# Quantum Metrology with Photoelectrons Vol. 3 \*Analysis methodologies\*

**Paul Hockett** 

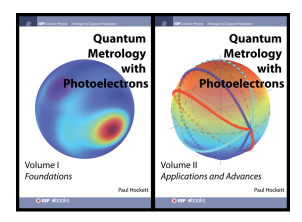
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Quantum Metrology with Photoelectrons Volume 3: *Analysis methodologies*, an open source executable book. This repository contains the source documents (mainly Jupyter Notebooks in Python) and notes for the book, as of Jan 2022 writing is in progress, and the current HTML build can be found online. The book is due to be finished in 2023, and will be published by IOP Press - see below for more details.

#### Series abstract

Photoionization is an interferometric process, in which multiple paths can contribute to the final continuum photoelectron wavefunction. At the simplest level, interferences between different final angular momentum states are manifest in the energy and angle resolved photoelectron spectra: metrology schemes making use of these interferograms are thus phase-sensitive, and provide a powerful route to detailed understanding of photoionization. In these cases, the continuum wavefunction (and underlying scattering dynamics) can be characterised. At a more complex level, such measurements can also provide a powerful probe for other processes of interest, leading to a more general class of quantum metrology built on phase-sensitive photoelectron imaging. Since the turn of the century, the increasing availability of photoelectron imaging experiments, along with the increasing sophistication of experimental techniques, and the availability of computational resources for analysis and numerics, has allowed for significant developments in such photoelectron metrology.

#### About the books



- Volume I covers the core physics of photoionization, including a range of computational examples. The material is presented as both reference and tutorial, and should appeal to readers of all levels. ISBN 978-1-6817-4684-5, http://iopscience.iop.org/book/978-1-6817-4684-5 (IOP Press, 2018)
- Volume II explores applications, and the development of quantum metrology schemes based on photoelectron measurements. The material is more technical, and will appeal more to the specialist reader. ISBN 978-1-6817-4688-3, http://iopscience.iop.org/book/978-1-6817-4688-3 (IOP Press, 2018)

Additional online resources for Vols. I & II can be found on OSF and Github.

Volume III in the series will continue this exploration, with a focus on numerical analysis techniques, forging a closer
link between experimental and theoretical results, and making the methodologies discussed directly accessible via
new software. The book is due for publication by IOP due in 2023; this volume is also open-source, with a live
HTML version at https://phockett.github.io/Quantum-Metrology-with-Photoelectrons-Vol3/ and source available
at https://github.com/phockett/Quantum-Metrology-with-Photoelectrons-Vol3.

For some additional details and motivations (including topical video), see the ePSdata project.

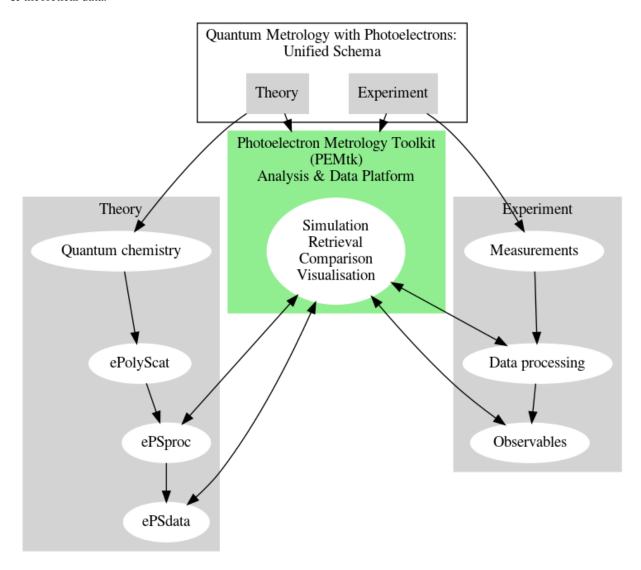
#### **Technical details**

This repository contains:

- doc-source: the source documents (mainly Jupyter Notebooks in Python)
- notes: additional notes for the book,
- the gh-pages branch contains the current HTML build, also available at https://phockett.github.io/ Quantum-Metrology-with-Photoelectrons-Vol3/

The project has been setup to use the Jupyter Book build-chain (which uses Sphinx on the back-end) to generate HTML and Latex outputs for publication from source Jupyter notebooks & markdown files.

The work *within* the book will make use of the Photoelectron Metrology Toolkit platform for working with experimental & theoretical data.



#### **Running code examples**

Each Jupyter notebook (\*.ipynb) can be treated as a stand-alone computational document. These can be run/used/modified independently with an appropriately setup python environment (details to follow).

#### **Building the book**

The full book can also be built from source:

- 1. Clone this repository
- 2. Run pip install -r requirements.txt (it is recommended you do this within a virtual environment)
- 3. (Optional) Edit the books source files located in the doc-source/directory
- 4. Run jupyter-book clean doc-source/ to remove any existing builds
- 5. For an HTML build:
  - Run jupyter-book build doc-source/
  - A fully-rendered HTML version of the book will be built in doc-source/\_build/html/.
- 6. For a LaTex & PDF build:
  - Run jupyter-book build doc-source/ --builder pdflatex
  - A fully-rendered HTML version of the book will be built in doc-source/\_build/latex/.

See https://jupyterbook.org/basics/building/index.html for more information.

#### **Credits**

This project is created using the open source Jupyter Book project and the executablebooks/cookiecutter-jupyter-book template.

To add: build env & main software packages (see automation for this...)



# Part I

# **Frontmatter**

#### **OVERVIEW**

#### 1.1 General overview

Vol. 3. will focus on analysis techniques for quantum metrology with photoelectrons, including:

- Interpreting experimental data.
- Extraction/reconstruction/determination of quantum mechanical properties (matrix elements, wavefunctions, density matrices) from experimental data.
- Comparison of experimental and theoretical data.
- New analysis methodologies & techniques.
- Introduction to newly-developed software platform (see below).

#### 1.2 Provisional contents

#### 1.2.1 Part 1: theory & software

General review & update of the topic, including recent theory developments.

- 1. Introduction
  - a. Topic overview.
  - b. Context of vol. 3 (following vols. 1 & 2).
  - c. Aims: Vol. 3 in the series will continue the exploration of quantum metrology with photoelectrons, with a focus on numerical analysis techniques, forging a closer link between experimental and theoretical results, and making the methodologies discussed directly accessible via a new software platform/ecosystem.
- 2. Quantum metrology software platform/ecosystem overview
  - a. Introduction to python packages for simulation, data analysis, and open-data.
  - b. Photoelectron metrology toolkit (PEMtk) package/platform for experimental data processing & analysis. (See *pemtk.readthedocs.io.*)
  - c. ePSproc package for theory & simulation. (See epsproc.readthedocs.io.)
  - d. ePSdata platform for data/results library (see ePSdata motivations).
- 3. General method development: geometric tensor treatment of photoionization, fitting & matrix-inversion techniques
  - a. Theory development overview tensor methods (e.g. ePSproc tensor methods)

- b. Direct molecular frame reconstruction via matrix-inversion methods (see Gregory, Margaret, Paul Hockett, Albert Stolow, and Varun Makhija. "Towards Molecular Frame Photoelectron Angular Distributions in Polyatomic Molecules from Lab Frame Coherent Rotational Wavepacket Evolution." *Journal of Physics B: Atomic, Molecular and Optical Physics* 54, no. 14 (July 2021): 145601.*DOI: 10.1088/1361-6455/ac135f.*)
- 4. Numerical implementation & analysis platform tools
  - a. Tensor methods implementation in ePSproc/PEMtk.
  - b. Information content analysis (inc. basis-set exploration, e.g. *PEMtk fitting demo*), see also vol. 2, sect. 12.1.
  - c. Density matrix analysis. (e.g. ePSproc density matrix method dev notes)
  - d. Generalised bootstrapping implementation in PEMtk (see vol. 2, sects. 11.3 & 12.3)

#### 1.2.2 Part 2: numerical examples

Open-source worked examples using the new software platform.

