Universitatide BARCELONA

# Mueller matrix polarimetry of anisotropic chiral media 

Oriol Arteaga Barriel



Departament de Física Aplicada i Òptica<br>Grup de Física i Enginyeria de Materials Amorfs i Nanoestructures

# Mueller matrix polarimetry of anisotropic chiral media 

Oriol Arteaga Barriel<br>Memòria presentada per optar al títol de Doctor en Física Tesi dirigida pel Dr. Adolf Canillas i Biosca materials

## Contents

Preface ..... vii
I. Backgrounds ..... 1

1. Light propagation fundamentals. Jones and Mueller descriptions ..... 3
1.1. Light propagation ..... 3
1.2. Plane waves and polarization of light ..... 5
1.3. Jones vectors and Jones matrices ..... 7
1.4. Stokes parameters and Mueller matrices ..... 9
1.4.1. Depolarization ..... 11
1.5. Basic polarization changes induced by transmissive optical elements ..... 12
1.5.1. Absolute phase and intensity changes ..... 13
1.5.2. Polarization changes ..... 14
2. Molecular and supramolecular optical activity ..... 21
2.1. Historical introduction ..... 21
2.2. Circular dichroism and circular birefringence in molecules ..... 23
2.2.1. Cotton effect ..... 25
2.2.2. Excitons ..... 26
2.2.3. J-aggregates ..... 27
2.3. Homochirality ..... 30
2.4. Standard methods to measure optical activity ..... 32
2.5. The concern of anisotropic media in chemistry ..... 33
II. Theory ..... 35
3. Light propagation in anisotropic optical active media ..... 37
3.1. Constitutive Equations. Introduction to Berreman matrix formalism ..... 38
3.2. The lamellar representation ..... 40
3.2.1. Calculation of the Mueller-Jones matrix ..... 44
3.2 .2 . The effect of the interfaces ..... 47
3.2.3. The controversy about Jones birefringence ( $\mathrm{LB}^{\prime}$ ) and Jones dichroism ( $L^{\prime}$ ) ..... 49
3.2.4. Twisted crystal ..... 52
4. Inversion and decomposition of Mueller matrices ..... 57
4.1. Analytic inversion of the Mueller-Jones polarization matrices for homogeneous media ..... 57
4.2. Inversion of a experimental Mueller matrix ..... 59
4.3. Decomposition of Mueller matrices ..... 60
4.3.1. Polar and Lu-Chipman decompositions ..... 60
4.3.2. Pseudopolar decomposition ..... 64
4.3.3. Symmetric decomposition ..... 73
III. Experimental sets ..... 75
5. Two-modulator generalized ellipsometer ..... 77
5.1. Introduction ..... 77
5.2. The photoelastic modulator ..... 78
5.3. Experimental configuration ..... 79
5.3.1. Optical components ..... 79
5.3.2. Electronic hardware ..... 81
5.4. Determination of the intensity with the Stokes-Mueller represen- tation ..... 84
5.5. Data measurement ..... 87
5.6. Calibration ..... 94
5.6.1. Using the calibration results into the measurement ..... 99
6. Quartz assisted two-modulator generalized ellipsometer ..... 101
6.1. Introduction ..... 101
6.2. Description of the quartz-assisted 2-MGE ..... 103
6.3. Effect of quartz plates in the measurement ..... 106
6.4. Calibration ..... 108
6.5. Measurement schemes ..... 110
6.5.1. Spectroscopic mode ..... 110
6.5.2. Spatially resolved mode ..... 113
IV. Experimental measurements ..... 117
7. Optical activity of $\alpha$-quartz ..... 119
7.1. Experimental details of the measurement ..... 120
7.2. Method of measurement ..... 122
7.2.1. A model for the dispersion of the gyration tensor components ..... 124
7.3. Data analysis of the Mueller matrix of quartz ..... 125
7.4. Results ..... 129
8. Chiral induction by hydrodynamic effects ..... 137
8.1. Spectroscopic measurements ..... 138
8.1.1. Porphyrin aggregates ..... 138
8.1.2. Rossette nanotubes ..... 144
8.2. Space-resolved measurements in square section stirred cuvettes ..... 146
8.3. Discussion ..... 148
9. Measurements on heterogenous solid-state samples ..... 155
9.1. Chiral domain in crystallizations ..... 155
9.1.1. Benzil polycrystalline films ..... 156
9.1.2. Other polycrystalline films ..... 158
9.2. Meteorites ..... 161
9.2.1. Motivation ..... 162
9.2.2. Experiment ..... 163
9.2.3. Results ..... 163
10. Conclusions ..... 169
10.1. Theory ..... 169
10.2. Experimental sets ..... 170
10.3. Experiments ..... 171
10.4. Future perspectives ..... 173
V. Appendices ..... 175
A. Resum en català ..... 177
A.1. Introducció ..... 177
A.2. Propagació de la llum en medis òpticament actius i anisòtrops ..... 178
A.3. Inversió i descomposició de les matrius de Mueller ..... 181
A.3.1. Inversió analítica d'un matriu de Mueller-Jones correspo- nent a un medi homogeni ..... 181
A.3.2. Descomposició de les matrius de Mueller ..... 182
A.4. Dispositius experimentals ..... 184
A.4.1. Rotació òptica aplicada a la mesura ..... 185
A.5. Mesures experimentals ..... 187
A.5.1. Quars ..... 187
A.5.2. Induccions quirals per efectes hidrodinàmics ..... 188
A.5.3. Mostres sòlides heterogènies ..... 189
A.5.4. Conclusions ..... 190
B. Computer software ..... 193
Bibliography ..... 203

## Preface

Optical activity is one of the most fascinating phenomena revealed by polarized light. It is observed when chiral materials interact with polarized electromagnetic beams and selectively couple with the left- or right-handed circularly polarized components. From a historic perspective, the study of optical activity has received more attention in chemistry than in any other field of science. Probably this is due to the dramatic consequences that the chirality of molecules or supramolecular structures has in life, as it is understood nowadays from a biochemistry perspective. Traditionally, among physicists, the interest in optical activity and its measurement has been relied to the study of the optical properties of anisotropic crystals. In any case, in applied physics and material science optical activity is usually not the focus of investigations.

