Manual: PEPICOBayes

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1 Introduction

PEPICOBayes is a program for reconstructing the probe spectrum from pump-probe and pump-only photoelectron-photoion coincidence (PEPICO) measurements applying Bayesian probability theory. The theory behind the program is described in reference [1]. The PEPICOBayes program is written in C and executed in Matlab. We offer a simple example Matlab-file to demonstrate the usage of the PEPICOBayes program.

2 Abbreviations

The abbreviations in this manual are similar to the abbreviations used in reference [1].

Abbreviation	Description
	Symbol that refers to $j \in \{1, 2\}$
$\mid j \mid$	$1 \cdots$ events due to pump-only laser pulse
	$2 \cdots$ events due to probe-only laser pulse
	Symbol that refers to $\rho \in \{\alpha, \beta\}$
ρ	$\alpha \cdots$ events in the pump-only experiment
	$\beta \cdots$ events in the pump-probe experiment
$n^{(ho)}$	measured count rates $n^{(\rho)} = \{n_{\mu\nu}^{(\rho)}\}$, dimension: $[\mathcal{N}_{\mu} \times \mathcal{N}_{\nu}]$
$q^{(j)}$	Monte Carlo time series of the spectrum $q^{(j)} = \{q_{\mu\nu}^{(j)}\},$
	dimension: $[\mathcal{N}_{\mu} \times \mathcal{N}_{\nu} \times N_{\text{sweep}}]$
$N_{ m run}$	Number of Monte Carlo steps between Monte Carlo measurements
$N_{ m sweep}$	Number of Monte Carlo measurements
\mathcal{N}_{μ} $\mathcal{N}_{ u}$	Number of elements in dimension μ
$\mathcal{N}_{ u}$	Number of elements in dimension ν
\mathcal{N}_p	Number of measurements
M	Number of measurements where N_e electrons and N_i ions were
N_{N_e,N_i}	detected.
π	Time series of experimental parameters: $\pi = \{\underline{\lambda}_1, \underline{\lambda}_2, \sigma_1, \sigma_2, \xi_i, \xi_e\},$
71	dimension: $[6 \times N_{\text{sweep}}]$
λ_j	Mean number of events in channel j
$\left egin{array}{c} \underline{\lambda}_j \ \overline{\xi}_i \end{array} ight $	Detection probability for ions
ξ_e	Detection probability for electrons

3 PEPICOBayes-Program

3.1 Function definition

The function returns four arguments and can be executed with six or eight arguments as input. Arguments seven and eight $(q_0^{(1)} \text{ and } q_0^{(2)})$ are start values for the spectra and optional. If no start spectra are passed the algorithm starts with flat spectra.

Input variables:

	Abbreviation	Variable name	Description	
1	$n^{(eta)}$	n_beta	measured pump-probe coincidence spectrum (dimension $[\mathcal{N}_{\mu} \times \mathcal{N}_{\nu}]$)	
2	$n^{(\alpha)}$	n_alpha	measured pump-only coincidence spectrum (dimension $[\mathcal{N}_{\mu} \times \mathcal{N}_{\nu}]$)	
3	$N_{ m run}$	Nrun	Number of Monte Carlo steps between measurements	
4	$N_{ m sweep}$	Nsweep	Number of Monte Carlo measurements	
5	parameter	parameter	$\begin{array}{l} \text{Parameter vector:} \\ \text{parameter := } \{\underline{\lambda}_{1}, \underline{\lambda}_{2}, \sigma_{1}, \sigma_{2}, \xi_{i}, \xi_{e}, \\ \sigma_{q^{(1)}}, \sigma_{q^{(\beta)}}, \sigma_{\underline{\lambda}_{1}}, \sigma_{\underline{\lambda}_{2}}, \sigma_{\sigma_{1}}, \sigma_{\sigma_{2}}, \sigma_{\xi_{i}}, \sigma_{\xi_{e}}, \\ \mathcal{N}_{p}^{(\beta)}, N_{00}^{(\beta)}, N_{01}^{(\beta)}, N_{02}^{(\beta)}, N_{03}^{(\beta)}, N_{10}^{(\beta)}, N_{11}^{(\beta)}, N_{12}^{(\beta)}, N_{20}^{(\beta)}, N_{21}^{(\beta)}, N_{30}^{(\beta)}, \\ \mathcal{N}_{p}^{(\alpha)}, N_{00}^{(\alpha)}, N_{01}^{(\alpha)}, N_{02}^{(\alpha)}, N_{03}^{(\alpha)}, N_{10}^{(\alpha)}, N_{11}^{(\alpha)}, N_{12}^{(\alpha)}, N_{20}^{(\alpha)}, N_{21}^{(\alpha)}, N_{30}^{(\alpha)} \} \end{array}$	
6	$q_0^{(1)}$		(optional) start values of $q^{(1)}$ (dimension $[\mathcal{N}_{\mu} \times \mathcal{N}_{\nu}]$)	
7	$q_0^{(2)}$		(optional) start values of $q^{(2)}$ (dimension $[\mathcal{N}_{\mu} \times \mathcal{N}_{\nu}]$)	