- 1. Quantum metrology example: generalised bootstrapping for a homonuclear diatomic scattering system (N2)\*
  - a. Experimental data overview & simulation.
  - b. Matrix element extraction (bootstrap protocol, see vol. 2, sects. 11.3 & 12.3) & statistical analysis.
  - c. Direct molecular frame reconstruction via matrix-inversion methods.
  - d. Comparison of methods.
  - e. Information content/quantum information analysis. (See vol. 2, sect. 12.1.)
- 2. Quantum metrology example: generalised bootstrapping for a heteronuclear scattering system (CO)\*
  - a. Experimental data overview & simulation.
  - b. Matrix element extraction (bootstrap protocol, see vol. 2, sects. 11.3 & 12.3) & statistical analysis.
  - c. Direct molecular frame reconstruction via matrix-inversion methods.
  - d. Comparison of methods.
  - e. Information content/quantum information analysis. (See vol. 2, sect. 12.1.)
- 3. Quantum metrology example: generalised bootstrapping and matrix-inversion methods for a complex/general asymmetric top scattering system (C2H4 (ethylene))\*
  - a. Experimental data overview & simulation.
  - b. Matrix element extraction (bootstrap protocol, see vol. 2, sects. 11.3 & 12.3) & statistical analysis.
  - c. Direct molecular frame reconstruction via matrix-inversion methods.
  - d. Comparison of methods.
  - e. Information content/quantum information analysis.
- 4. Future directions & outlook
- 5. Summary & conclusions
- \* Exact choice of "simple" and "complex" systems may change, but should include a homonuclear diatomic and/or heteronuclear diatomic, and symmetric and asymmetric top polyatomic systems. May also include an atomic example.

# Part II Theory & software

**CHAPTER** 

**TWO** 

#### INTRODUCTION

The overall aim of Vol. 3 is to expand, explore, and illustrate, quantum metrology with photoelectrons: specifically, the application of new python-based tools to tackle problems in matrix element retrieval. The book itself is written as a set of Jupyter Notebooks, hence all the material herein is available directly to readers, and can be run locally to further explore the topic, or adapt the methodology to new problems

Whilst this volume aims to provide a self-contained text, and computational examples which may be used without extensive background knowledge, a brief introduction to the core physics and some recent extensions is presented herein. The unfamiliar reader is referred to Volume 1 of the series for a more detailed presentation, and gateway to the literature [1]. Following the topical introduction, the remainder of Part I introduces the main computational and software tools, recent theory developments, and concludes with a general overview for approaching matrix element retrieval numerically.

Part II details the application of these tools to a few specific cases, starting with a (relatively) simple homonuclear diatomic example, then escalating to a polyatomic asymetric top case.

### 2.1 Topical introduction

#### 2.2 Context & aims for Vol. 3

As noted previously, Vol. 3 is somewhat distinct from the previous volumes in the series; although involving computational elements, Vols. 1 & 2 [1, 2] are more traditional publications. The material presented in this volume aims to continue the exploration of quantum metrology with photoelectrons, with a focus on numerical analysis techniques, forging a closer link between experimental and theoretical results, and making the methodologies discussed directly accessible via a new software platform/ecosystem. In order to fulfil this aim, Vol. 3 is a computational/computable document, with code directly available to readers. Each chapter or section is composed of a Jupyter Notebook (.ipynb), each of which can be modified and used independently.

To facilitate code transparency and reuse, the book is available via a Github repository, Quantum Metrology Vol. 3. An HTML version is also available, which includes interactive figures. A full introduction to the relevant tool-chain, including installation instructions, can be found in Chapter 3: *Quantum metrology software platform/ecosystem overview*.

Quantum Metrology with Photoelectrons Vol. 3 *Analysis methodologies*				

# QUANTUM METROLOGY SOFTWARE PLATFORM/ECOSYSTEM OVERVIEW

#### **STUB**

In recent years, a unified Python codebase/ecosystem/platform has been in development to tackle various aspects of photoionization problems, including *ab initio* computations and experimental data handling, and (generalised) matrix element retrieval methods. The eponymous *Quantum Metrology with Photoelectrons* platform is introduced here, and is used for the analysis herein. The main aim of the platform is to provide a unifying data platform, and analysis routines, for photoelectron metrology, including new methods and tools, as well as a unifying bridge between these and existing tools. Fig. 3.1 provides a general overview of some of the main tools and tasks/layers.

As of late 2022, the new parts of the platform - primarily the Photoelectron Metrology Toolkit [3] library - implement general data handling (although not a full experimental analysis toolchain), matrix element handling and retrieval, which will be the main topic of this volume. In the future, it is hoped that the platform will be extended to other theoretical and experimental methods, including full experimental data handling.

## 3.1 Analysis components

The two main components of the platform for analysis tasks, as used herein, are:

- The Photoelectron Metrology Toolkit [3] (PEMtk) codebase aims to provide various general data handling routines for photoionization problems. At the time of writing, simulation of observables and fitting routines are implemented, along with some basic utility functions. Much of this is detailed herein, and more technical details and ongoing documentation case be found in the PEMtk documentation [4].
- The ePSproc codebase [5, 6, 7] aims to provide methods for post-processing with *ab initio* radial dipole matrix elements from ePolyScat (ePS) [8, 9, 10, 11], or equivalent matrix elements from other sources (dedicated support for R-matrix results from the RMT suite [12, 13] is in development). The core functionality includes the computation of AF and MF observables. Manual computation without known matrix elements is also possible, e.g. for investigating limiting cases, or data analysis and fitting hence these routines also provide the backend functionality for PEMtk fitting routines. Again more technical details can be found in the ePSproc documentation [7].

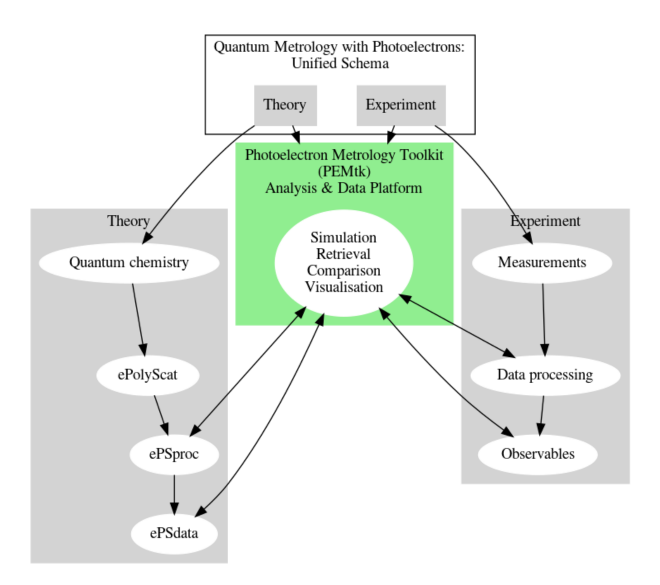


Fig. 3.1: Quantum metrology with photoelectrons ecosystem overview.

#### 3.2 Additional tools

Other tools listed in Fig. 3.1 include:

- Quantum chemistry layer. The starting point for *ab initio* computations. For the examples herein, all computations made use of Gamess ("The General Atomic and Molecular Electronic Structure System") [14, 15] for electronic structure computations, and inputs to ePolyScat.
- ePolyScat (ePS) [8, 9, 10, 11] is an open-source tool for numerical computation of electron-molecule scattering & photoionization by Lucchese & coworkers. All matrix elements used herein were obtained via ePS calculations. For more details see ePolyScat website and manual [11] and Refs. [8, 9, 10].
- ePSdata [16] is an open-data/open-science collection of ePS + ePSproc results.
  - ePSdata collects ePS datasets, post-processed via ePSproc (Python) in Jupyter notebooks, for a full opendata/open-science transparent pipeline.
  - Source notebooks are available on the ePSdata [16] Github project repository, and notebooks + datasets via ePSdata Zenodo [17]. Each notebook + dataset is given a Zenodo DOI for full traceability, and notebooks are versioned on Github.
  - Note: ePSdata may also be linked or mirrored on the existing ePolyScat Collected Results OSF project, but will effectively supercede those pages.
  - All results are released under Creative Commons Attribution-NonCommercial-ShareAlike 4.0 (CC BY-NC-SA 4.0) license, and are part of an ongoing Open Science initiative.

## 3.3 Docker deployments

A Docker-based distribution of various codes for tackling photoionization problems is also available from the Open Photoionization Docker Stacks [18] project, which aims to make a range of these tools more accessible to interested researchers, and fully cross-platform/portable. The project currently includes Docker builds for ePS, ePSproc and PEMtk.

#### 3.4 General discussion

Note that, at the time of writing, rotational wavepacket simulation is not yet implemented in the PEMtk suite, and these must be obtained via other codes. An intial build of the limapack suite for rotational wavepacket simulations is currently part of the Open Photoionization Docker Stacks [18], but has yet to be tested.

