Fast dictionary-based approach for mass spectrometry data analysis

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Joint work with



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Motivation

Mass Spectrometry (1/2)

■ Mass Spectrometry :

- A technique used to measure the characteristics, the chemical composition and the structure of a sample or molecule.
- 1919, Joseph John Thomson.

■ Utility area:

- Pharmaceutical: drug discovery, combinatorial chemistry, pharmacokinetics, drug metabolism.
- Clinical: neonatal screening, haemoglobin analysis, drug testing.
- Environmental : water quality, food contamination.
- Geological : oil composition.
- Biotechnology: analysis of proteins, peptides.

Mass Spectrometry (2/2)

■ Principle of measure :



Aim:

Analyse efficiently big MS data.

Outline

- Problem statement
- Observation model
 - Mathematical model
 - Dictionary-based strategy
 - III-posed problem
- Optimization strategy
 - Variational formulation
 - Primal dual algorithm
- Practical Implementation
 - Dictionary construction
 - Circulant approximation
- Application
 - Synthetic results
 - Experimental results
 - MS spectrum analysis

Problem statement

Some reminders on chemistry

Atom?

- Atomic number = Number of protons.
- Mass number = Number of protons + neutrons.

■ Molecule?

Set of atoms linked together.

■ Isotopic pattern?

An atom can be present under different forms with different numbers of neutron, called **isotopes**.

Each stable isotope is present in the nature with a specific abundance.

Atomic Structure



Protein class (1/2)

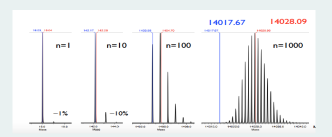
■ Protein formula $(C_{N_C}H_{N_H}O_{N_O}N_{N_N}S_{N_S})$

Atom Name	Atom Symbol	Mass (in Dalton)	Relative Abundance	
Carbon	12 _C	12 (by definition)	0.9893	
	13 _C	13.0033548378	0.0107	
Hydrogen	¹ H	1.00782503207	0.999885	
	² H	2.0141017778	0.000115	
Oxygen	¹⁶ O	15.99491461956	0.99757	
	¹⁷ O	16.99913170	0.00038	
	¹⁸ O	17.9991610	0.00205	
Nitrogen	¹⁴ N	14.0030740048	0.99636	
	¹⁵ N	15.0001088982	0.00364	
Sulfur	³² S	31.97207100	0.9499	
	³³ S	32.97145876	0.0075	
	³⁴ S	33.96786690	0.0425	

Isotopic mass and natural abundance of atoms found in proteins (by definition)

Protein class (2/2)

■ Example of Alkane molecule C_nH_{2n+2} $(n \in \mathbb{N}^*)$:



→ The larger the molecule, the larger is the number of different isotopes present, associated with specific probabilities of appearance.

How to find the position of the mono-isotopic peak from a large pattern distribution?



Mathematical model

For a given chemical sample, composed of P different molecules with monoisotopic mass $m_p^{\text{iso}} \in (0, +\infty)$, charge state $z_p \in \mathbb{N}^*$ and abundance $a_p \in (0, +\infty)$, for $p \in \{1, \dots, P\}$:

$$y = \sum_{p=1}^{P} a_{p} D(m_{p}^{iso}, z_{p}) + n$$
 (1)

- y : acquired MS spectrum.
- $D(m_p^{iso}, z_p)$: mass distribution associated to monoisotopic mass m_p^{iso} and charge state z_p .
- n : acquisition noise.
 - Discrete measurements on a grid with size M :

$$\mathbf{y} = \sum_{p=1}^{P} \mathbf{a}_{p} \mathbf{d}(\mathbf{m}_{p}^{\mathsf{iso}}, \mathbf{z}_{p}) + \mathbf{n}$$
 (2)

with $\mathbf{y} \in \mathbb{R}^M$, $\mathbf{d}(m_p^{\text{iso}}, z_p) \in [0, +\infty[^M \text{ and } \mathbf{n} \in \mathbb{R}^M]$.

