Liquid-Chromatography Retention Order Prediction for Metabolite Identification

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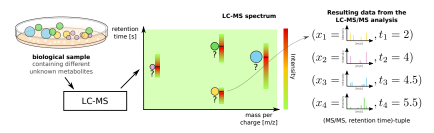
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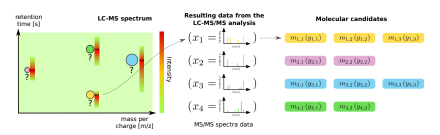
Metabolite identification

- Small molecules (< 1000 Da) involved in biological processes
- Identification of metabolites present in a biological sample
- Widely used analysis workflow: Liquid chromatography (LC) combined with tandem mass spectrometry (MS/MS)



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State-of-the-art MS/MS based metabolite identification



Identification workflow

- 1. For each MS/MS spectra x_i define a set of molecular candidate structures $\{m_{i,1}, m_{i,2}, \ldots\}$. using the molecular mass
- 2. Assign a "MS/MS matching score" $y_{i,j}$ to each candidate. Input-Ouput-Kernel-Regression[Bro+16]
- 3. Higest scoring candidate $m_{i,j}$ is the identification for spectra x_i .

Retention times (RTs) for metabolite identification

- Retention times are *valuable* orthogonal information [Rut+16; SNV15; Aic+15] distinction of diastereoisomers
- State-of-the-art machine learning metabolite identification methods use only MS/MS information [Bro+16; Düh+15]

Challenges utilizing RTs

- 1. Measurements are LC-system specific.
- 2. Public datasets are relatively *small* and originate from *heterogeneous systems*

Proposed method to tackle the challenges

- 1. **Predict the pairwise retention order** of molecular candidate structures using preference learning
- Prediction model can be trained on multiple retention time datasets arising from heterogeneous LC-systems.
- Retention orders are largely preserved across LC-systems [SNV15].
- 2. Integrating predicted candidate retention orders and MS/MS based scores to *jointly* identify a set of metabolites.

Retention order pairs for preference learning

Notation

- Molecule m_i from the molecular space \mathcal{M}
- $t_i \in \mathbb{R}_+$ its retention time
- $s_i \in \mathcal{S}$ chromatographic system it has been measured with

Pairwise molecule preference

- m_i is preferred over m_i when it elutes before m_i , i.e. $t_i < t_i$
- Set of pairwise preferences of given LC-system s is defined as:

$$\mathcal{P}(s) = \{(i,j) | s_i = s_j = s, t_i < t_j\}$$

• Set of pairwise preferences from *multiple* LC-systems:

$$\mathcal{P} = \bigcup_{s \in S} \mathcal{P}(s)$$

We want to learn a pairwise retention order prediction function:

$$f(m_i = 0), m_j = 0) = \begin{cases} 1 & m_i = 0 \\ -1 & \text{otherwise} \end{cases}$$
 elutes before $m_j = 0$

Kernelized RankSVM prediction model

$$f(m_i, m_j) = \operatorname{sign}(\mathbf{w}^T(\phi(m_j) - \phi(m_i)))$$

- w are the RankSVM [Joa02; KLL14] model parameters
- $\phi: \mathcal{M} \to \mathcal{F}_m$ feature map associated with k_m embedding the molecular structures into a feature space.
- Kernel function $k_m: \mathcal{M} \times \mathcal{M} \to \mathcal{R}$ encodes similarity between molecular structures

Training the RankSVM for retention order prediction

We optimize \mathbf{w} considering the pairwise preferences \mathcal{P} from (possibly) different chromatographic systems:

$$\min_{\mathbf{w},\xi} \quad \frac{1}{2} \mathbf{w}^T \mathbf{w} + C \sum_{(i,j) \in \mathcal{P}} \xi_{ij}
s.t. \quad \mathbf{w}^T (\phi(m_j) - \phi(m_i)) \ge 1 - \xi_{ij}, \forall (i,j) \in \mathcal{P}
\xi_{ii} > 0, \forall (i,j) \in \mathcal{P},$$

with C > 0 being the regularization parameter.