3.2. Additional tools



**CHAPTER** 

#### **FOUR**

#### THEORY

- · Observables: photoelectron flux in the LF and MF
- Photoionization dynamics
- Tensor formulation of photoionization
- Information content

## 4.1 Observables: photoelectron flux in the LF and MF

The observables of interest - the photoelectron flux as a function of energy, ejection angle, and time - can be written quite generally as an expansion in spherical harmonics:

$$\bar{I}(\epsilon, t, \theta, \phi) = \sum_{L=0}^{2n} \sum_{M=-L}^{L} \bar{\beta}_{L,M}(\epsilon, t) Y_{L,M}(\theta, \phi) \tag{4.1}$$

Here the flux in the laboratory frame (LF) or aligned frame (AF) is denoted  $I(\epsilon,t,\theta,\phi)$ , with the bar signifying ensemble averaging, and the molecular frame flux by  $I(\epsilon,t,\theta,\phi)$ . Similarly, the expansion parameters  $\bar{\beta}_{L,M}(\epsilon,t)$  include a bar for the LF/AF case. These observables are generally termed photoelectron angular distributions (PADs), often with a prefix denoting the reference frame, e.g. LFPADs, MFPADs, and the associated expansion parameters  $\bar{\beta}_{L,M}(\epsilon,t)$  are generically termed "anisotropy" parameters. The polar coordinate system  $(\theta,\phi)$  is referenced to an experimentally-defined axis in the LF/AF case (usually defined by the laser polarization), and the molecular symmetry axis in the MF. Some arbitrary examples are given in Fig. 4.1, which illustrates both a range of distributions of increasing complexity, and some basic code to set  $\beta_{L,M}$  parameters and visualise them; the values used as tabulated in Fig. 4.2.

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```
# And GLUE for display later with caption
# from myst_nb import glue
# glue("padExamplePlot", figObj[0], display=False);
# Glue with Plotly wrapper.
# gluePlotly("padExamplePlot", figObj[0]) # Working in Render test notebook, but
not here? Issue with subplots?

# Test in separate cell...
gluePlotly("padExamplePlot", figObj[0]) # Working in Render test notebook, but not
where? Issue with subplots?
```

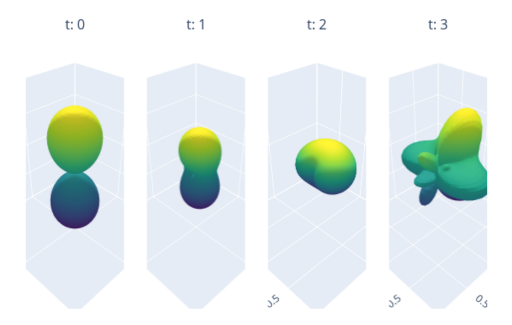


Fig. 4.1: Examples of angular distributions (expansions in spherical harmonics  $Y_{L,M}$ ), for a range of cases. Note that up-down asymmetry is associated with odd-l contributions, and breaking of cylindrical symmetry with  $m \neq 0$  terms.

```
t 0 1 2 3
1 m
0 0 1.0 1.0 1.0 1.0
1 0 0.0 0.5 0.8 1.0
2 0 1.0 0.5 0.0 0.0
4 -2 0.0 0.0 0.0 0.5
2 0.0 0.0 0.0 0.5
```

Fig. 4.2: Values used for the plots in Fig. 4.1

In general, the spherical harmonic rank and order (L,M) of Eq. (4.1) are constrained by experimental factors in the LF/AF, and n is effectively limited by the molecular alignment (which is correlated with the photon-order for gas phase experiments, or conservation of angular momentum in the LF more generally [19]), but in the MF is defined by the maximum continuum angular momentum  $n = l_{max}$  imparted by the scattering event [20].

For basic cases these limits may be low: for instance, a simple 1-photon photoionization event (n=1) from an isotropic ensemble (zero net ensemble angular momentum) defines  $L_{max}=2$ ; for cylindrically symmetric cases (i.e.  $D_{\infty h}$  symmetry) M=0 only. For MF cases,  $l_{max}=4$  is often given as a reasonable rule-of-thumb for the continuum - hence  $L_{max}=8$  - although in practice higher-l may be populated. Some realistic example cases are discussed later (**PART II**), see also ref. [1] for more discussion and complex examples.

In general, these observables may also be dependent on various other parameters; in Eq. (4.1) two such parameters,  $(\epsilon, t)$ , are included, as the usual variables of interest. Usually  $\epsilon$  denotes the photoelectron energy, and t is used in the case of time-dependent (usually pump-probe) measurements. As discussed below (Sect. 4.2), the origin of such dependencies may be complicated but, in general, the associated photoionization matrix elements are energy-dependent, and time-dependence may also appear for a number of intrinsic or extrinsic (experimental) reasons, e.g. electronic or nuclear dynamics, rotational (alignment) dynamics, electric field dynamics etc. In many cases only one particular aspect may be of interest, so t can be used as a generic label to index changes as per Fig. 4.1.

#### 4.1.1 Symmetrized harmonics

Symmetrized (or generalised) harmonics, which essentially provide correctly symmetrized expansions of spherical harmonics  $(Y_{lm})$  functions for a given irreducible representation,  $\Gamma$ , can be defined by linear combinations of spherical harmonics (Refs. [21, 22, 23] as below):

$$X_{hl}^{\Gamma\mu*}(\theta,\phi) = \sum_{\lambda} b_{hl\lambda}^{\Gamma\mu} Y_{l,\lambda}(\theta,\phi) \tag{4.1} \label{eq:4.1}$$

where:

- $\Gamma$  is an irreducible representation,
- $(l, \lambda)$  define the usual spherical harmonic indicies (rank, order)
- $b_{hI\lambda}^{\Gamma\mu}$  are symmetrization coefficients,
- index μ allows for indexing of degenerate components,
- h indexs cases where multiple components are required with all other quantum numbers identical.

The exact form of these coefficients will depend on the point-group of the system, see, e.g. Refs. [23, 24]. Numerical routines for the generation of symmetrized harmonics are implemented in PEMtk: point-groups, character table generation and symmetrization (computing  $b_{hl\lambda}^{\Gamma\mu}$  parameters) is handled by libmsym [25, 26]; additional handling also makes use of pySHtools [27, 28]. A brief example is given below, see the PEMtk documentation [4] for more details.

```
# Import class
from pemtk.sym.symHarm import symHarm

# Compute hamronics for Td, lmax=4
sym = 'Td'
lmax=4

symObj = symHarm(sym,lmax)

# Character tables can be displayed
symObj.printCharacterTable()

# Glue items for later
```

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```
glue("symHarmPG2", f"${sym}$", display=False)
glue("symHarmPG", sym, display=False)
glue("symHarmLmax", lmax, display=False)
```

```
# The full set of expansion parameters can be tabulated
# pd.set_option('display.max_rows', 1)
symObj.displayXlm() # Display values (note this defaults to REAL harmonics)
# symObj.displayXlm(YlmType='comp') # Display values for COMPLEX harmonic expansion.
```

```
# To plot using ePSproc/PEMtk class, these values can be converted to ePSproc BLM_
⇔data type...
# Run conversion - the default is to set the coeffs to the 'BLM' data type
symObj.toePSproc()
# Set to new key in data class
data.data['symHarm'] = {}
for dataType in ['BLM']: #['matE', 'BLM']:
   data.data['symHarm'][dataType] = symObj.coeffs[dataType]['b (comp)'] # Select_
→expansion in complex harmonics
    data.data['symHarm'][dataType].attrs = symObj.coeffs[dataType].attrs
# Plot full harmonics expansions, plots by symmetry
# Note 'squeeze=True' to force drop of singleton dims may be required.
# data.padPlot(keys='symHarm',dataType='BLM', facetDims = ['Cont'], squeeze = True, ...
⇔backend=plotBackend)
data.padPlot(keys='symHarm',dataType='BLM', facetDims = ['Cont'], squeeze = True, _
abackend=plotBackend, plotFlag=False, returnFlag=True) # Working
figObj = data.data['symHarm']['plots']['BLM']['polar'][0]
# And GLUE for display later with caption
# from myst_nb import glue
# glue("padExamplePlot2", figObj, display=False);
gluePlotly("symHarmPADs", figObj)
```

## 4.2 Photoionization dynamics

The core physics of photoionization has been covered extensively in the literature, and only a very brief overview is provided here with sufficient detail to introduce the metrology/reconstruction/retrieval problem; the reader is referred to Vol. 1 [1] (and refs. therein) for further details and general discussion.

Photoionization can be described by the coupling of an initial state of the system to a particular final state (photoion(s) plus free photoelectron(s)), coupled by an electric field/photon. Very generically, this can be written as a matrix element  $\langle \Psi_i | \hat{\Gamma}(\mathbf{E}) | \Psi_f \rangle$ , where  $\hat{\Gamma}(\mathbf{E})$  defines the light-matter coupling operator (depending on the electric field  $\mathbf{E}$ ), and  $\Psi_i$ ,  $\Psi_f$  the total wavefunctions of the initial and final states respectively.