Dictionary-based strategy

Protein molecule: Averagine model [Senko et al., 1994]

$$\begin{array}{cccc}
\mathcal{A}: & \mathbb{R}_{+} \times \mathbb{N}^{*} & \rightarrow & \mathbb{R}_{+} \\
 & (m, z) & \rightarrow & \mathbf{d}(m, z)
\end{array} \tag{3}$$

For M candidate values of isotopic masses and Z candidate charge values, we define a grid with size T = MZ, and we define the dictionary $\mathbf{D} \in \mathbb{R}^{M \times T}$:

$$D = [D_{1 \le j \le M, 1}, D_{1 \le j \le M, 2}, ..., D_{1 \le j \le M, T}]$$
(4)

ullet *i*-th column of $oldsymbol{D}$ determined by the averagine model at *i*-th grid position :

$$(D)_{1 \le i \le M, i} = d(m_i, z_i)$$

Inverse problem

Problem

$$y = D\overline{x} + n' \tag{5}$$

where $\mathbf{y} \in \mathbb{R}^M$, $\mathbf{D} \in \mathbb{R}^{M \times T}$, $\overline{\mathbf{x}} \in \mathbb{R}_+^T$ and $\mathbf{n}' \in \mathbb{R}^M$.

- \times y : measure.
- D : large scale ill-conditioned matrix.
- X T = MZ: very large size.
- \mathbf{x} \mathbf{x} : unknown sparse vector with positive entries.
- x n' : high noise level.

⇒ Ill-posed inverse problem in high dimensions.

Aim

Find $\hat{\mathbf{x}}$ from \mathbf{y} and \mathbf{D} such that $\hat{\mathbf{x}} \simeq \overline{\mathbf{x}}$ (6)

Optimization strategy

Variational formulation

$$\underset{\mathbf{x} \in \mathbb{R}^{T}}{\operatorname{minimize}} \ \Phi(\mathbf{x}) \quad \text{subject to} \quad \|\mathbf{D}\mathbf{x} - \mathbf{y}\| \le \eta \tag{7}$$

- $\Phi: \mathbb{R}^T \mapsto]-\infty, +\infty]$ is a proper, lower semicontinuous (lsc), convex regularization function used to enforce positivity and sparsity on the solution.
- $\eta > 0$ is a parameter that depends on the noise characteristics.

Proximity operator

Let $\Phi : \mathbb{R} \to]-\infty, +\infty]$ a lsc proper and convex function. The proximity operator of Φ is defined as [Moreau, 1965]

http://proximity-operator.net/ [Cherni et al., 2017]:

$$\operatorname{prox}_{\Phi}(x) : \mathbb{R}^{\mathbb{N}} \to \mathbb{R}^{\mathbb{N}}$$
$$x \to \underset{y \in \mathbb{R}^{\mathbb{N}}}{\operatorname{argmin}} \left(\Phi(y) + \frac{1}{2} \|y - x\|^2 \right)$$

Primal Dual algorithm [Chambolle and Pock, 2011]

Initialisation $\mathbf{u}^{(0)} \in \mathbb{R}^M, \mathbf{x}^{(0)} \in \mathbb{R}^T$

$$0 < \sigma < \|\mathbf{D}\|^2 / \tau$$

 $\rho \in (0, 2), \ \tau > 0$

Minimisation

For k = 0, 1, ...

$$\begin{split} \tilde{\mathbf{x}}^{(k)} &= \mathsf{prox}_{\tau \Phi} \big(\mathbf{x}^{(k-1)} - \tau \mathbf{D}^{\mathsf{T}} \big(\mathbf{u}^{(k-1)} \big) \big) \\ \mathbf{v}^{(k)} &= \mathbf{u}^{(k-1)} + \sigma \mathbf{D} \big(2 \tilde{\mathbf{x}}^{(k)} - \mathbf{x}^{(k-1)} \big) \\ \tilde{\mathbf{u}}^{(k)} &= \mathbf{v}^{(k)} - \sigma \mathsf{proj}_{\| - \mathbf{y} \| \le \eta} \big(\mathbf{v}^{(k)} \big/ \sigma \big) \\ \mathbf{x}^{(k)} &= \mathbf{x}^{(k-1)} + \rho \big(\tilde{\mathbf{x}}^{(k)} - \mathbf{x}^{(k-1)} \big) \\ \mathbf{u}^{(k)} &= \mathbf{u}^{(k-1)} + \rho \big(\tilde{\mathbf{u}}^{(k)} - \mathbf{u}^{(k-1)} \big) \end{split}$$