Ellipsometry or, more generally, polarimetry are optical characterization techniques centered in the study and interpretation of polarized electromagnetic beams after interacting with media. Optically active materials have always been in the spotlight of these techniques, but only in comparatively few occasions the emphasis has been put on the specific problems that the measurements of optical activity and their interpretation presents. In spite of the steadily growing interest in media of complex response to polarized light, often specialists on polarimetry are not aware of the new challenges that the study of optical activity in supramolecular structures offers, or in the long-term problems that crystallographers have had to measure optical rotation for directions out of the optical axis in crystals. Sometimes their notions about optical activity do not go beyond the very simple and classic example of a solution of sugar molecules in water that rotates an incoming polarized light by a certain angle that depends on the concentration of sugar.

When this thesis started, around five years ago, we had no idea that at the end it would be so much focused in the study of optical activity. At that time I hardly knew what optical activity was, and I could not imagine how much I would enjoy working on it. Our starting point was the long-term experience that our research group had with ellipsometry and with self-built spectroscopic ellipsometers, and our initial plans was to renew them to start working with anisotropic samples. At the moment of writing this preface we realize that the term "Mueller polarimetry" describes better the type of measurements we have employed than the term "ellipsometry", as this later tends to be associated, although not exclusively, to the reflection of light at thin films and interfaces. In spite of this precision in this work he will indistinctly use either "ellipsometry" or "polarimetry" to refer
to the type of measurements we have performed.
This work uses polarimetry to study the phenomenon of interaction of polarized light with chiral samples. The development of polarimetry as a measurement technique and the interpretation of the corresponding data that it offers falls mostly within the domain of optics or electromagnetism. However most of the work we have done has been motivated by the study of problems that nowadays mostly belong to the domain of chemistry. The interaction between aspects from chemitry related to optical activity in molecular or supramolecular systems and aspects from physics related to light polarization will be a constant during this work. Therefore this is a multidisciplinary work, in which the references to questions about molecular optical activity will be very frequent despite having been written under the perspective of physics. Due to the diversity of aspects we have have worked with and to their associated multidisciplinary we are aware that this thesis is neither a good introductory text for newcomers to polarimetry nor a introductory explanation about molecular optical activity. In contrast, we hope that those readers with some background on polarization optics will find here some innovative and nonconventional application of polarimetry, and that readers with some experience in the study optical activity will find here how the Mueller matrix polarimetry can be useful to study complex optical active media.

I also take advantage of this preface to give a personal vision about the historical perspective of the study of the interaction of samples with optical activity with polarized light. During the 19th century this was a subject of feverous scientific investigation because much the progress made in the study and comprehension of light polarization and its interaction with materials came from the study of optical activity. This intense investigation on the subject was kept during the first part of 20th century. However, and although the work with optical active media in polarization optics has never stopped, during the second part of the 20th century and also during the first years of the 21st century the active research on new or in refined methods of measurement has apparently decreased. In contrast, during these decades the measurement of optical activity emerged as a routine characterization technique and was worldwide popularized in chemistry laboratories. As a result, measurements of optical activity became more numerous than ever before, but perhaps not many efforts were put on how to improve them. This can leave the sensation, in my opinion misleading, that everything in this field is already known and there is no place for new discoveries. The specialization in modern science has probably implied that the current researchers that are more interested in optical activity are not specialists in polarized light and communications between different scientific communities are not always fluid. This is probably the reason why, surprisingly for an applied research in the 21st century, to develop the part of the work we have performed with crystals I had to resort many references that were many decades old to find there still unsurpassed reports about measurements of the optical activity of certain crystals. Fortunately,
at the time of finishing writing this thesis, I keep the gratifying feeling that there is still much to be done in this field, and I will be satisfied if at least a part of this sensation is transmitted to the reader.

## Organization

This thesis is structured so as to give a complete account of the measurement of samples with optical activity with Mueller polarimetry, covering aspects that range from theory of light propagation in anisotropic media to details of instrumental design. We have made an effort to make this thesis as self-contained as possible and we have included several review material for this purpose.

The work developed in this thesis can be divided in three big areas: study and interpretation of light propagation though an anisotropic optical active media in terms of the transmissive Mueller matrix, design and construction of an appropriate instrument for the measurements of the Mueller matrix and the experimental results on particular anisotropic samples showing optical activity. We consider that this work introduces some degree of novelty in each of these three aspects and if there is a particular one that suscitate more interest will probably depend on the reader's background. Globally this thesis is divided in four parts and two appendices.

Part I is formed by the two first preliminary chapters. In chapter 1 we introduce some basic concepts about light polarization and light propagation in optical media. In chapter 2 we give a general review on questions related to the measurements of molecular and supramolecular optical activity that serve as a motivation for later chapters.

Part II is devoted to development of the theory that will be later used for the interpretation of experimental data. Chapter 3 focuses on study of light propagation through anisotropic optical active media. The difficulty associated to this study is outlined in this chapter and special emphasis is put in presenting a differential matrix method formalism. Chapter 4 deals with the explicit problem of interpretation of the measured Mueller matrix. Two main alternatives are offered: an analytic inversion method and a decomposition method based on the algebra of Mueller matrices.

Part III is dedicated to the presentation of the experimental technique used in this thesis for the determination of the Mueller matrices. In chapter 5 we introduce the instrument we have built for the spectroscopic measurement of the experimental Mueller matrices, the two-modulator generalized ellipsometer (2-MGE). Aspects about the design of the instrument as well as calibration procedures are presented. Chapter 6 describes a novel method that supposes a further development of the 2-MGE capabilities, because permits it us to obtain measurements with a high lateral resolution. It enables the characterization of samples
which are in-plane heterogeneous.
Part IV includes some of the most significative experimental investigations about optical activity developed in the framework of this thesis. It begins with chapter 7, which presents the spectroscopic measurement of the gyration tensor of quartz crystal. Chapter 8 is dedicated to a series of experiments developed on solutions containing supramolecular structures that show a reversible chiral induction by effect of hydrodynamic flows. In chapter 9 , heterogenous solid state samples are studied using spatially resolved Mueller matrix mappings. Finally, in chapter 10 we summarize the main conclusions of the work.

Appendix A is a Catalan summary of the thesis, while appendix B is an informal revision of the software written for the development of several of the tasks cited in previous chapters.