The size of a Monte Carlo step of variable $T \in \{q^{(1)}, q^{(2)}, \underline{\lambda}_1, \underline{\lambda}_2, \sigma_1, \sigma_2, \xi_i, \xi_e\}$ is Gaussian distributed with mean value zero and standard deviation σ_T . σ_T should be adjusted that $p_{\rm acc}$ is at the order of 50% in order to efficiently sample the probability distribution. Note that the standard deviation of for the step distribution for σ_j is $\min\{\sigma_{\sigma_j}, \sigma_j\}$. This is important if the fluctuations are very small and allows to visit phase space with a very small σ_j . For numerical stability the smallest value of σ_j is 10^{-8} .

Output variables:

	Abbreviation	Variable name	Description
1	$q^{(1)}$	q1_timeseries	Time series of $q^{(1)}$ (dimension $[\mathcal{N}_{\mu} \times \mathcal{N}_{\nu} \times N_{\text{sweep}}]$)
2	$q^{(2)}$	$q2_timeseries$	Time series of $q^{(2)}$ (dimension $[\mathcal{N}_{\mu} \times \mathcal{N}_{\nu} \times N_{\text{sweep}}]$)
3	π	Pi	Time series of π (dimension [6 x N_{sweep}])
4	$p_{ m acc}$	p_acc	Vector of acceptance rates with 6 entries: $p_{\rm acc}(1): \text{ Acceptance rate of } \underline{\lambda}_1$ $p_{\rm acc}(2): \text{ Acceptance rate of } \underline{\lambda}_2$ $p_{\rm acc}(3): \text{ Acceptance rate of } \sigma_1$ $p_{\rm acc}(4): \text{ Acceptance rate of } \sigma_2$ $p_{\rm acc}(5): \text{ Acceptance rate of } \xi_i$ $p_{\rm acc}(6): \text{ Acceptance rate of } \xi_e$ $p_{\rm acc}(7): \text{ Acceptance rate of } q^{(1)}$ $p_{\rm acc}(8): \text{ Acceptance rate of } q^{(2)}$

Note that the first entry in the time series is the start value of the corresponding variable.