There are many flavours of this fundamental light-matter interaction, depending on system and coupling. For metrology, the focus is currently on the simplest case of single-photon absorption, in the weak field (or purturbative), dipolar regime, resulting in a single photoelectron. (For more discussion of various approximations in photoionzation, see Refs. [29, 30].) In this case the core physics is well defined, and tractable (albeit non-trivial), via the separation of matrix elements into radial (energy) and angular-momentum (geometric) terms pertaining to couplings between various elements of the

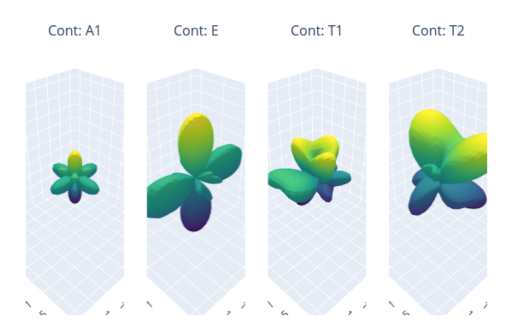


Fig. 4.3: Examples of angular distributions from expansions in symmetrized harmonics  $X_{hl}^{\Gamma\mu*}(\theta,\phi)$ , for all irreducible representations in Td symmetry  $(l_{max}$  =4)

problem; the retrieval of such matrix elements is a well-defined problem, making use of analytic terms in combination with fitting methodologies as explored herein. Again, more extensive background and discussion can be found in *Quantum Metrology* Vol. 1 [1], and references therein. [Add some more refs here?]

The basic case also provides a strong foundation for extension into more complex light-matter interactions, in particular cases with shaped laser-fields (i.e. a time-dependent coupling  $\hat{\Gamma}(\mathbf{E},\mathbf{t})$ ) and multi-photon processes (which require multiple matrix elements). Note, however, that non-perturbative (strong field) light-matter interactions are, typically, not amenable to description in a separable picture in this manner. In such cases the laser field, molecular and continuum properties are strongly coupled, and are typically treated numerically in a fully time-dependent manner (although some separation of terms may work in some cases).

Underlying the photoelectron observables is the photoelectron continuum state  $|\mathbf{k}\rangle$ , prepared via photoionization. The photoelectron momentum vector is denoted generally by  $\mathbf{k}=k\mathbf{\hat{k}}$ , in the MF. The ionization matrix elements associated with this transition provide the set of quantum amplitudes completely defining the final continuum scattering state,

$$\left|\Psi_{f}\right\rangle = \sum \int \left|\Psi_{+};\mathbf{k}\right\rangle \left\langle\Psi_{+};\mathbf{k}|\Psi_{f}\right\rangle d\mathbf{k},\tag{4.2}$$

where the sum is over states of the molecular ion  $|\Psi_{+}\rangle$ . The number of ionic states accessed depends on the nature of the ionizing pulse and interaction. For the dipolar case,

$$\hat{\Gamma}(\mathbf{E}) = \hat{\mu}.\mathbf{E} \tag{4.3}$$

Hence,

$$\left\langle \Psi_{+};\mathbf{k}|\Psi_{f}\right\rangle =\left\langle \Psi_{+};\,\mathbf{k}|\hat{\mu}.\mathbf{E}|\Psi_{i}\right\rangle \tag{4.4}$$

Where the notation implies a perturbative photoionization event from an initial state i to a particular ion plus electron state following absorption of a photon  $h\nu$ ,  $|\Psi_i\rangle + h\nu \rightarrow |\Psi_+; \mathbf{k}\rangle$ , and  $\hat{\mu}.\mathbf{E}$  is the usual dipole interaction term [31], which includes a sum over all electrons s defined in position space as  $\mathbf{r}_{\mathbf{c}}$ :

$$\hat{\mu} = -e\sum_{s} \mathbf{r_s} \tag{4.5}$$

The position space photoelectron wavefunction is typically expressed in the "partial wave" basis, expanded as (asymptotic) continuum eignstates of orbital angular momentum, with angular momentum components (l,m) (note lower case notation for the partial wave components, distinct from upper-case for the similar terms (L,M) in the observables),

$$\Psi_{\mathbf{k}}(r) \equiv \langle r | \mathbf{k} \rangle = \sum_{lm} Y_{lm}(\hat{\mathbf{k}}) \psi_{lm}(r, k)$$
(4.6)

where r are MF electronic coordinates and  $Y_{lm}(\mathbf{\hat{k}})$  are the spherical harmonics.

Similarly, the ionization dipole matrix elements can be separated generally into radial (energy-dependent or 'dynamical' terms) and geometric (angular momentum) parts (this separation is essentially the Wigner-Eckart Theorem, see Ref. [32] for general discussion), and written generally as (using notation similar to [33]):

$$\langle \Psi_{+}; \mathbf{k} | \hat{\mu}.\mathbf{E} | \Psi_{i} \rangle = \sum_{lm} \gamma_{l,m} \mathbf{r}_{k,l,m}$$
(4.7)

Provided that the geometric part of the matrix elements  $\gamma_{l,m}$  - which includes the geometric rotations into the LF arising from the dot product in Eq. (4.7) and other angular-momentum coupling terms - are know, knowledge of the so-called radial (or reduced) dipole matrix elements, at a given k thus equates to a full description of the system dynamics (and, hence, the observables).

For the simplest treatment, the radial matrix element can be approximated as a 1-electron integral involving the initial electronic state (orbital), and final continuum photoelectron wavefunction:

$$\mathbf{r}_{k,l,m} = \int \psi_{lm}(r,k)r\Psi_i(r)dr \tag{4.8}$$

As noted above, the geometric terms  $\gamma_{l,m}$  are analytical functions which can be computed for a given case - minimally requiring knowledge of the molecular symmetry and polarization geometry, although other factors may also play a role (see Sect. 4.3.2 for details).

The photoelectron angular distribution (PAD) at a given  $(\epsilon, t)$  can then be determined by the squared projection of  $|\Psi_f\rangle$  onto a specific state  $|\Psi_{\perp}; \mathbf{k}\rangle$ ,

and therefore the amplitudes in Eq. (4.7) also determine the observable anisotropy parameters  $\beta_{L,M}(\epsilon,t)$  (Eqn. (4.1)). (Note that the photoelectron energy  $\epsilon$  and momentum k are used somewhat interchangeably herein, with the former usually preferred in reference to observables.) Note, also, that in the treatment above there is no time-dependence incorporated in the notation; however, a time-dependent treatment readily follows, and may be incorporated either as explicit time-dependent modulations in the expansion of the wavefunctions for a given case, or implicitly in the radial matrix elements. Examples of the former include, e.g. a rotational or vibrational wavepacket, or a time-dependent laser field. The rotational wavepacket case is discussed herein (see Sect. 4.3.2). The radial matrix elements are a sensitive function of molecular geometry and electronic configuration in general, hence may be considered to be responsive to molecular dynamics, although they are formally time-independent in a Born-Oppenheimer basis - for further general discussion and examples see Ref. [34]; discussions of more complex cases with electronic and nuclear dynamics can be found in Refs. [30, 35, 36, 37].

Typically, for reconstruction experiments, a given measurement will be selected to simplify this as much as possible by, e.g., populating only a single ionic state (or states for which the corresponding observables are experimentally energetically-resolvable), and with a bandwidth  $d\mathbf{k}$  which is small enough such that the matrix elements can be assumed constant. Importantly, the angle-resolved observables are sensitive to the magnitudes and (relative) phases of these matrix elements, and can be considered as angular interferograms

## 4.3 Tensor formulation of photoionization

A number of authors have treated MFPADs and related problems [REFS];

herein, a geometric tensor based formalism is developed, which is close in spirit to the treatments given by Underwood and co-workers [37, 38, 39], but further separates various sets of physical parameters into dedicated tensors; this allows for a unified theoretical and numerical treatment, where the latter computes properties as tensor variables which can be further manipulated and investigated. Furthermore, the tensors can readily be converted to a density matrix representation [32, 40], which is more natural for some quantities, and also emphasizes the link to quantum state tomography and other quantum information techniques. Much of the theoretical background, as well as application to aspects of the current problem, can be found in the textbooks of Blum [40] and Zare [32].