## Part I.

## Backgrounds

## Chapter 1.

## Light propagation fundamentals. Jones and Mueller descriptions

With the formulation of the electromagnetic theory by James C. Maxwell it was shown that light is an electromagnetic wave with a frequency within a particular range. At that time it was already known from previous works on optics (e.g. Young and Fresnel) that when light propagates through an optical medium shows rather complicated behavior due to refraction and absorption phenomena. The electromagnetic theory has permitted the development of a rigorous theoretical framework for the successful analysis of the optical properties of a wide range of media. Nowadays the interest in this topic has not decreased because this is not a finished research, specially for anisotropic and/or inhomogeneous media.

In this chapter we give a short theoretical background on the propagation of light in a medium and we review some basic topics about polarization optics. This revision focuses on the presentation of Maxwell's equations and the constitutive relations for different type of media and on the different vector representations for polarized light. We finish with a phenomenological description of the various anisotropic polarization-dependent effects that can be present in a medium.

### 1.1. Light propagation

Maxwell's equations in differential form are given by:

$$
\begin{align*}
& \nabla \times \mathbf{E}(\mathbf{r}, t)=-\frac{\partial \mathbf{B}(\mathbf{r}, t)}{\partial t}  \tag{1.1a}\\
& \nabla \times \mathbf{H}(\mathbf{r}, t)=\mathbf{J}(\mathbf{r}, t)+\frac{\partial \mathbf{D}(\mathbf{r}, t)}{\partial t}  \tag{1.1b}\\
& \nabla \cdot \mathbf{D}(\mathbf{r}, t)=\rho(\mathbf{r}, t)  \tag{1.1c}\\
& \nabla \cdot \mathbf{B}(\mathbf{r}, t)=0 \tag{1.1d}
\end{align*}
$$

where $\mathbf{E}$ is the electric field, $\mathbf{H}$ the magnetic field, $\mathbf{D}$ the electric displacement, $\mathbf{B}$ the magnetic flux density, $\mathbf{J}$ the current density and $\rho$ the free charge density.

Relations between the physical quantities appearing in Eqs. (1.1) (i.e. between $\mathbf{E}$ and $\mathbf{D}$ and between $\mathbf{H}$ and $\mathbf{B}$ ) are required to solve the Maxwell equations. They are known as constitutive relations, and they are established by the physical
properties of the medium in which light propagates. Maxwell's equations are generally held to be inviolable and, therefore, the properties of matter enter solely through the constitutive equations. In free space the constitutive relations are:

$$
\begin{align*}
\mathbf{D} & =\varepsilon_{0} \mathbf{E}  \tag{1.2}\\
\mathbf{B} & =\mu_{0} \mathbf{H} \tag{1.3}
\end{align*}
$$

where $\varepsilon_{0}$ and $\mu_{0}$ are the permittivity and the permeability of vacuum, respectively. In general, the constitutive relations in regions occupied by matter have the form

$$
\begin{gather*}
\mathbf{D}=\varepsilon_{0} \mathbf{E}+\mathbf{P},  \tag{1.4}\\
\mathbf{B}=\mu_{0}(\mathbf{H}+\mathbf{M}), \tag{1.5}
\end{gather*}
$$

where $\mathbf{P}$ and $\mathbf{M}$ are, respectively, the electric and the magnetic polarizations and they can be interpreted as the average electric and magnetic dipole moment per unit volume. In a homogeneous linear isotropic dielectric medium, the electric polarization is parallel and proportional to the electric field:

$$
\begin{equation*}
\mathbf{P}=\varepsilon_{0} \chi_{e} \mathbf{E} \tag{1.6a}
\end{equation*}
$$

so that

$$
\begin{equation*}
\mathbf{D}=\varepsilon_{0}\left(1+\chi_{e}\right) \mathbf{E}=\varepsilon \mathbf{E}, \tag{1.6b}
\end{equation*}
$$

where $\chi_{e}$ is the electric susceptibility that is always positive and $\varepsilon$, the permittivity. Similarly in isotropic homogeneous linear magnetic media, the magnetic polarization is parallel and proportional to the magnetic field.

$$
\begin{equation*}
\mathbf{M}=\chi_{m} \mathbf{H} \tag{1.7a}
\end{equation*}
$$

so that

$$
\begin{equation*}
\mathbf{B}=\mu_{0}\left(1+\chi_{m}\right) \mathbf{H}=\mu \mathbf{H} \tag{1.7b}
\end{equation*}
$$

where $\chi_{m}$ is the magnetic susceptibility that can be positive and negative; and $\mu$ is the permeability.

In an anisotropic material, the polarization and the electric field are not necessarily in the same direction. For these materials the electric and magnetic susceptibilities are in general tensors, which means that the permittivity $\varepsilon$ and the permeability $\boldsymbol{\mu}$ are tensors:

$$
\begin{align*}
\mathbf{D} & =\varepsilon \mathbf{E}  \tag{1.8a}\\
\mathbf{B} & =\boldsymbol{\mu} \mathbf{H} \tag{1.8b}
\end{align*}
$$

In this introductory section we will restrict our attention to the study of Maxwell's equations with linear isotropic constitutive relations. [Eqs. (1.6b) and
(1.7b)]. Maxwell's equations (1.1) with these constitutive relations and in a source-free space (i.e. without current or charge densities) can be written as

$$
\begin{align*}
& \nabla \times \mathbf{E}(\mathbf{r}, t)=-\mu \frac{\partial \mathbf{H}(\mathbf{r}, t)}{\partial t},  \tag{1.9a}\\
& \nabla \times \mathbf{H}(\mathbf{r}, t)=\varepsilon \frac{\partial \mathbf{E}(\mathbf{r}, t)}{\partial t},  \tag{1.9b}\\
& \nabla \cdot \mathbf{E}(\mathbf{r}, t)=0  \tag{1.9c}\\
& \nabla \cdot \mathbf{H}(\mathbf{r}, t)=0 \tag{1.9d}
\end{align*}
$$

Taking the curl at both sides of Eq. (1.9a) and substituting Eq. (1.9b) we get

$$
\begin{equation*}
\nabla \times(\nabla \times \mathbf{E})=-\mu \varepsilon \frac{\partial^{2} \mathbf{E}}{\partial t^{2}} \tag{1.10}
\end{equation*}
$$

Applying the identity $\left[\nabla \times \nabla \times=\nabla(\nabla \cdot)-\nabla^{2}\right]$ and using Eq. (1.9c) we obtain the wave equation:

$$
\begin{equation*}
\nabla^{2} \mathbf{E}-\frac{1}{u^{2}} \frac{\partial^{2} \mathbf{E}}{\partial t^{2}}=0 \tag{1.11}
\end{equation*}
$$

where $u=(\mu \varepsilon)^{-1 / 2}$ is the phase velocity of light propagating in the medium [in vacuum $\left.u=\left(\mu_{0} \varepsilon_{0}\right)^{-1 / 2} \equiv c\right] . n=c / u$ is referred to as the index of refraction.

