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# Probabilistic framework for integration of mass spectrum and retention time information in small molecule identification



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## 1. Small Molecule Identification in Untargeted Metabolomics

- Challenge in untargeted metabolomics studies: Identification of the small molecules present in a biological sample
- LC-MS<sup>2</sup> widely used analysis platform: Liquid chromatography (LC) coupled with tandem mass spectrometry (MS<sup>2</sup>) (Fig. 1)
- Most machine learning approaches for small molecule identification only utilize MS<sup>2</sup> information [4, 3]
- LC retention time (RT) information can aid small molecule identification [6, 8]
- Challenges utilizing RT information: (1) LC-system specific RTs and (2) public RT databases are limited in size and molecule coverage

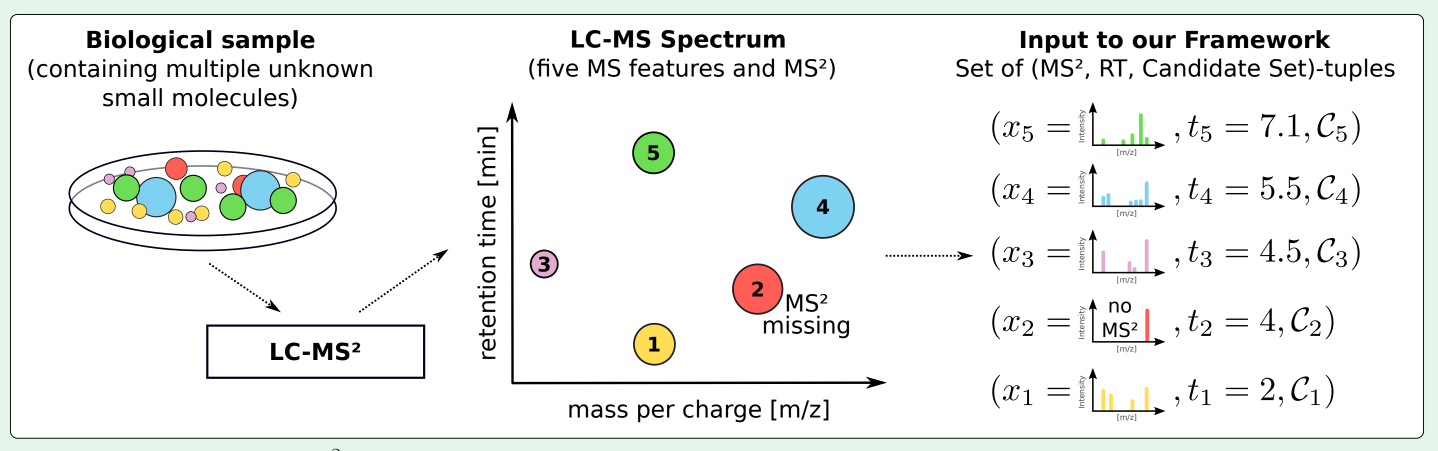


Fig. 1: LC-MS $^2$  analysis pipeline and resulting data used as input for our framework.

### 2. Retention Time (RT) Utilization for Small Molecule Identification

#### Comparison of measured RTs with (in-house) reference RTs

- References databases (DB) typically small and LC-system specific
- RT mapping between LC-systems possible, but requires DB overlaps [8]
- 2) Comparison of measured RTs with predicted ones [1]
- RT prediction models allow to get RTs for practically every molecular structure
- Majority of prediction models trained for a single LC-system only
- Compare RT-proxy or retention order with observed RT information [6, 2]
- Allows for predictions within an whole LC-system family, e.g. reversed-phase
- Retention orders largely preserved across LC-setups and prediction possible

#### Our proposed approach:

- Exploitation of all pairwise observed retention orders in an LC-MS<sup>2</sup> datasets
- Comparison of observed and predicted retention orders to down- and up-vote potential molecular structures of the unknown molecules.