Learned model

$$\mathbf{w}^T \phi(m_i) < \mathbf{w}^T \phi(m_i), \text{ if } (i,j) \in \mathcal{P}$$

Evaluating retention order prediction

Dataset

- 1098 retention times of 946 unique molecular structures
- 5 different reversed phase LC-systems (denoted with \hat{S})
- We use counting MACCS dictionary fingerprints with MinMax-Kernel [Ral+05].

Evaluation measure and protocol

• Pairwise prediction accuracy for a target system $s \in \hat{S}$:

$$Acc(s) \equiv \frac{|\{(i,j) \in \mathcal{P}(s) \mid \mathbf{w}^T \phi(m_i) < \mathbf{w}^T \phi(m_j)\}|}{\mathcal{P}(s)}$$

- Accuracy accessed using repeated 10-fold cross-validation.
- no test molecular structure in the training set

Can pairwise predictor benefit from information of different systems?

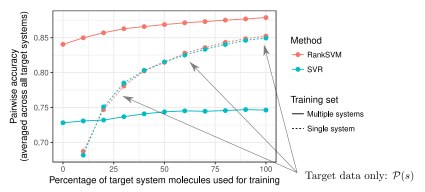
Compare performance of different training sets

- Single system, target data only: $\mathcal{P}(s)$
- Multiple systems, *no* target data: $\mathcal{P} \setminus \mathcal{P}(s)$
- ullet Multiple systems, all available data: ${\cal P}$
- Varying percentage of target system molecules used for training

Comparison method

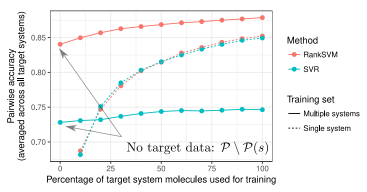
- Support Vector Regression (SVR) trained on retention times directly [Aic+15].
- Multiple systems: Retention times are considered jointly.

Application setting: Training retention times only available from single target system.



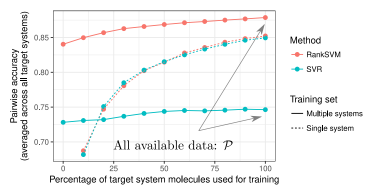
- Increasing amount of training data improves prediction.
- RankSVM and SVR perform equally.

Application Setting: Training retention times only available from not target system.



- Performance of single system without data from the target.
- RankSVM outperforms SVR by considering retention orders.

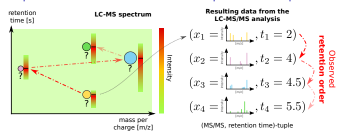
Application Setting: Training retention times from target and others systems available.



- Considering target and non-target systems' data outperforms single system.
- RankSVM again outperforms SVR.

Predicted retention orders for metabolite identification

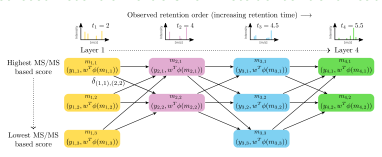
Exploit observed retention order in complex LC-MS experiment



Identification workflow

- 1. execute 1. (query candidates) and 2. (predict matching scores)
- 2. Predict retention orders between all candidates $m_{i,j}$ and $m_{i+1,s}$ of MS/MS spectra of consecutively eluting molecules. x_1 and x_2
- 3. Find the *overall* most consistent metabolite identification using the shorest path algorithm.