For a monochromatic wave the time variation of the electric field vector is sinusoidal: $\mathbf{E}(\mathbf{r}, t)=E(r) e^{i \omega t}$ where $\omega$ is the angular frequency. Substituting it into Eq. (1.11) and introducing the wavenumber, $k=\omega / u$, we get

$$
\begin{equation*}
\nabla^{2} \mathbf{E}(\mathbf{r})+k^{2} \mathbf{E}(\mathbf{r})=0, \tag{1.12}
\end{equation*}
$$

which is known as the Helmholtz equation.
In this thesis we will focus our attention on chiral anisotropic media. Such media, also called gyrotropic media, exhibit optical activity and the physics underlying their interaction with the electromagnetic fields is more complex than for nongyrotropic systems. If a medium is chiral and anisotropic its constitutive relations are different than those presented in Eqs. (1.8). We discuss light propagation through this type of media in detail in Chapter 3.

### 1.2. Plane waves and polarization of light

A plane wave is a constant-frequency wave whose wavefronts are infinitely parallel planes of constant amplitude normal to the phase velocity vector. A uniform plane wave is a particular solution of Maxwell's equations in a homogeneous region of the space.

Given a plane wave with a direction of propagation $\hat{\mathbf{k}}$, Maxwell's equations impose that the electric and magnetic field should be perpendicular to the direction of propagation and one to each other. Conventionally, when considering
polarization, only the electric field vector is described, since the magnetic field is perpendicular to the electric field and proportional to it. For instance, the electric field vector of lightwave propagating along the z-axis ( $\hat{\mathbf{k}} \equiv \hat{\mathbf{z}}$ ) is

$$
\begin{equation*}
\mathbf{E}(z, t)=\left(E_{x 0} e^{i \delta_{x}} \hat{\mathbf{x}}+E_{y 0} e^{i \delta_{y}} \hat{\mathbf{y}}\right) e^{[i(\omega t-k z)]} \tag{1.13}
\end{equation*}
$$

where the amplitudes $E_{x 0}$ and $E_{y 0}$ are real numbers. The polarization state is given by the relative difference in magnitude and phase between these components. Note that we have chosen the $e^{i \omega t}$ time-dependence instead of $e^{-i \omega t}$. This results in negative imaginary parts of the dielectric functions of all treated materials.

If we consider the electric field at a certain point $z$ as a function of time, Eq. (1.13) is the representation of a ellipse in the plane $x y$ (Fig. 1.1). The parameters that describe the ellipse are the azimuth $\theta \in\left[-\frac{\pi}{2}, \frac{\pi}{2}\right]$ and the ellipticity ${ }^{1}$ $\epsilon \in\left[-\frac{\pi}{4}, \frac{\pi}{4}\right]$. For a vanishing ellipticity the two components of the electric field are in phase and the light wave is linearly polarized. When $\epsilon= \pm \frac{\pi}{4}$ the polarization is circular. When $\epsilon$ does not satisfy any of the above conditions the light wave is elliptically polarized.


Figure 1.1.: The polarization state can be described by two characteristic angles of the polarization ellipse: the azimuth $\theta$, and the ellipticity $\epsilon$.

The handedness of the ellipse of polarization determines the sense in which the ellipse is described. The polarization is right-handed if the field vector rotates

[^0]clockwise when looking against the direction of $\hat{\mathbf{k}}$ (i.e. looking "into the beam" for a travelling wave). Similarly, the polarization is left-handed for a counterclockwise rotation sense.

### 1.3. Jones vectors and Jones matrices

The Jones vector is a direct way to describe polarized light. It was introduced by the American Physicist R. Clark Jones in 1941 [1]. The Jones vector can be built by noting that Eq. (1.13) can be casted in the form of a column vector:

$$
\mathbf{E}(z, t)=\left[\begin{array}{c}
E_{x 0} \exp \left[i\left(\omega t-k z+\delta_{x}\right)\right]  \tag{1.14}\\
E_{y 0} \exp \left[i\left(\omega t-k z+\delta_{y}\right)\right]
\end{array}\right]=\exp [i(\omega t-k z)]\left[\begin{array}{c}
E_{x 0} e^{i \delta_{x}} \\
E_{y 0} e^{i \delta_{y}}
\end{array}\right] .
$$

To express a Jones vector the term $\exp [i(\omega t-k z)]$ is usually omitted and Eq. (1.14) is thus simplified to

$$
\mathbf{E}(z, t)=\left[\begin{array}{l}
E_{x}  \tag{1.15}\\
E_{y}
\end{array}\right]
$$

where $E_{x}$ and $E_{y}$ are complex amplitudes:

$$
\begin{align*}
& E_{x}=E_{x 0} e^{i \delta_{x}}  \tag{1.16a}\\
& E_{y}=E_{y 0} e^{i \delta_{y}} \tag{1.16~b}
\end{align*}
$$

The Jones vector is a direct representation of a single, monochromatic, uniform and transverse-electric plane wave. It contains complete information about the amplitudes and phases of the field components, hence about the polarization of the wave. Jones vectors describe completely polarized light. As an strictly monochromatic wave is always completely polarized, i. e. $E_{x 0}, E_{y 0}, \delta_{x}$ and $\delta_{y}$ are constants, there is always a Jones vector associated to a monochromatic wave. However, real life light beams are, at best, quasi-monochromatic, which means that some slow fluctuations of amplitudes and phases of the fields may be present. A quasi-monochromatic wave can be regarded as a superposition of mutually incoherent monochromatic light beams whose frequencies vary in a narrow bandwidth $\delta \omega$ around a central frequency $\omega_{0}$. A quasi-monochromatic wave behaves like a monochromatic wave for times much shorter than the coherence time of the wave and, within this limit, the description of the polarization states of quasi-monochromatic light in terms of the Jones vectors is valid.