# 3. LC-MS<sup>2</sup> Experiment Data: Input and Output of our Framework

• **Input:** Preprocessed LC-MS<sup>2</sup> data, i.e. after peak-picking and alignment (Fig.1):

$$\mathcal{D} = \{(x_i, t_i, \mathcal{C}_i)\}_{i=1}^N$$

- $x_i$ : MS Information; MS<sup>2</sup>, or MS<sup>1</sup> (precursor m/z) if no fragmentation available
- $t_i$ : Measured RT
- $\mathcal{C}_i$ : Molecular candidate sets, e.g. molecular structures found in PubChem by exact mass search
- N: Number of MS features
- Precomputed MS scoring assumed: MS<sup>2</sup> scores, e.g. by CSI:FingerID [4], MetFrag [6] or IOKR [3], or deviation of candidate and precursor mass for MS<sup>1</sup>
- Output: Ranking of the molecular candidates in  $m_{ir} \in \mathcal{C}_i$  for each MS feature i
- Ranking based on MS and RT information

#### 4. Our Probabilistic Framework to integrate MS and RT Information

- Graphical model G superimposed on the LC-MS<sup>2</sup> data (Fig. 2)
- Let G = (V, E) be complete graph with a **node**  $i \in V$  for each MS feature, and an **edge**  $(i, j) \in E$  for each feature pair
- Discrete random variable  $z_i \in \mathcal{Z}_i = \{1, \dots, n_i\}$  associated with each node  $(n_i = |\mathcal{C}_i|)$
- Candidate assignment for the complete data  $\mathbf{z} = \{z_i \mid i \in V\} \in \mathcal{Z}_1 \times \ldots \times \mathcal{Z}_N = \mathcal{Z}_N$
- Intuitively: Random variable  $z_i$  denotes the candidate  $m_{ir} \in \mathcal{C}_i$  assigned to feature i.
- Pairwise Markov Random Field as probabilistic model [5]:

$$p(\mathbf{z}) = \frac{1}{Z} \prod_{i \in V} \psi_i(z_i) \prod_{(i,j) \in E} \psi_{ij}(z_i, z_j)$$

- Potential functions:  $\psi_i(z_i)$  MS score and  $\psi_{ij}(z_i,z_i)$  match of observed and **predicted** retention order
- Molecular candidates ranked based on max-marginals [5] (Fig. 2):

$$p_{\max}(z_i = r) = \max_{\{\mathbf{z}' \in \mathcal{Z} \mid z_i' = r\}} p(\mathbf{z'})$$

• Intuitively: Maximum marginal probability of a candidate assignment z with  $z_i = r$ .