Within this treatment, the observables can be defined in a series of simplified forms, emphasizing the quantities of interest for a given problem. Some details are defined in the following subsections,

#### 4.3.1 Channel functions

A simple form of the equations, amenable to fitting, is to write the observables in terms of "channel functions", which define the ionization continuum for a given case and set of parameters u (e.g. defined for the MF, or defined for a specific experimental configuration),

$$\beta_{L,M}^{u} = \sum_{\zeta,\zeta'} \Upsilon_{L,M}^{u,\zeta\zeta'} \mathbb{I}^{\zeta\zeta'}$$

$$\tag{4.9}$$

Where  $\zeta, \zeta'$  collect all the required quantum numbers, and define all (coherent) pairs of components. The term  $\mathbb{I}^{\zeta\zeta'}$  denotes the coherent square of the ionization matrix elements:

$$\mathbb{I}^{\zeta,\zeta} = I^{\zeta}(\epsilon)I^{\zeta'*}(\epsilon) \tag{4.10}$$

This is effectively a convolution equation (cf. refs. [38, 41]) with channel functions, for a given "experiment" u, summed over all terms  $\zeta$ ,  $\zeta'$ . Aside from the change in notation (which is here chosen to match the formalism of Refs. [8, 9, 10]),

these matrix elements are essentially identical to the simplified (radial) forms  $\mathbf{r}_{k,l,m}$  defined in Eqn. (4.7), in the case where  $\zeta = k, l, m$ . These complex matrix elements can also be equivalently defined in a magnitude, phase form:

$$I^{\zeta}(\epsilon) \equiv \mathbf{r}_{\zeta} \equiv r_{\zeta} e^{i\phi_{\zeta}} \tag{4.11}$$

This tensorial form is numerically implemented in the ePSproc [6] codebase, and is in contradistinction to standard numerical routines in which the requisite terms are usually computed from vectorial and/or nested summations, which can be somewhat opaque to detailed interpretation, and typically implement the full computation of the observables in one monolithic computational routine. The Photoelectron Metrology Toolkit [3] codebase implements matrix element retrieval based on the tensor formalism, with pre-computation of all the geometric tensor components (channel functions) prior to a fitting protocol for matrix element analysis, essentially a fit to Eqn. (4.9), with terms  $I^{\zeta}(\epsilon)$  as the unknowns (in magnitude, phase form per (4.11)). The main computational cost of a tensor-based approach is that more RAM is required to store the full set of tensor variables; however, the method is computationally efficient since it is inherently parallel (as compared to a traditional, serial loop-based solution), hence may lead to significantly faster evaluation of observables. Furthermore, the method allows for the computational routines to match the formalism quite closely, and investigation of the properties of the channel functions for a given problem in general terms, as well as for specific experimental cases.

TODO: numerical examples here or below. TODO: benchmarks, or link to them (see test fitting notebooks...?).

#### 4.3.2 Full tensor expansion

In more detail, the channel functions can be given as a set of tensors, defining each aspect of the problem.

For the MF:

$$\begin{split} \beta_{L,-M}^{\mu_{i},\mu_{f}}(\epsilon) &= & (-1)^{M} \sum_{P,R',R} (2P+1)^{\frac{1}{2}} E_{P-R}(\hat{e};\mu_{0}) \\ &\times & \sum_{l,m,\mu} \sum_{l',m',\mu'} (-1)^{(\mu'-\mu_{0})} \Lambda_{R',R}(R_{\hat{n}};\mu,P,R,R') B_{L,-M}(l,l',m,m') \\ &\times & I_{l,m,\mu}^{p_{i}\mu_{i},p_{f}\mu_{f}}(\epsilon) I_{l',m',\mu'}^{p_{i}\mu_{i},p_{f}\mu_{f}*}(\epsilon) \end{split} \tag{4.12}$$

And the LF/AF as:

$$\bar{\beta}_{L,-M}^{\mu_{i},\mu_{f}}(E,t) = \frac{(-1)^{M} \sum_{P,R',R} [P]^{\frac{1}{2}} E_{P-R}(\hat{e};\mu_{0})}{\sum_{l,m,\mu} \sum_{l',m',\mu'} (-1)^{(\mu'-\mu_{0})} \Lambda_{R'}(\mu,P,R') B_{L,S-R'}(l,l',m,m')} \times I_{l,m,\mu}^{p_{i}\mu_{i},p_{f}\mu_{f}}(\epsilon) I_{l',m',\mu'}^{p_{i}\mu_{i},p_{f}\mu_{f}*}(\epsilon) \sum_{K,Q,S} \Delta_{L,M}(K,Q,S) A_{Q,S}^{K}(t)}$$

$$(4.13)$$

In both cases a set of geometric tensor terms are required,

these terms provide details of:

- $E_{P-R}(\hat{e}; \mu_0)$ : polarization geometry & coupling with the electric field.
- $B_{L,S-R'}(l,l',m,m')$ : geometric coupling of the partial waves into the  $\beta_{L,M}$  terms (spherical tensors).
- $\Lambda_{R'}(\mu, P, R')$ : frame couplings and rotations.
- $\Delta_{L,M}(K,Q,S)$ : alignment frame coupling.
- $A_{O.S}^{K}(t)$ : ensemble alignment described as a set of axis distribution moments (ADMs).

And  $I_{l,m,\mu}^{p_i\mu_i,p_f\mu_f}(\epsilon)$  are the (radial) dipole ionization matrix elements, as a function of energy  $\epsilon$ . These matrix elements are essentially identical to the simplified forms  $r_{k,l,m}$  defined in Eqn. (4.7), except with additional indices to label symmetry and polarization components defined by a set of partial-waves  $\{l,m\}$ , for polarization component  $\mu$  (denoting the photon angular momentum components) and channels (symmetries) labelled by initial and final state indexes  $p_i\mu_i, p_f\mu_f$ . The notation here follows that used by ePolyScat (ePS) [8, 9, 10, 11], and these matrix elements again represent the quantities to be obtained numerically from data analysis, or from an ePolyScat (or similar) calculation.

[Numerical example here, or already included above somewhere]

Note that, in this case as given, time-dependence arises purely from the  $A_{Q,S}^K(t)$  terms in the AF case, and the electric field term currently describes only the photon angular momentum coupling, although can in principle also describe time-dependent/shaped fields. Similarly, a time-dependent initial state (e.g. a vibrational wavepacket) could also describe a time-dependent MF case.

It should be emphasized, however, that the underlying physical quantities are essentially identical in all the theoretical approaches, with a set of coupled angular-momenta defining the geometrical part of the photoionization problem, despite these differences in the details of the theory and notation.

#### 4.3.3 Density matrix representation

The density operator associated with the continuum state in Eq. (4.2) is easily written as  $\hat{\rho} = |\Psi_c\rangle\langle\Psi_c|$ . In the channel function basis, this leads to a density matrix given by the radial matrix elements:

$$\rho^{\zeta\zeta'} = \mathbb{I}^{\zeta,\zeta'} \tag{4.14}$$

Since the matrix elements characterise the scattering event, the density matrix provides an equivalent characterisation of the scattering event.

Further discussion can also be found in the literature, see, e.g., Ref. [40] for general discussion, Ref. [33] for application in pump-probe schemes.

TODO: numerical examples here

#### 4.4 Information content

As discussed in *Quantum Metrology* Vol. 2 [2], the information content of a single observable might be regarded as simply the number of contributing  $\beta_{L,M}$  parameters. In set notation:

$$M = \mathsf{n}\{\beta_{I,M}\}\tag{4.15}$$

where M is the information content of the measurement, defined as  $n\{...\}$  the cardinality (number of elements) of the set of contributing parameters. A set of measurements, made for some experimental variable u, will then have a total information content:

$$M_u = \sum_u \mathbf{n} \{\beta^u_{L,M}\}$$

In the case where a single measurement contains multiple  $\beta_{L,M}$ , e.g. as a function of energy  $\epsilon$  or time t, the information content will naturally be larger:

$$\begin{split} M_u &= & \sum_{u,k,t} \mathbf{n} \{\beta^u_{L,M}(\epsilon,t)\} \\ &= & M_u \times M \end{split}$$

where the second line pertains if each measurement has the same native information content, independent of u. It may be that the variable k is continuous (e.g. photoelectron energy), but in practice it will usually be discretized in some fashion by the measurement.

In terms of purely experimental methodologies, a larger  $M_u$  clearly defines a richer experimental measurement which explores more of the total measurement space spanned by the full set of  $\{\beta_{L,M}^u(k,t)\}$ . However, in this basic definition a larger  $M_u$  does not necessarily indicate a higher information content for quantum retrieval applications. The reason for this is simply down to the complexity of the problem (cf. Eq. (4.9)), in which many couplings define the sensitivity of the observable to the underlying system properties of interest. In this sense, more measurements, and larger M, may only add redundancy, rather than new information.