Light intensity can be obtained by pre-multiplying the Jones vector $\mathbf{E}$ by its Hermitian adjoint $\mathbf{E}^{\dagger}{ }^{2}$ :

$$
\begin{equation*}
I=\mathbf{E}^{\dagger} \mathbf{E}=E_{x}^{*} E_{x}+E_{y}^{*} E_{y} \tag{1.17}
\end{equation*}
$$

[^1]The overall effect of the interaction of light with a medium or an optical system can be described by a $2 \times 2$ complex matrix $\mathbf{J}$, referred so as to Jones matrix:

$$
\left[\begin{array}{c}
E_{x}^{\prime}  \tag{1.18}\\
E_{u}^{\prime}
\end{array}\right]=\mathbf{J}\left[\begin{array}{c}
E_{x} \\
E_{y}
\end{array}\right] .
$$

$E_{x, y}$ and $E_{x, y}^{\prime}$ are, respectively, the electric vector components of incident and outgoing light.

The Jones matrix $\mathbf{J}$ is composed of four elements which in general are complex. Therefore, to completely define a Jones matrix 8 independent real parameters are needed. The Jones matrix describing a series of optical elements is equal to the matrix product of the Jones matrix of each element. If we focus our attention on one incident and one emergent plane wave, the overall effect of an entire cascade of $N$ optical elements is described by

$$
\begin{equation*}
\mathbf{E}_{\mathbf{o}}=\mathbf{J}_{\mathbf{N}} \mathbf{J}_{\mathbf{N}-\mathbf{1}} \ldots \mathbf{J}_{\mathbf{I I}} \mathbf{J}_{\mathbf{I}} \mathbf{E}_{\mathbf{i}}=\mathbf{J}_{\mathbf{c o m b}} \mathbf{E}_{\mathbf{i}}, \tag{1.19}
\end{equation*}
$$

where $\mathbf{J}_{\mathbf{c o m b}}=\mathbf{J}_{\mathbf{N}} \mathbf{J}_{\mathbf{N}-\mathbf{1}} \ldots \mathbf{J}_{\mathbf{I I}} \mathbf{J}_{\mathbf{I}}$. The incident plane wave encounters the optical element I first, then element II, etc.

Jones matrices are related to the particular coordinate system used to specify the orientations of the incident and emergent light beams. The effect of rotating both the input and output coordinate system by the same angle $\alpha$ (positive for a counter-clockwise rotation looking against the direction of propagation ${ }^{3}$ ) is to transform the Cartesian Jones matrix of the optical system according to:

$$
\mathbf{J}^{\prime}=\mathbf{R}(\alpha) \mathbf{J R}(-\alpha), \quad \text { where } \quad \mathbf{R}(\alpha)=\left[\begin{array}{cc}
\cos \alpha & \sin \alpha  \tag{1.20}\\
-\sin \alpha & \cos \alpha
\end{array}\right]
$$

In this equation the polarization element remains fixed but the coordinate system rotates by $\alpha$. In the case in which the coordinate system remains fixed but there is a rotation of the polarization element by an angle $\alpha$ such that the angle of incidence is unchanged (for a normal-incidence beam, rotating the element about the normal), the resulting Jones matrix is

$$
\begin{equation*}
\mathbf{J}_{\alpha}=\mathbf{R}(-\alpha) \mathbf{J R}(\alpha) . \tag{1.21}
\end{equation*}
$$

Quasi-monochromatic radiation is not necessarily completely polarized. It can also be unpolarized or partially polarized and, in practice, partially polarized light is frequently found experimentally. The Jones formalism can only handle completely polarized light. A different formalism that can be also applied to partially polarized or unpolarized ration is described in the next section.

[^2]
### 1.4. Stokes parameters and Mueller matrices

The polarization state of light can also be specified by a set of parameters that form the so-called Stokes vector $\mathbf{S}$ :

$$
\mathbf{S}=\left[\begin{array}{c}
S_{0}  \tag{1.22}\\
S_{1} \\
S_{2} \\
S_{3}
\end{array}\right]=\left[\begin{array}{c}
I \\
Q \\
U \\
V
\end{array}\right]=\left[\begin{array}{c}
<E_{x}^{*} E_{x}+E_{y}^{*} E_{y}> \\
<E_{x}^{*} E_{x}-E_{y}^{*} E_{y}> \\
<E_{y}^{*} E_{x}+E_{x}^{*} E_{y}> \\
<i\left(E_{y}^{*} E_{x}-E_{x}^{*} E_{y}\right)>
\end{array}\right] .
$$

The angle brackets $<\ldots>$ denote a time average and it is introduced to account for the fluctuations in the amplitude and phase of quasi-monochromatic light. The first Stokes parameter, $I$, is the total intensity of electromagnetic radiation, while the other three parameters describe the polarization state of radiation. For every Stokes vector the Stokes parameters satisfy the inequality

$$
\begin{equation*}
I^{2} \geq Q^{2}+U^{2}+V^{2} \tag{1.23}
\end{equation*}
$$

in which the equality is only satisfied for completely polarized light. The degree of polarization $p$ is defined from the Stokes parameters as

$$
\begin{equation*}
p=\frac{\sqrt{Q^{2}+U^{2}+V^{2}}}{I} . \tag{1.24}
\end{equation*}
$$

Thus, for completely polarized light $p=1$, for unpolarized light $p=0$, and for partially polarized $0<p<1$.

The Stokes parameters were introduced by Stokes in the XIX century. However the matrix method for manipulating Stokes vectors was developed by Hans Mueller much later, in the 1940's [2]. The Stokes vector representation is directly connected with experiment, because it uses real and measurable intensities instead of complex numbers. This has caused that Mueller matrices have become popular when performing experimental work in polarization optics; in fact, many modern optical instruments are able to partially or completely determine the Mueller matrix of an optical sample.

A Mueller matrix $\mathbf{M}$ is a $4 \times 4$ matrix with 16 real elements that connects the input and output Stokes vectors after the interaction of electromagnetic radiation with an optical medium.

$$
\begin{equation*}
\mathbf{S}^{o u t}=\mathbf{M S}^{i n} . \tag{1.25}
\end{equation*}
$$

The upper-left element of Mueller matrix $\mathbf{M}, m_{00}$, is a gain for unpolarized incident light and it must hold the inequality $m_{00}>0$. A normalized Mueller matrix is obtained by scaling the matrix such that the upper left element is equal to one.

$$
\mathbf{M}^{*}=\left[\begin{array}{cccc}
1 & m_{01} / m_{00} & m_{02} / m_{00} & m_{03} / m_{00}  \tag{1.26}\\
m_{10} / m_{00} & m_{11} / m_{00} & m_{12} / m_{00} & m_{13} / m_{00} \\
m_{20} / m_{00} & m_{21} / m_{00} & m_{22} / m_{00} & m_{23} / m_{00} \\
m_{30} / m_{00} & m_{31} / m_{00} & m_{32} / m_{00} & m_{33} / m_{00}
\end{array}\right]
$$

The elements of a normalized Mueller matrix have values between -1 and 1. All the experimental Mueller matrices that will be shown in this work are normalized.