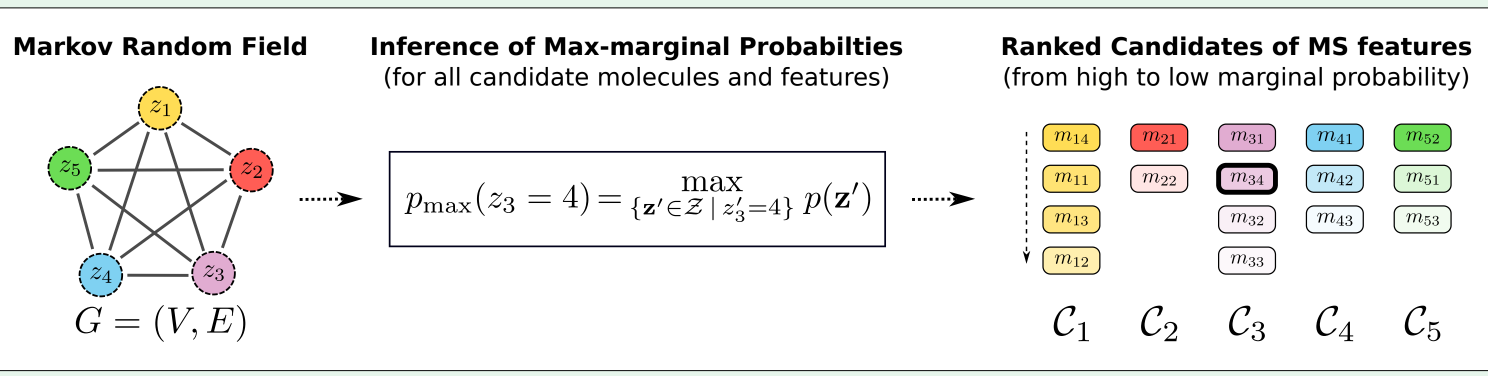


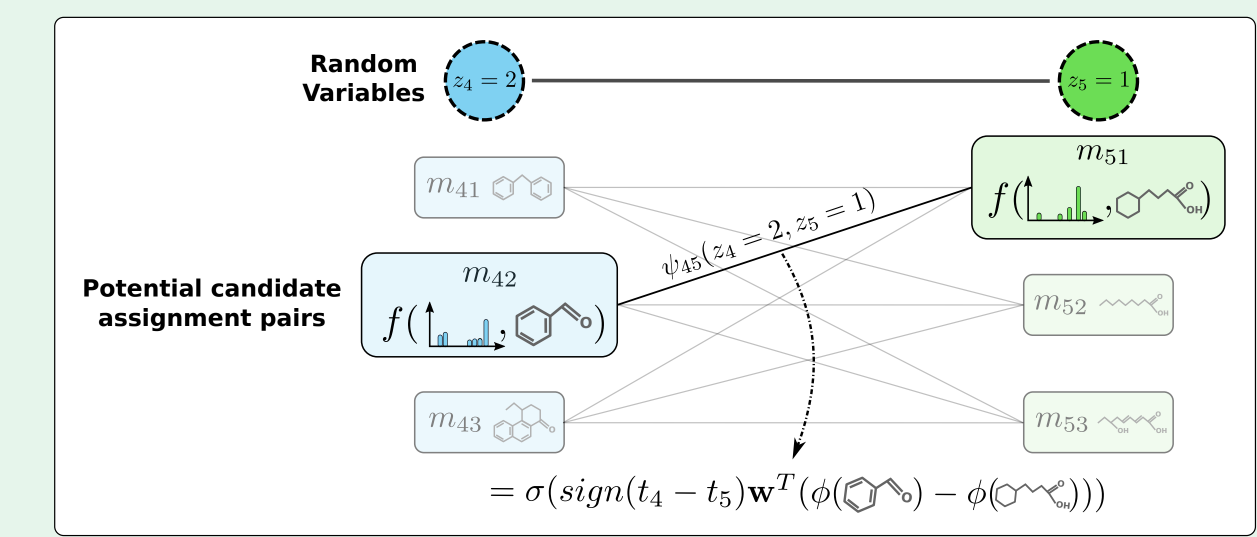
Fig. 2: MRF probability distribution and candidate ranking, e.g. MS feature i=3 and candidate 4 ( $m_{34}$ ).

### 5. Encoding MS and Retention Order Information: $\psi_i$ and $\psi_{ii}$

- Goodness of the candidate assignment z modeled by potential functions  $\psi$  (Fig. 3)
- Node potential  $\psi_i: \mathcal{Z}_i \to \mathbb{R}_{>0}$ :  $\psi_i(z_i = r) = f(x_i, m_{ir})$
- o f returns the MS matching score  $\in (0,1]$  of spectrum  $x_i$  and candidate  $m_{ir}$
- Edge potential  $\psi_{ij}: \mathcal{Z}_i \times \mathcal{Z}_j \to \mathbb{R}_{>0}$ , with  $\sigma$  being the sigmoid function:

$$\psi_{ij}(z_i = r, z_j = s) = \sigma(\underbrace{\operatorname{sign}(t_i - t_j)}_{\text{observed}} \cdot \underbrace{\mathbf{w}^T(\phi(m_{ir}) - \phi(m_{js}))}_{\text{predicted retention order}})$$

- Intuitively: Matching observed and predicted retention orders receive high scores.
- Retention order prediction using Ranking Support Vector Machine (RankSVM) w [2]
- Candidate molecules  $m_{ir}$  representation using non-linear features  $\phi$



3: Example: Node and edge score for all candidate pairs of feature i=4 and j=5.