A more complete accounting of information content would, therefore, also include the channel couplings, i.e. sensitivity/dependence of the observable to a given system property, in some manner. For the case of a time-dependent measurement, arising from a rotational wavepacket, this can be written as:

$$M_u = \mathbf{n}\{\Upsilon^u_{LM}(\epsilon,t)\}$$

In this case, each  $(\epsilon, t)$  is treated as an independent measurement with unique information content, although there may be redundancy as a function of t depending on the nature of the rotational wavepacket and channel functions.

(Note this is in distinction to previously demonstrated cases where the time-dependence was created from a shaped laser-field, and was integrated over in the measurements, which provided a coherently-multiplexed case, see refs. [42, 43, 44] for details.)

# Part III

# **Backmatter**

QUARTER	
CHAPTER	
EIVE	
FIVE	

# **BIBLIOGRAPHY**



Part IV

**Test pages** 

# **BUILD VERSIONS AND CONFIG TESTS**

## 6.1 Versions

```
* sparse not found, sparse matrix forms not available.

* natsort not found, some sorting functions not available.

* Setting plotter defaults with epsproc.basicPlotters.setPlotters(). Run directly—to modify, or change options in local env.

* Set Holoviews with bokeh.

* pyevtk not found, VTK export not available.
```

OMP: Info #271: omp\_set\_nested routine deprecated, please use omp\_set\_max\_active\_ \( \)elevels instead.

```
Date: Wed Nov 23 08:54:19 2022 EST
              OS : Linux
          CPU(s) : 64
        Machine: x86_64
    Architecture : 64bit
            RAM : 62.8 GiB
     Environment : Jupyter
     File system : btrfs
Python 3.9.7 | packaged by conda-forge | (default, Sep 29 2021, 19:20:46)
[GCC 9.4.0]
           pemtk : 0.0.1
         epsproc: 1.3.2-dev
         xarray : 2022.3.0
      jupyterlab: 3.2.1
         plotly : 5.11.0
       holoviews : 1.15.2
           numpy: 1.20.3
           scipy : 1.7.1
         IPython : 7.28.0
      matplotlib : 3.4.3
```

```
scooby: 0.7.0
```

```
!jupyter-book --version
```

```
Jupyter Book : 0.13.1
External ToC : 0.2.4
MyST-Parser : 0.15.2
MyST-NB : 0.13.2
Sphinx Book Theme : 0.3.3
Jupyter-Cache : 0.4.3
NbClient : 0.5.4
```

#### 6.2 Docker build env

To do

# 6.3 Book versions

```
QMpath = '/home/jovyan/QM3'
!git -C {QMpath} branch
!git -C {QMpath} log --format="%H" -n 1
```

```
gh-pages
* main
cd035142e0d76265dbd9e883c520d74253c4d688
```

```
# Check current remote commits
!git ls-remote --heads https://github.com/phockett/Quantum-Metrology-with-
--Photoelectrons-Vol3
```

```
8c82e989a15742b4cfebd54571d03ce49f686a10 refs/heads/gh-pages
4801fde561b70425fa5b364534ab4168e3490bd2 refs/heads/main
```

# 6.4 Github pkg versions

Note - can't get versions for local pip installs from repo(?).

```
from pathlib import Path
import epsproc as ep
ep.__file__
```

```
'/opt/conda/lib/python3.9/site-packages/epsproc/__init__.py'
```

```
# Check current Git commit for local ePSproc version - NOTE THIS ONLY WORKS FOR_
⇔INSTALLED FROM GIT CLONES
# from pathlib import Path
# import epsproc as ep
!git -C {Path(ep.__file__).parent} branch
!git -C {Path(ep.__file__).parent} log --format="%H" -n 1
  fatal: not a git repository (or any of the parent directories): .git
  fatal: not a git repository (or any of the parent directories): .git
# Check current remote commits
!git ls-remote --heads https://github.com/phockett/ePSproc
  92c661789a7d2927f2b53d7266f57de70b3834fa
                                                  refs/heads/dependabot/pip/notes/
   ⇔envs/envs-versioned/mistune-2.0.3
  fe1e9540c7b91fe571f60562acd31d8e489d491e
                                                 refs/heads/dependabot/pip/notes/
   ⇔envs/envs-versioned/nbconvert-6.5.1
  70b80a1e3a54de91c2bfe3b6be82d611fcfd5f43
                                                 refs/heads/dependabot/pip/notes/
   ⇔envs/envs-versioned/pillow-9.3.0
  b8d23a0273a78fd0a24deb6803d8ad048124c501
                                                 refs/heads/dev
  1c0b8fd409648f07c85f4f20628b5ea7627e0c4e
                                                 refs/heads/master
  69cd89ce5bc0ad6d465a4bd8df6fba15d3fd1aee
                                                  refs/heads/numba-tests
  ea30878c842f09d525fbf39fa269fa2302a13b57
                                                  refs/heads/revert-9-master
  baf0be0c962e8ab3c3df57c8f70f0e939f99cbd7
                                                 refs/heads/testDev
# Check current remote commits
!git ls-remote --heads https://github.com/phockett/PEMtk
  bb51d8b601eee2486a5a8757f081549398e56e10
                                                  refs/heads/master
  3f4686dffdbb310f15692f978ba36d6a3d15e8d3
                                                  refs/heads/mfFittingDev
```

#### 6.5 Full conda env

```
!conda list
```

```
# packages in environment at /opt/conda:
                                                Build Channel
# Name
                        Version
_libgcc_mutex
                        0.1
                                         conda_forge conda-forge
_openmp_mutex
                        4 5
                                             1_llvm conda-forge
alabaster
                       0.7.12
                                               pypi_0 pypi
alembic
                       1.7.4
                                         pyhd8ed1ab_0 conda-forge
alsa-lib
                       1.2.3.2
                                          h166bdaf_0 conda-forge
                                     py_1 conda-forge
py39hf3d152e_1 conda-forge
altair
                       4.1.0
ansi2html
                       1.8.0
                                      py39hf3d152e_0 conda-forge
                       3.3.4
anyio
                                         pyh9f0ad1d_0 conda-forge
appdirs
                       1.4.4
                                       py39h3811e60_0 conda-forge
pyhd8ed1ab_0 conda-forge
argon2-cffi
                       21.1.0
                       0.9.27
asteval
                                                                (continues on next page)
```

			(continued from previous page)
astropy	5.0.4	py39hd257fcd_0	conda-forge
async_generator	1.10	py_0	conda-forge
atk-1.0	2.36.0	h3371d22_4	conda-forge
attrs	21.2.0	pyhd8ed1ab_0	conda-forge
babel	2.9.1	pyh44b312d_0	conda-forge
backcall	0.2.0	pyh9f0ad1d_0	conda-forge
backports	1.0	py_2	conda-forge
backports.functools_lru_c		pyhd8ed1ab_	3
beautifulsoup4	4.10.0	pyha770c72_0	conda-forge
blas	2.112	openblas	conda-forge
blas-devel	3.9.0	12_linux64_openbla	
bleach	4.1.0	pyhd8ed1ab_0	conda-forge
blinker	1.4	py_1	conda-forge
blosc	1.21.0	h9c3ff4c_0	conda-forge
bokeh	2.4.1	py39hf3d152e_1	conda-forge
boost-cpp	1.74.0	h312852a_4	conda-forge
bottleneck	1.3.2	py39hce5d2b2_4	conda-forge
brotli	1.0.9	h7f98852_5	conda-forge
brotli-bin	1.0.9	h7f98852_5	conda-forge
brotli-python		py39h5a03fae_7	3
	1.0.9		conda-forge
brotlipy	0.7.0	py39h3811e60_1001	conda-forge
brunsli	0.1	h9c3ff4c_0	conda-forge
bzip2	1.0.8	h7f98852_4	conda-forge
c-ares	1.18.1	h7f98852_0	conda-forge
c-blosc2	2.0.4	h5f21a17_1	conda-forge
ca-certificates	2022.9.24	ha878542_0	conda-forge
cached-property	1.5.2	hd8ed1ab_1	conda-forge
cached_property	1.5.2	pyha770c72_1	conda-forge
cairo	1.16.0	h6cf1ce9_1008	conda-forge
cartopy	0.20.1	py39he7aa91e_2	conda-forge
certifi	2022.9.24	pyhd8ed1ab_0	conda-forge
certipy	0.1.3	py_0	conda-forge
cffi	1.14.6	py39h4bc2ebd_1	conda-forge
cfitsio	4.0.0	h9a35b8e_0	conda-forge
cftime	1.6.0	py39hd257fcd_1	conda-forge
chardet	4.0.0	py39hf3d152e_1	conda-forge
charls	2.2.0	h9c3ff4c_0	conda-forge
charset-normalizer	2.0.0	pyhd8ed1ab_0	conda-forge
chrpath	0.16	h7f98852_1002	conda-forge
click	8.0.3	py39hf3d152e_0	conda-forge
cloudpickle	2.0.0	pyhd8ed1ab_0	conda-forge
colorama	0.4.4	pyh9f0ad1d_0	conda-forge
colorcet	3.0.1	pyhd8ed1ab_0	conda-forge
conda	4.10.3	py39hf3d152e_2	conda-forge
conda-package-handling	1.7.3	py39h3811e60_0	conda-forge
configurable-http-proxy	4.5.0	node15_he6ea98c_0	conda-forge
cryptography	35.0.0		conda-forge
curl	7.79.1	h2574ce0_1	conda-forge
cycler	0.10.0	py_2	conda-forge
cython	0.29.24	py39he80948d_0	conda-forge
cytoolz	0.11.0	py39h3811e60_3	conda-forge
dash	2.7.0	pyhd8ed1ab_0	conda-forge
dask		pyhd8ed1ab_0	conda-forge
	71171 111 11		
	2021.10.0		
dask-core	2021.10.0	pyhd8ed1ab_0	conda-forge

