Supplementary Figure S1-2 shows the complete heatmaps for charge +1 and +2 respectively.

![Figure 2](image)

**Figure 3**

Benchmarking the efficiency of fragmentation graph construction algorithms. We compare (A) the average fragmentation graph construction time and (B) maximum memory consumption of molDiscovery with Dereplicator+ on the AntiMarin database over molecules of various masses. Note that we randomly selected 200 molecules from the AntiMarin database for each 200 Da mass window to compute average running time per molecule as well as maximum memory usage. For cases exceeding the memory limitation of 15 GB, no data point is shown.

![Figure 3](image)
Benchmarking the database search accuracy on (A) GNPS spectral library (4,781 molecules) and (B) GNPS spectral library excluding CSI:FingerID training data (1,739 molecules). For each spectrum, we rank the matching scores given by molDiscovery, Dereplicator+, MAGMa+ 1.0.1, CFM-ID 2.0, and CSI:FingerID 1.4.3, (from SIRIUS 4.4.2 [36]). If there is a tie, we assign the average rank to all the matches in the tie. Then we check if the correct molecule-spectrum match is among the top k (k = 1, 3, 5, 10) matches. The x-axis is the rank k, and the y-axis is the mean proportion of correct molecule-spectrum matches in the test datasets that are among the top k matches.
Figure 5

Accuracy of molDiscovery on tandem mass spectra of different fragmentation modes. MolDiscovery models were trained on the GNPS spectra augmented with MoNA spectra labeled with either CID or HCD fragmentation modes. The two models are then tested on both CID and HCD data. Matches were ranked according to score. The fraction of spectra for which the correct molecule was in the top k (k = 1, 3, 5, 10) matches are shown along the y-axis.
Figure 6

Accuracy of Dereplicator+ and molDiscovery on 163 doubly-charged spectra from MoNA. The molDiscovery model was initially trained on 4,781 singly-charged spectra from GNPS and further trained on 813 spectra identified as doubly charged spectra by Dereplicator+. 
Figure 7

Database search accuracy of molDiscovery, Dereplicator+, CSI:FingerID, MAGMa+, and CFM-ID on molecules of (A) different superclasses from ClassyFire [40] and (B) different mass ranges on GNPS spectral library compounds excluding CSI:FingerID training data.
Figure 8

Performance of molDiscovery and Dereplicator+ on GNPS spectral datasets. The curves show the number of (A) small molecule-spectrum matches and (B) unique compounds identified by molDiscovery and Dereplicator+ in the search of 8 million spectra from 46 GNPS spectral datasets against 719,958 compounds of AllDB at different FDR levels.
Figure 9

(A) Biosynthetic gene cluster of longicatenamycin family. MolDiscovery identified longicatenamycin variants at FDR 0% in Streptomyces rimosus NRRL B-8076. After searching its genome using antiSMASH, we detected an NRP BGC with adenylation domains which show high specificity to the amino acids residues of longicatenamycin. (B) Molecular structures of the two longicatenamycin variants identified by molDiscovery at FDR 0%.
Figure 10

Overview of fragmentation graph construction algorithm. (A) A small molecule. (B) Metabolite graph of the molecule. (C) Indexed metabolite graph. (D) Construction of the fragmentation graph of the small molecule. We assign an index to each component of a molecule and represent each of its fragments as a binary vector, where 1 and 0 indicate presence and absence of the components. Each bridge/2-cut of a parent fragment gives two child fragments. (E) Higher depth fragments can be computed as overlaps of their parents with the depth one fragments.

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- molDiscoverySupplementary08282020.pdf