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PEPICOBayes (n^{(\beta)}, n^{(\alpha)}, N_{\text{run}}, N_{\text{sweep}}, \text{ parameter}, q_0^{(1)}, q_0^{(2)}) read all input variables for k < N_{\text{sweep}} for i < N_{\text{run}} suggest Monte Carlo step in \underline{\lambda}_1, \underline{\lambda}_2, \sigma_1, \sigma_2, \xi_e, \xi_i, q^{(1)} or q^{(2)} compute Monte Carlo step probability if update is accepted overwrite corresponding variable with the new value else do nothing end end copy \underline{\lambda}_1, \underline{\lambda}_2, \sigma_1, \sigma_2, \xi_e, \xi_i, q^{(1)} and q^{(2)} into output variables at Monte Carlo time k end end
```

3.2 Pseudocode of the algorithm

A Metropolis Hastings Monte Carlo algorithm is used to sample the probability distribution for $q^{(1)}$, $q^{(2)}$ and π .

3.3 Start value for π

The start values of π can for example be approximated by the formula given by Stert et. al. [2]. In summary this means solving the following equation for λ

$$\frac{N_e N_i}{\lambda \mathcal{N}_p} \left(1 + \lambda \left(1 - \frac{N_e}{\lambda \mathcal{N}_p} \right) \left(1 - \frac{N_i}{\lambda \mathcal{N}_p} \right) \right) e^{-\lambda \left(1 - \left(1 - \frac{N_e}{\lambda \mathcal{N}_p} \right) \left(1 - \frac{N_i}{\lambda \mathcal{N}_p} \right) \right)} \approx N_{11} \tag{1}$$

and insert the resulting λ into

$$\xi_e \approx \frac{N_e}{\lambda \mathcal{N}_p} \tag{2}$$

$$\xi_i \approx \frac{N_i}{\lambda \mathcal{N}_p} \tag{3}$$

 N_{11} is the total number of measured coincidencies and N_e and N_i are the total number of detected electrons and ions, respectively.

4 Results of example.m

In the example two ion types $(\mathcal{N}_{\mu}=2)$ and seven different electron flight times $(\mathcal{N}_{\nu}=2)$ were detected.

The result of the converged Monte Carlo computation for the example data should look like figure 3. Converged time series are shown in figure 1 and 2 for demonstration purposes. Check for correlations by evaluating the autocorrelation function or with techniques like jackknife or binning.

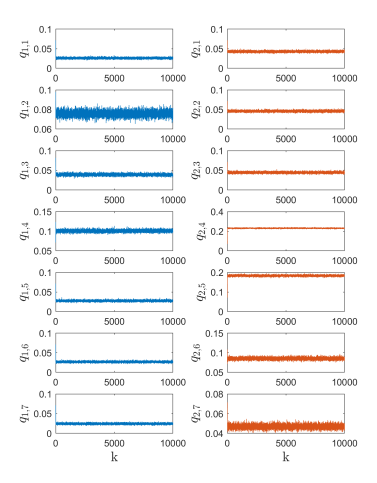


Figure 1: Time series of $q^{(2)}$.

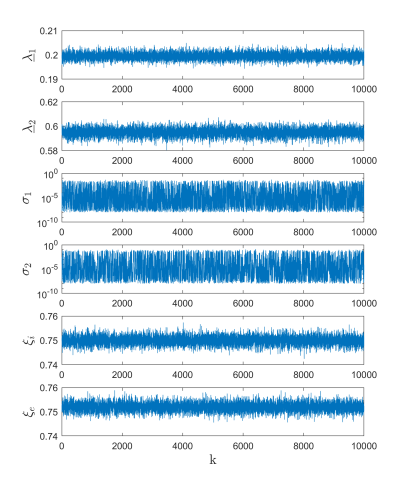


Figure 2: Time series of π . The real values are $\lambda_1 = 0.2$, $\lambda_2 = 0.6$, $\sigma_1 = 0$, $\sigma_2 = 0$, $\xi_i = 0.75$, $\xi_e = 0.75$. The algorithm can only evaluate that σ_j is very small but cant pin it down because in this regime it has no influence.

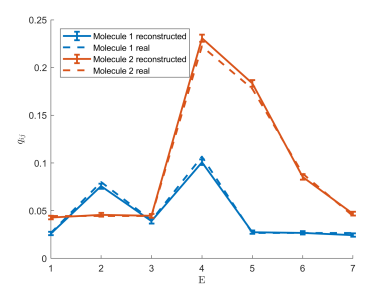


Figure 3: Reconstructed spectrum $q^{(2)}$.

5 Troubleshooting

5.1 Reconstruction is not converged

If the result is not converged the last entry of the time series of $q^{(1)}$, $q^{(2)}$ and π can be used as start parameter for a new computation. Extend the Monte Carlo chain until the desired results are converged.

5.2 C file in MATLAB

In the downloaded folder are pre-compiled c files (tested with Windows7, Windows10 and Debian 8.9 - MATLAB version 2013a, 2016b). If the pre-compiled files dont work see https://de.mathworks.com/help/matlab/matlab_external/what-you-need-to-build-mex-files.html, follow the instructions and compile the c file on your computer.

References

- [1] M. Rumetshofer, P. Heim, B. Thaler, W.E. Ernst, M Koch, and W. von der Linden. Analysis of femtosecond time-resolved photoelectron-photoion coincidence measurements applying bayesian probability theory.
- [2] V. Stert, W. Radloff, C.P. Schulz, and I.V. Hertel. Ultrafast photoelectron spectroscopy: Femtosecond pump-probe coincidence detection of ammonia cluster ions and electrons. *The European Physical Journal D-Atomic, Molecular, Optical and Plasma Physics*, 5(1):97–106, 1999.