			(continued from previous pa
lebugpy	1.4.1	py39he80948d_0	conda-forge
lecorator	5.1.0	pyhd8ed1ab_0	conda-forge
lefusedxml	0.7.1	pyhd8ed1ab_0	conda-forge
lill	0.3.4	pyhd8ed1ab_0	conda-forge
listributed	2021.10.0	py39hf3d152e_0	conda-forge
locutils	0.17.1	pypi_0	pypi
lucc0	0.23.0	py39h5a03fae_0	conda-forge
entrypoints	0.3	py39hde42818_1002	conda-forge
psproc	1.3.2.dev0	pypi_0	pypi
xpat	2.4.8	h27087fc_0	conda-forge
fmpeg	4.4.0	h6987444_4	conda-forge
ftw	3.3.10	nompi_h77c792f_102	conda-forge
lask	2.1.3	pyhd8ed1ab_0	conda-forge
lask-compress	1.13	pyhd8ed1ab_0	conda-forge
ont-ttf-dejavu-sans-mono	2.37	hab24e00_0	conda-forge
ont-ttf-inconsolata	3.000	h77eed37_0	conda-forge
ont-ttf-source-code-pro	2.038	h77eed37_0	conda-forge
ont-ttf-ubuntu	0.83	hab24e00_0	conda-forge
ontconfig	2.14.0	h8e229c2_0	conda-forge
onts-conda-ecosystem	1	0	conda-forge
onts-conda-forge	1	0	conda-forge
reetype	2.10.4	h0708190_1	conda-forge
reexl	1.0.6	h7f98852_0	conda-forge
ribidi	1.0.10	h36c2ea0_0	conda-forge
sspec	2021.10.1	pyhd8ed1ab_0	conda-forge
uture	0.18.2	pyhd8ed1ab_6	conda-forge
dal	3.3.3	py39h0494519_2	conda-forge
dk-pixbuf	2.42.6	h04a7f16_0	conda-forge
eos	3.10.0	h9c3ff4c_0	conda-forge
eotiff	1.7.0	hcfb7246_3	conda-forge
ettext	0.19.8.1	h73d1719_1008	conda-forge
hostscript	9.54.0	h27087fc_2	conda-forge
			_
hp-import	2.1.0	pypi_0	pypi
iflib	5.2.1	h36c2ea0_2	conda-forge
itdb	4.0.9	pypi_0	pypi
itpython	3.1.29	pypi_0	pypi
lib	2.70.2	h780b84a_4	conda-forge
lib-tools	2.70.2	h780b84a_4	conda-forge
mp	6.2.1	h58526e2_0	conda-forge
mpy2	2.1.0b5	py39h78fa15d_0	conda-forge
mt	6.2.0	h7e416ca_3	conda-forge
nuplot	5.4.3	hedd7fda_0	conda-forge
nutls	3.6.13	h85f3911_1	conda-forge
raphicsmagick	1.3.37	hdc87540_0	conda-forge
raphite2	1.3.13	h58526e2_1001	conda-forge
reenlet	1.1.2	py39he80948d_0	conda-forge
shhg-gmt	2.3.7	ha770c72_1003	conda-forge
st-plugins-base	1.18.5	hf529b03_3	conda-forge
streamer	1.18.5	h9f60fe5_3	conda-forge
tk2	2.24.33	h539f30e_1	conda-forge
5ру	3.4.0	nompi_py39h7e08c79_	_101 conda-forge
arfbuzz	3.1.1	h83ec7ef_0	conda-forge
df4	4.2.15	h10796ff_3	conda-forge
	1.12.1	nompi_h2750804_101	conda-forge
df5			_
df5 eapdict	1.0.1	ру_0	conda-forge

			(continued from previous p
nvplot	0.8.1	py_0	pyviz
icu	68.2	h9c3ff4c_0	conda-forge
idna	3.1	pyhd3deb0d_0	conda-forge
imagecodecs	2021.8.26	py39h571908b_2	conda-forge
imageio	2.9.0	py_0	conda-forge
imagesize	1.4.1	pypi_0	pypi
importlib-metadata	4.8.1	py39hf3d152e_0	conda-forge
importlib_metadata	4.8.1	hd8ed1ab_1	conda-forge
importlib_resources	5.3.0	pyhd8ed1ab_0	conda-forge
ipykernel	6.4.2	py39hef51801_0	conda-forge
ipympl	0.8.2	pyhd8ed1ab_0	conda-forge
ipython	7.28.0	py39hef51801_0	conda-forge
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python-json-logger			_
python-kaleido	0.2.1	pyhd8ed1ab_0	conda-forge
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### **BIBLIOGRAPHY**

- [1] Paul Hockett. *Quantum Metrology with Photoelectrons, Volume 1: Foundations.* IOP Publishing, 2018. ISBN 978-1-68174-684-5. doi:10.1088/978-1-6817-4684-5.
- [2] Paul Hockett. *Quantum Metrology with Photoelectrons, Volume 2: Applications and Advances.* IOP Publishing, 2018. ISBN 978-1-68174-688-3. doi:10.1088/978-1-6817-4688-3.
- [3] Paul Hockett. Photoelectron Metrology Toolkit (PEMtk) Github repository. 2021. URL: https://github.com/phockett/PEMtk (visited on 2022-02-18).
- [4] Paul Hockett. PEMtk the Photoelectron Metrology Toolkit documentation. 2021. URL: https://pemtk.readthedocs.io (visited on 2022-02-18).
- [5] Paul Hockett. ePSproc: Post-processing suite for ePolyScat electron-molecule scattering calculations. *Authorea*, 2016. doi:10.6084/m9.figshare.3545639.
- [6] Paul Hockett. ePSproc: Post-processing for ePolyScat (Github repository). Github, 2016. doi:10.6084/m9.figshare.3545639.
- [7] Paul Hockett. ePSproc: Post-processing for ePolyScat documentation. 2020. URL: https://epsproc.readthedocs.io (visited on 2022-02-18).
- [8] Robert R. Lucchese, Kazuo Takatsuka, and Vincent McKoy. Applications of the Schwinger variational principle to electron-molecule collisions and molecular photoionization. *Physics Reports*, 131(3):147–221, January 1986. doi:10.1016/0370-1573(86)90147-X.
- [9] F. A. Gianturco, R. R. Lucchese, and N. Sanna. Calculation of low-energy elastic cross sections for electron-CF4 scattering. *The Journal of Chemical Physics*, 100(9):6464, May 1994. doi:10.1063/1.467237.
- [10] Alexandra P P Natalense and Robert R Lucchese. Cross section and asymmetry parameter calculation for sulfur 1s photoionization of SF[sub 6]. *The Journal of Chemical Physics*, 111(12):5344, 1999. doi:10.1063/1.479794.
- [11] R R Lucchese. ePolyScat User's Manual. URL: https://epolyscat.droppages.com/ (visited on 2022-04-26).
- [12] Andrew C. Brown, Gregory S. J. Armstrong, Jakub Benda, Daniel D. A. Clarke, Jack Wragg, Kathryn R. Hamilton, Zdeněk Mašín, Jimena D. Gorfinkiel, and Hugo W. van der Hart. RMT: R-matrix with time-dependence. Solving the semi-relativistic, time-dependent Schrödinger equation for general, multielectron atoms and molecules in intense, ultrashort, arbitrarily polarized laser pulses. *Computer Physics Communications*, 250:107062, May 2020. arXiv:1905.06156, doi:10.1016/j.cpc.2019.107062.
- [13] Andrew C. Brown, Gregory S. J. Armstrong, Jakub Benda, Daniel D. A. Clarke, Jack Wragg, Kathryn R. Hamilton, Zdeněk Mašín, Jimena D. Gorfinkiel, and Hugo W. van der Hart. RMT: R-matrix with time-dependence (repository). May 2020. URL: https://gitlab.com/Uk-amor/RMT/rmt (visited on 2022-11-09), arXiv:1905.06156.
- [14] Michael W Schmidt, Kim K Baldridge, Jerry A Boatz, Steven T Elbert, Mark S Gordon, Jan H Jensen, Shiro Koseki, Nikita Matsunaga, Kiet A Nguyen, Shujun Su, Theresa L Windus, Michel Dupuis, and John A Montgomery. General