For every $(\mathbf{y}, \mathbf{v}) \in (\mathbb{R}^N)^2$:

$$\operatorname{proj}_{\|\cdot - \mathbf{y}\| \le \eta}(\mathbf{v}) = \mathbf{v} + (\mathbf{v} - \mathbf{y}) \min \left(\frac{\eta}{\|\mathbf{v} - \mathbf{v}\|}, 1 \right) - \mathbf{y}. \tag{8}$$

► The convergence of the iterates $(\mathbf{x}^{(k)})_{(k \in N)}$ is ensured [Chambolle and Pock, 2011].

Practical Implementation

Dictionary construction

Given a range of masses $[m_{\min}, m_{\max}]$ and charges $[z_{\min}, z_{\max}]$, we define a regular grid:

$$(\forall i \in \{1,\ldots,T\}) \quad m_i = m_{\min} + (j-1)m_{\max}, \tag{9}$$

$$z_i = z_{\min} + (\ell - 1)z_{\max}, \qquad (10)$$

with the convention $i = \ell M + j$, $j \in \{1, ..., M\}$ and $\ell \in \{1, ..., Z\}$.

- The *i*-th column of **D** is taken as $d(m_i, z_i)$.
- $\mathbf{d}(m_i, z_i)$ corresponds to a sampled version of D(m, z) on the mass grid with size M.

Circulant approximation

- ▶ Difficulty
 - X Very large value for MZ.
 - ⇒ Large dictionary D which presents a computational challenge and large memory resources.
- Assumptions
 - Similar isotopic mass patterns mainly differ by a sample shift of peaks positions.
 - Isotopic patterns are sparse with positives entries.

Proposal

 \rightarrow Decompose the mass axis into windows with width $L \leq M$.

Circulant approximation

► Initial dictionary

$$D = [D_{1 \le i \le M,1}, D_{1 \le i \le M,2}, ..., D_{1 \le i \le M,T}] \quad T = MZ$$
 (11)

$$D = [D_1|...|D_{\ell}|...|D_Z]_{1 < \ell < Z}$$
 (12)

▶ Circulant model

Each \mathbf{D}_{ℓ} is approximated by the following block diagonal (BDiag) matrix made of M/L blocks assumed to be circulant (Circ) matrices with first line $\overline{\mathbf{d}}_{s,\ell}$, $s \in \{1,\ldots,M/L\}$:

$$\overline{\mathbf{D}}_{\ell} = \mathsf{BDiag}\left(\left[\mathsf{Circ}\left(\overline{\mathbf{d}}_{s,\ell}\right)\right]_{1 \le s \le M/L}\right).$$
 (13)

- \checkmark Fast computation of products (D, D^T) using fast Fourier tools.
- ✓ High reduction of the memory requirements.

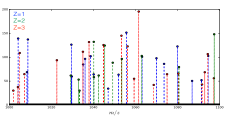
$$\overline{\mathbf{D}} = \left[\overline{\mathbf{D}}_1 | \dots | \overline{\mathbf{D}}_{\ell} | \dots | \overline{\mathbf{D}}_Z\right]_{1 < \ell < Z} \tag{14}$$

Application

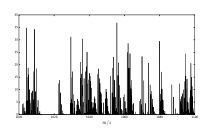
Synthetic results (1/4)

Simulated data

- Signal A:
 - M = 5000
 - Z = 3, $z_{min} = 1$, $z_{max} = 3$
 - P = 50 proteins
 - Mass axis = [1000, 1100]



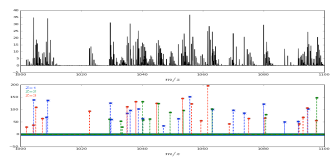
Original signal



Measured data

Synthetic results (2/4)

- Parameters dataset : Gaussian noise, iid with std $\sigma = 10^{-2}$, $\tau = \theta \sigma \sqrt{M}$. $\theta \simeq 1$.
- Parameters algorithm : $\rho = 1.9$, $\tau = \|\mathbf{D}\|^{-1}$, $\sigma = 0.9\tau$, maximum iterations number=1000.