A rotation of the coordinate system by an angle $\alpha$ (positive for a counterclockwise rotation looking against the direction of propagation) can be expressed as a matrix multiplication, in analogy to the coordinate rotation of Jones matrices presented in Eq. (1.20):

$$
\mathbf{M}^{\prime}=\mathbf{R}(\alpha) \mathbf{M R}(-\alpha), \quad \text { where } \quad \mathbf{R}(\alpha)=\left[\begin{array}{cccc}
1 & 0 & 0 & 0  \tag{1.27}\\
0 & \cos 2 \alpha & \sin 2 \alpha & 0 \\
0 & -\sin 2 \alpha & \cos 2 \alpha & 0 \\
0 & 0 & 0 & 1
\end{array}\right]
$$

On the other hand, if the polarization element described by the Mueller matrix is rotated by an angle $\alpha$, then the resulting Mueller matrix is

$$
\begin{equation*}
\mathbf{M}_{\alpha}=\mathbf{R}(-\alpha) \mathbf{M R}(\alpha) . \tag{1.28}
\end{equation*}
$$

The inequality in Eq.(1.23) is important because it allows the classification of the character of light-medium interaction. For completely polarized input light, if the output light satisfies Eq.(1.23) as an equality, then we say that the medium is non-depolarizing (i.e. the exiting beam is still completely polarized).

For every Jones matrix there exists a corresponding Mueller matrix. To emphasize this connection, we will call those Mueller matrices which can be derived from a Jones matrix as "Mueller-Jones" matrices. The Mueller-Jones matrix corresponding to any Jones matrix can be calculated using the following relation [3]:

$$
\begin{equation*}
\mathbf{M}=A\left(\mathbf{J} \otimes \mathbf{J}^{*}\right) A^{-1}, \tag{1.29}
\end{equation*}
$$

where $\otimes$ denotes de Kronecker product and $A$ is

$$
A=\frac{1}{\sqrt{2}}\left(\begin{array}{cccc}
1 & 0 & 0 & 1  \tag{1.30}\\
1 & 0 & 0 & -1 \\
0 & 1 & 1 & 0 \\
0 & i & -i & 0
\end{array}\right)
$$

A general Jones matrix has 8 independent parameters, the absolute phase information is lost in the Mueller-Jones matrix, leading to only 7 independent matrix elements for a Mueller-Jones matrix derived from a Jones matrix. Therefore, it is evident that in a Mueller-Jones matrix there exist 9 interrelations o constraints among its 16 elements $^{4}$. In the case of normalized Jones or Mueller-Jones matrices the number of independent parameters is 6 , because both the absolute phase and absolute intensity information are lost. It has been demonstrated [6] that a

[^3]necessary and sufficient condition for a Mueller matrix $\mathbf{M}$ to be a Mueller-Jones matrix is:
\[

$$
\begin{equation*}
\sum_{i=0}^{3} \sum_{j=0}^{3} m_{i j}^{2}=4 m_{00}^{2} \tag{1.31}
\end{equation*}
$$

\]

A $4 \times 4$ real matrix is not a physically realizable Mueller matrix if it can operate on an incident Stokes vector to produce a vector with a degree of polarization greater than one, which is a physically unrealizable polarization state. Experimentally it is possible to obtain a physically unrealizable Mueller matrix because of the noise inherent to experimental data and since the matrices of several ideal optical elements (polarizers, retarders) lie in the boundary between physically realizable Mueller matrices and nonphysical unrealizable matrices. The conditions for physical realizability have been thoroughly studied in the literature and several necessary conditions for the physical realizability have been published [7]. One of the most cited criteria for Mueller matrix physically realizability is that the eigenvalues of the associated coherency matrix need to be non-negative [7].

### 1.4.1. Depolarization

Depolarization is the reduction of the degree of polarization of light. Depolarization can be the result of decorrelation of the phases and the amplitudes of the electric vector and selective absorption of polarization state [8]. In fact, a measured Mueller matrix is a mixture of pure (nondepolarizing) states, depolarization and a certain amount of noise [9]. Despite depolarization phenomenon is involved in many experimental measurements, depolarization is perhaps the least understood characteristic concerning polarized light and the problem of light-medium interaction in presence of depolarization is studied with considerably less degree of accuracy than the nondepolarizing case. In polarimetry the causes of depolarization can be multiple, many of them are due to some type of incoherent scattering introduced by the sample or the incoherent superposition of the light interacting from different materials. Another source of depolarization is caused by finite spectral resolution of the instruments (determined in most cases by the monochromator). Instead of being perfectly monochromatic they integrate over a range of wavelengths, resulting in quasi-depolarization.

Some media depolarize all the input polarization states equally (isotropic depolarization). Other media depolarize some polarization states but may not depolarize some particular polarization states (anisotropic depolarization). The Mueller matrix of a isotropic depolarizer is:

$$
\left[\begin{array}{llll}
1 & 0 & 0 & 0  \tag{1.32}\\
0 & p & 0 & 0 \\
0 & 0 & p & 0 \\
0 & 0 & 0 & p
\end{array}\right]
$$

When $p=0$ one has the Mueller matrix of the ideal depolarizer.
To investigate the depolarization features of a medium it is useful to use the polarization fraction $\beta$ of a Mueller matrix that was introduced by Gil and Bernabeu $[10,11]$. It is the euclidian distance of the normalized Mueller matrix from the ideal depolarizer:

$$
\begin{equation*}
\beta=\frac{\sqrt{\left(\sum_{i j} m_{i j}^{2}\right)-m_{00}^{2}}}{\sqrt{3} m_{00}} \tag{1.33}
\end{equation*}
$$

here $\beta$ varies from zero for an ideal depolarizer to 1 for all nondepolarizing Mueller matrices. All intermediate values are associated with partial depolarization.