## 6. Experiments and Results

- Evaluation datasets: CASMI 2016 [7], EA subset from MassBank used by [6]
- 681 (MS<sup>2</sup>, RT)-tuples with each 310 candidates (median statistic)
- Datasets cover two different LC columns and flow gradients
- RankSVM training data: 1248 RTs from PredRed [8] and CASMI 2016 training
- No evaluation set molecule in RankSVM tranining set
- **Performance measure:** Top-k accuracy, percentage of correct molecular candidates at rank  $\leq k$

### **Experiment 1: Comparison to MetFrag + LogP (RT Proxy) Prediction**

MetFrag relaunched [6]: Prediction of LogP values for candidates, linear model mapping measured RTs to LogPs, candidate re-ranking based on LogP deviation

Method	Top-1	Top-5	Top-10	Top-20
$MS^2 + RT (Our)$	21.3	52.9	64.0	74.3
MS <sup>2</sup> + RT (MetFrag & LogP)	20.5	49.1	61.2	72.6
Only $MS^2$ (baseline)	16.7	49.5	60.4	70.6

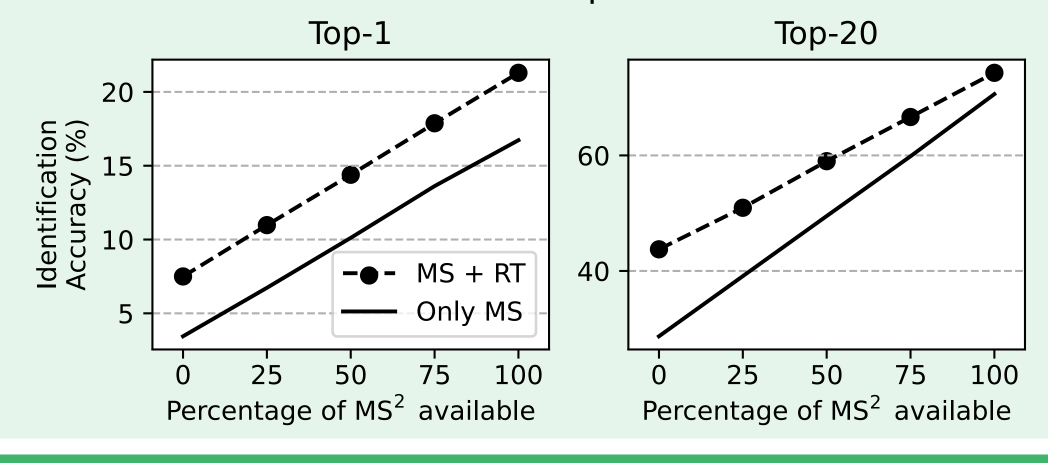
### **Experiment 2: Performance with different MS<sup>2</sup>-Scoring Methods**

MetFrag (in-silico fragmenter scores) and IOKR [3] as  $MS^2$ -scoring methods

MS <sup>2</sup> -Scorer	Method	Top-1	Top-5	Top-10	Top-20
MetFrag	$MS^2 + RT$ (our)		52.9	64.0	74.3
	Only MS <sup>2</sup> (baseline)	16.7	49.5	60.4	70.6
IOKR	<b>\</b>	26.7		62.5	70.3
	Only MS <sup>2</sup> (baseline)	25.1	49.5	60.3	67.6

#### **Experiment 3: Missing MS<sup>2</sup> Spectra**

- Simulating missing MS<sup>2</sup> information: Varying from 0% to 100% MS<sup>2</sup>
- If only MS<sup>1</sup>: Use mass deviation between precursor and candidate molecule



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