Reconstruction results of the signal A: (top) input data y, (bottom) exact spectrum (dots), restored spectrum with exact dictionary (dashed line), and with its block-circulant approximation for L=10 (asterisks).

Synthetic results (3/4)

■ Impact of noise? Memory storage?

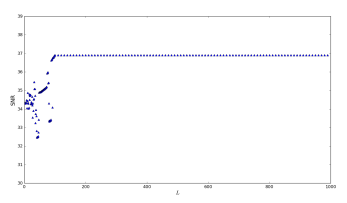
σ.	Exact dictionary approach			Block-circulant approximation		
σ	SNR	Time	Memory storage	SNR	Time	Memory
1	16.18	303.33		15.57	127.85	
0.1	35.73	206.84	572	35.43	44.48	0.53
0.01	39.56	377.80		38.38	290.56	

SNR (in dB), computation time (in s) and memory storage (in MB) for the restoration of signal A for various values of noise level. Block-circulant approximation $\overline{\mathbf{D}}$ is tested for L = 10.

- Good quality reconstruction even with high noise level.
- ▶ Block-circulant approximation is faster than exact dictionary approach with limited deterioration of the results quality.

Synthetic results (4/4)

 \blacksquare Influence of L?



SNR of the restored signal A with $\sigma = 10^{-2}$ using $\overline{\mathbf{D}}$ for various L values

▶ Reconstruction quality stable to the value of *L*.

Experimental results

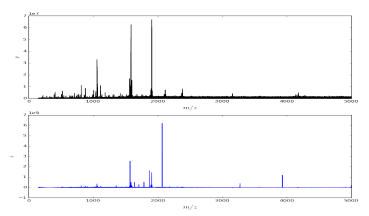
Parameters dataset

- peptide EVEALEKKVAALESKVQALEKKVEALEHG-NH2
- $3 \mu M (C_{140} H_{240} N_{38} O_{45}).$
- Trimer form within 50 mM of NH₄OAc
- BRUKER Solarix 15 T, FT-ICR instrument, ESI source.
- N = 8000000

■ Parameters algorithm

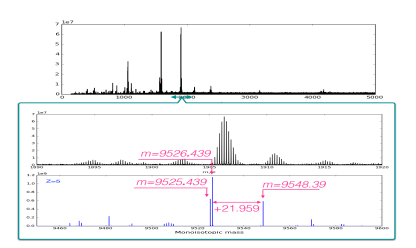
- \bullet σ : estimated from an empty frame of the measured signal.
- maximum iterations number: 2000

Experimental results



Analysis of the real FT-ICR MS spectrum of peptide in trimer form : (top) acquired data, (bottom) recovered spectrum z = 5 using block-circulant approximated dictionary with L = 10

MS spectrum analysis



- Theoretical monoisotopic mass of the peptide : m = 9526.337 Daltons.
- Theoretical position of the Sodium : m = +21.982 Daltons.

Conclusion & perspectives

Conclusion & perspectives

- √ New dictionary-based approach for MS data analysis based on proteomic averagine model.
- $\sqrt{}$ Penalized cost function promoting sparsity and positivity, minimized with efficient primal-dual scheme with sought convergence guarantees.
- √ Block-circulant approximation of the dictionary-based approach.
- ⇒ Efficient analysis of synthetic and real MS spectra.
- ⇒ Fast approach devoted to the big data scale.
- ⇒ Limited memory resources required.

Conclusion & perspectives

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- √ Block-circulant approximation of the dictionary-based approach.
- ⇒ Efficient analysis of synthetic and real MS spectra.
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- ⇒ Limited memory resources required.

▶ Extend this approach to the processing of multidimensional MS spectra.

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