- atomic and molecular electronic structure system. *Journal of Computational Chemistry*, 14(11):1347–1363, 1993. doi:10.1002/jcc.540141112.
- [15] Mark S. Gordon. Gamess website. URL: http://www.msg.ameslab.gov/gamess/.
- [16] Paul Hockett. ePS data: Photoionization calculations archive. 2019. URL: https://phockett.github.io/ePSdata/ (visited on 2022-02-16).
- [17] Paul Hockett. ePSdata repositories on Zenodo. 2019. URL: https://zenodo.org/search?page=1&size=20&q=hockett&keywords=Data.
- [18] Paul Hockett. Open Photoionization Docker Stacks. URL: https://github.com/phockett/open-photoionization-docker-stacks (visited on 2022-08-04).
- [19] C. Yang. On the Angular Distribution in Nuclear Reactions and Coincidence Measurements. *Physical Review*, 74(7):764–772, October 1948. doi:10.1103/PhysRev.74.764.
- [20] D Dill. Fixed-molecule photoelectron angular distributions. *The Journal of Chemical Physics*, 65(3):1130–1133, 1976. doi:10.1063/1.433187.
- [21] S. L. Altmann and C. J. Bradley. On the Symmetries of Spherical Harmonics. *Philosophical Transactions of the Royal Society A: Mathematical, Physical and Engineering Sciences*, 255(1054):199–215, January 1963. doi:10.1098/rsta.1963.0002.
- [22] SL Altmann and AP Cracknell. Lattice harmonics I. Cubic groups. *Reviews of Modern Physics*, 37(1):19–32, 1965. URL: http://rmp.aps.org/abstract/RMP/v37/i1/p19\_1 (visited on 2013-06-12).
- [23] N Chandra. Photoelectron spectroscopic studies of polyatomic molecules. I. Theory. *Journal of Physics B: Atomic and Molecular Physics*, 20(14):3405–3415, July 1987. doi:10.1088/0022-3700/20/14/013.
- [24] Katharine L. Reid and Ivan Powis. Symmetry considerations in molecular photoionization: Fixed molecule photoelectron angular distributions in C3v molecules as observed in photoelectron–photoion coincidence experiments. *The Journal of Chemical Physics*, 100(2):1066, 1994. doi:10.1063/1.466638.
- [25] Marcus Johansson and Valera Veryazov. Automatic procedure for generating symmetry adapted wavefunctions. *Journal of Cheminformatics*, 9(1):8, February 2017. doi:10.1186/s13321-017-0193-3.
- [26] Marcus Johansson. Libmsym Github. July 2022. URL: https://github.com/mcodev31/libmsym (visited on 2022-08-03).
- [27] Mark A. Wieczorek and Matthias Meschede. SHtools Github. SHTOOLS, August 2022. URL: https://github.com/SHTOOLS/SHTOOLS (visited on 2022-08-03).
- [28] Mark A. Wieczorek and Matthias Meschede. SHTools: Tools for Working with Spherical Harmonics. *Geochemistry*, *Geophysics*, *Geosystems*, 19(8):2574–2592, August 2018. doi:10.1029/2018GC007529.
- [29] Tamar Seideman. Time-resolved photoelectron angular distributions: concepts, applications, and directions. *Annual review of physical chemistry*, 53:41–65, January 2002. doi:10.1146/annurev.physchem.53.082101.130051.
- [30] Tamar Seideman. Time-resolved photoelectron angular distributions as a probe of coupled polyatomic dynamics. *Physical Review A*, 64(4):042504, September 2001. doi:10.1103/PhysRevA.64.042504.
- [31] Christopher Gerry and Peter Knight. Introductory Quantum Optics. Cambridge University Press, 2005.
- [32] Richard N Zare. Angular Momentum: Understanding Spatial Aspects in Chemistry and Physics. John Wiley & Sons, 1988.
- [33] Katharine L. Reid, David J. Leahy, and Richard N. Zare. Effect of breaking cylindrical symmetry on photoelectron angular distributions resulting from resonance-enhanced two-photon ionization. *The Journal of Chemical Physics*, 95(3):1746, 1991. doi:10.1063/1.461023.
- [34] Guorong Wu, Paul Hockett, and Albert Stolow. Time-resolved photoelectron spectroscopy: from wavepackets to observables. *Physical chemistry chemical physics : PCCP*, 13(41):18447–67, November 2011. doi:10.1039/c1cp22031d.

46 Bibliography

- [35] Yasuki Arasaki, Kazuo Takatsuka, Kwanghsi Wang, and Vincent McKoy. Probing wavepacket dynamics with femtosecond energy- and angle-resolved photoelectron spectroscopy. *Journal of Electron Spectroscopy and Related Phenomena*, 108(1-3):89–98, 2000. doi:DOI: 10.1016/S0368-2048(00)00148-1.
- [36] Toshinori Suzuki and Benjamin J. Whitaker. Non-adiabatic effects in chemistry revealed by time-resolved charged-particle imaging. *International Reviews in Physical Chemistry*, 20(3):313–356, July 2001. doi:10.1080/01442350110045046.
- [37] Albert Stolow and Jonathan G. Underwood. Time-Resolved Photoelectron Spectroscopy of Non-Adiabatic Dynamics in Polyatomic Molecules. In Stuart A. Rice, editor, *Advances in Chemical Physics*, volume 139, pages 497–584. John Wiley & Sons, Inc., Hoboken, NJ, USA, March 2008. doi:10.1002/9780470259498.ch6.
- [38] Katharine L. Reid and Jonathan G. Underwood. Extracting molecular axis alignment from photoelectron angular distributions. *The Journal of Chemical Physics*, 112(8):3643, 2000. doi:10.1063/1.480517.
- [39] Jonathan G. Underwood and Katharine L. Reid. Time-resolved photoelectron angular distributions as a probe of intramolecular dynamics: Connecting the molecular frame and the laboratory frame. *The Journal of Chemical Physics*, 113(3):1067, 2000. doi:10.1063/1.481918.
- [40] Karl Blum. Density Matrix Theory and Applications. Volume 64. Springer Berlin Heidelberg, Berlin, Heidelberg, 3rd editio edition, 2012. ISBN 978-3-642-20560-6. doi:10.1007/978-3-642-20561-3.
- [41] Margaret Gregory, Paul Hockett, Albert Stolow, and Varun Makhija. Towards molecular frame photoelectron angular distributions in polyatomic molecules from lab frame coherent rotational wavepacket evolution. *Journal of Physics B: Atomic, Molecular and Optical Physics*, 54(14):145601, July 2021. arXiv:2012.04561, doi:10.1088/1361-6455/ac135f.
- [42] P. Hockett, M. Wollenhaupt, C. Lux, and T. Baumert. Complete Photoionization Experiments via Ultrafast Coherent Control with Polarization Multiplexing. *Physical Review Letters*, 112(22):223001, June 2014. doi:10.1103/PhysRevLett.112.223001.
- [43] Paul Hockett, Matthias Wollenhaupt, Christian Lux, and Thomas Baumert. Complete photoionization experiments via ultrafast coherent control with polarization multiplexing. II. Numerics and analysis methodologies. *Physical Review A*, 92(1):013411, July 2015. doi:10.1103/PhysRevA.92.013411.
- [44] P Hockett, M Wollenhaupt, and T Baumert. Coherent control of photoelectron wavepacket angular interferograms. *Journal of Physics B: Atomic, Molecular and Optical Physics*, 48(21):214004, November 2015. doi:10.1088/0953-4075/48/21/214004.

Bibliography 47