Eq. (1.33) is the figure of merit that we will use through this work to quantify depolarization of experimental Mueller matrices. Nevertheless it is not the only available criterion for depolarizing in the literature. Another used criterion was introduced by Cloude [12] and is based in the normalized eigenvalues of the coherency matrix associated with the Mueller matrix to define a "entropy" parameter that is understood as a degree of depolarization. Recently, Ossikovki has published two alternative criteria [13] that are claimed to offer a better identification of depolarization for certain classes of Mueller matrices.

### 1.5. Basic polarization changes induced by transmissive optical elements

In this section we will consider a simplified phenomelogical approach to describe the various forms in which an optical element can modify the polarization of a light beam that traverses it. Let us consider a monochromatic plane wave propagating along the positive direction of the $z$ axis of an $x, y, z$ orthogonal, right-handed, Cartesian coordinate laboratory system. Further, we will assume that the light beam traverses the optical element and does not change the direction of propagation; thus the electric field components are always in the $x y$ plane.

When an electromagnetic wave passes through a nondepolarizing medium it can experiment changes in intensity, phase and polarization, all of them defined relative to the case where there is no medium. Although in this work we are almost exclusively interested in the changes of polarization, we cannot forget that for every medium there can be also a change in intensity and absolute phase that are independent of the polarization of light used to illuminate the medium.

With these premises we will consider light propagation through a dichroic (diattenuating) and birefringent (retarding) anisotropic and nondepolarizing medium. In this medium eight different types of optical behaviour are measurable in the $x y z$ laboratory coordinate system: six of them are related to changes in polarization, one to changes in the absolute phase, and another one to changes in the
intensity. These can be described using Jones vectors since they contain complete information about the amplitudes and phases of the field components.

### 1.5.1. Absolute phase and intensity changes

We start presenting the simple case of free propagation of a plane wave of light through a medium that does not induce any change in the polarization. Light propagates through this medium at a different speed with respect to the case where there was no medium. Also absorption processes can change the amplitude of the components of the field. The complex index of refraction $\tilde{n}$ can be used to calculate the complex phase shift induced by the medium:

$$
\begin{equation*}
\mathbf{E}^{\prime}=\mathbf{E} e^{-i \frac{2 \pi}{\lambda} \tilde{n} l}=\mathbf{E} e^{-i \frac{2 \pi}{\lambda} n l} e^{-\frac{2 \pi}{\lambda} k l} \equiv \mathbf{E} e^{-i \chi} \tag{1.34}
\end{equation*}
$$

where the complex index of refraction is $\tilde{n} \equiv n-i k$ and $\chi \equiv \eta-i \kappa$. Therefore:

$$
\begin{align*}
\eta & =\frac{2 \pi}{\lambda} n l,  \tag{1.35a}\\
\kappa & =\frac{2 \pi}{\lambda} k l . \tag{1.35b}
\end{align*}
$$

The Jones vector transformation of Eq. (1.34) in matrix form is:

$$
\left[\begin{array}{c}
E_{x}^{\prime}  \tag{1.36}\\
E_{y}^{\prime}
\end{array}\right]=\left[\begin{array}{cc}
e^{-i \chi} & 0 \\
0 & e^{-i \chi}
\end{array}\right]\left[\begin{array}{l}
E_{x} \\
E_{y}
\end{array}\right]
$$

The term $e^{-i \frac{2 \pi}{\lambda} n l}$ of Eq.(1.34) expresses the simple fact that the wave has been retarded by $\frac{2 \pi}{\lambda} n l$, while the term $e^{-\frac{2 \pi}{\lambda} k l}$ expresses the amplitude attenuation of the Jones vector components.

The factor $e^{-\frac{2 \pi}{\lambda} k l}$ is related to the well-known Beer-Lambert law, which is used to study the absorption of light when it travels through a material. This law states that there is a logarithmic dependence between the transmission $T$ and the product of the absorption coefficient of a substance $\alpha$ and the the distance the light travels through the material, $l$ :

$$
\begin{equation*}
T=\frac{I}{I_{0}}=10^{-\alpha l}=10^{-A} \tag{1.37}
\end{equation*}
$$

This equation can be also written in the base of natural logarithms

$$
\begin{equation*}
T=\frac{I}{I_{0}}=e^{-\alpha_{e} l}=e^{-A_{e}} . \tag{1.38}
\end{equation*}
$$

Consequently the relation between $A$ and $A_{e}$ is $A=\ln 10 A_{e}$. Eq. (1.38) can be rewritten as $I=I_{0} e^{-A_{e}}$, and as the intensity of a wave is the square of its amplitude $A$ we have $A=A_{0} e^{-A_{e} / 2}$ and we get the relation:

$$
\begin{equation*}
A_{e}=\frac{4 \pi}{\lambda} k l \tag{1.39}
\end{equation*}
$$

that written in terms of $\alpha_{e}$ is $\alpha_{e}=4 \pi k / \lambda$. By default all the absorptions given in this work will displayed in the base of natural logarithms.


Figure 1.2.: Scheme of Cartesian coordinate system used for the description of the optical effects. The optical element is supposed to be a plane-parallel section of width $l$.

### 1.5.2. Polarization changes

## Linear birefringence

Suppose that the medium through which the wave is propagating is not isotropic, but rather uniaxially linearly birefringent, which means that there is one axis of symmetry, called the optic axis, which imposes constraints upon the propagation of light beams within the material. Two particular modes of propagation can be described, either as an ordinary beam polarized in a plane perpendicular to the optic axis, or as an extraordinary beam polarized in a plane containing the optic axis ${ }^{5}$. Each one of the beams has an associated refractive index and the difference between these refractive index is eventually known as linear birefringence.

When the value of the linear birefringence of a certain material is given in the literature, it is implicitly assumed that this value corresponds to the natural basis of the material (a coordinate system that corresponds with its optic axis),

[^4]so that the given value of the linear birefringence is the difference of refractive indices for extraordinary, $n_{e}$, and ordinary polarizations, $n_{o}$. Nevertheless, during experiments, the orientation of the optic axis for a certain sample is in general unknown and the measurements take place in a laboratory coordinate frame not coincident with the optical axis. For optical calculus it is useful to define the laboratory axes shown in Fig. 1.2: $x, y, 45^{\circ}$ to $x$ and $135^{\circ}$ to $x$ because these axes permit to define two parameters, for the projection of linear birefringence along $x$ and another for the projection along $45^{\circ}$ that come in linearly in the Mueller matrix calculus.