Predicted retention orders for metabolite identification



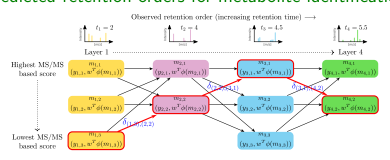
Edges connecting candidates of consecutive layers with edge weight:

$$\delta_{(i,j),(i+1,s)} = -y_{i+1,s} + D \cdot \max(0, \underbrace{\mathbf{w}^T(\phi(m_{i,j}) - \phi(m_{i+1,s}))}_{\text{RankSVM order penalty}})),$$

 $D \ge 0$ weight on order penmalty: $\max(...) > 0$ if observed \ne predicted order.

 Candidates along the shortest path from first to last layer: mos consistent identification.

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Experiments metabolite identification

Dataset

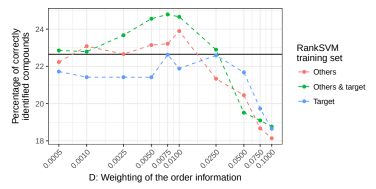
- 342 reversed phase LC-retention times
- o for 120 MS/MS spectra available \rightarrow (MS/MS, RT)-tuple
- \circ remaining 222 RTs are used for RankSVM training (s_{Impact})
- Additionally we use the 5 datasets \hat{S} of the previous experiments for RankSVM training

Evaluation measure and protocol

- randomly sample 1000 times 80 (MS/MS, RT)-tuples
- We construct the graph containing the candidates run the shortest path algorithm.
- Percentage of correct identifications for different values of D
- Comparison to baseline performance when D=0

Experiments metabolite identification

Baseline performance 22.7%: (D = 0, only MS/MS spectra used, black line)



- Improved identification accuracy for Others (23.9%) and Others & target (24.8%)
- RankSVM trained only on the target data cannot improve.

Summary

- Proposed a method for predicting liquid chromatographic orders using RankSVM.
- Prediction model can be trained on retention time data from different chromatographic systems.
- Proposed method to integrate predicted retention orders and MS/MS scores for metabolite identification in LC-MS setting
- Metabolite identification accuracy can be improved using predicted retention orders.

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Source code available

https://version.aalto.fi/ gitlab/bache1/retention_order_ prediction



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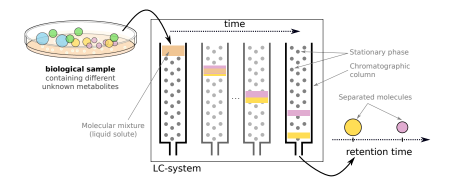


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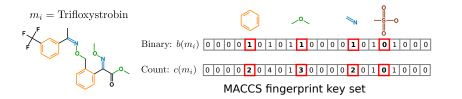


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Liquid-Chromatography principle



Molecules represented using MACCS dictionary fingerprints



Kernels used for the feature embedding in RankSVM

• Binary: Tanimoto kernel [Ral+05]

$$k_m(m_i, m_j) = \frac{|b(m_i) \cap b(m_j)|}{|b(m_i) \cup b(m_i)|}$$

• Count: MinMax kernel [Ral+05]

$$k_{m}(m_{i}, m_{j}) = \frac{\sum_{s=1}^{N_{sub}} \min(c_{s}(m_{i}), c_{s}(m_{j}))}{\sum_{s=1}^{N_{sub}} \max(c_{s}(m_{i}), c_{s}(m_{i}))}$$

Compare binary and counting molecular fingerprints

- Pairwise prediction accuracy $(\pm 2\sigma)$ for different target systems
- RankSVM models trained using single system $\mathcal{P}(s)$.

Target system s	Binary MACCS	Counting MACCS
Eawag_XBridgeC18	$0.796(\pm0.015)$	$0.844(\pm0.011)$
$FEM_{J}long$	$0.882(\pm0.016)$	$0.905 (\pm 0.015)$
RIKEN	$0.826(\pm0.024)$	$0.848 (\pm 0.017)$
$UFZ_{-}Phenomenex$	$0.790(\pm0.027)$	$0.802 (\pm 0.017)$
LIFE_old	$0.842(\pm0.050)$	$0.862 (\pm 0.035)$