In an arbitrarily oriented linearly birefringent medium the if wave that propagates in the $z$ direction is linearly polarized parallel to the $x$ axis, the wave will experiment a specific refractive index $n_{x}$ and will travel at a speed $c / n_{x}$, where $c$ is the speed of light in vacuum. On the other hand, if the wave is linearly polarized orthogonal to the $x$ axis, the wave will experience a different refractive index $n_{y}$ and will travel at a speed $c / n_{y}$. Again, if the wave is polarized at $45^{\circ}$ to the $x$ direction it will experience a refractive index $n_{45}$, and if it is orthogonal to this last direction the refractive index will be $n_{135}$. It is worth to stress that $n_{x}, n_{y}, n_{45}$ and $n_{135}$ are not independent between them; they all can be related to $n_{e}$ and $n_{o}$ if the orientation of the optical axis in our laboratory coordinate frame is known, as it will be further discussed in Section 3.2.3. Using these newly defined refractive indices, the effect of wave propagation in birefringent media can be described using the following two parameters :

- the horizontal linear birefringence projection,

$$
\begin{equation*}
\mathrm{LB}=\frac{2 \pi}{\lambda}\left(n_{x}-n_{y}\right) l \tag{1.40}
\end{equation*}
$$

- and the $45^{\circ}$ linear birefringence projection,

$$
\begin{equation*}
\mathrm{LB}^{\prime}=\frac{2 \pi}{\lambda}\left(n_{45}-n_{135}\right) l \tag{1.41}
\end{equation*}
$$

where $l$ is the travelled distance.
The effect of the horizontal linear birefringence on a Jones vector is given by the Jones matrix.

$$
\mathbf{J}_{\mathrm{LB}}=\left[\begin{array}{cc}
e^{-i \frac{2 \pi}{\lambda} n_{x} l} & 0  \tag{1.42}\\
0 & e^{-i \frac{2 \pi}{\lambda} n_{y} l}
\end{array}\right]
$$

and, alternatively, this matrix can be written in a more convenient form:

$$
\begin{align*}
\mathbf{J}_{\mathrm{LB}} & =e^{-i \frac{2 \pi}{\lambda} n_{x} l}\left[\begin{array}{cc}
1 & 0 \\
0 & e^{i \frac{2 \pi}{\lambda}\left(n_{x}-n_{y}\right) l}
\end{array}\right]=e^{-i \frac{\pi}{\lambda}\left(n_{x}+n_{y}\right) l}\left[\begin{array}{cc}
e^{-i \frac{\pi}{\lambda}\left(n_{x}-n_{y}\right) l} & 0 \\
0 & e^{i \frac{\pi}{\lambda}\left(n_{x}-n_{y}\right) l}
\end{array}\right] \\
& =e^{-i \eta / 2}\left[\begin{array}{cc}
e^{-i \mathrm{LB} / 2} & 0 \\
0 & e^{i \mathrm{LB} / 2}
\end{array}\right], \tag{1.43}
\end{align*}
$$

where the outside scalar factor represents an isotropic phase delay.
Leaving aside the isotropic factor, the effect of the $45^{\circ}$ linear birefringence on a Jones vector in the coordinate axes $x-y$ can be easily calculated by a coordinate rotation of $-45^{\circ}$ from the $45^{\circ}-135^{\circ}$ axes to the $x-y$ axes:

$$
\mathbf{J}_{\mathrm{LB}^{\prime}}=\mathbf{R}\left(-45^{\circ}\right)\left[\begin{array}{cc}
e^{-i \mathrm{LB}^{\prime} / 2} & 0  \tag{1.44}\\
0 & e^{i \mathrm{LB}^{\prime} / 2}
\end{array}\right] \mathbf{R}\left(45^{\circ}\right)=\left[\begin{array}{cc}
\cos \mathrm{LB}^{\prime} / 2 & -i \sin \mathrm{LB}^{\prime} / 2 \\
-i \sin \mathrm{LB}^{\prime} / 2 & \cos \mathrm{LB}^{\prime} / 2
\end{array}\right]
$$

If the maximum value of the birefringence, $\mathrm{LB}_{\theta}$, is obtained at angle $\theta$ then LB and $\mathrm{LB}^{\prime}$ are related to $\mathrm{LB}_{\theta}$ and $\theta$ by:

$$
\begin{align*}
& \mathrm{LB}=\mathrm{LB}_{\theta} \cos 2 \theta  \tag{1.45a}\\
& \mathrm{LB}^{\prime}=\mathrm{LB}_{\theta} \sin 2 \theta \tag{1.45b}
\end{align*}
$$

It is important to stress that neither of the quantities defined LB or $\mathrm{LB}^{\prime}$ and that we, respectively, name as horizontal or $45^{\circ}$ linear birefringence are equivalent to the common definition of linear birefringence that is made in crystal optics bibliography ${ }^{6}$. Our definitions of LB or $\mathrm{LB}^{\prime}$ are always given with reference to the laboratory coordinate system instead of from the own optic-axis based coordinate system of the anisotropic medium. In other words, values of LB and $\mathrm{LB}^{\prime}$ change if the orientation of the anisotropic medium changes.

## Linear dichroism

If the wave propagates through a linearly dichroic medium, a linearly-polarized wave travelling in a direction perpendicular to the optic axis will be attenuated by different amounts depending upon the direction of the vibration of the transverse electric field with respect to the optic axis. Following an analogous argumentation to that presented in the previous section let $k_{x}$ and $k_{y}$ respectively denote the extinction coefficients of the medium for light linearly polarized parallel and orthogonal to the $x$ axis, while $k_{45}$ and $k_{145}$ are the extinction coefficients for light linearly polarized parallel and orthogonal to the $45^{\circ}$ axis. With these coefficients we can define the following parameters:

- the horizontal linear dichroism projection,

$$
\begin{equation*}
\mathrm{LD}=\frac{2 \pi}{\lambda}\left(k_{x}-k_{y}\right) l \tag{1.46}
\end{equation*}
$$

- the $45^{\circ}$ linear dichroism projection,

$$
\begin{equation*}
\mathrm{LD}^{\prime}=\frac{2 \pi}{\lambda}\left(k_{45}-k_{135}\right) l . \tag{1.47}
\end{equation*}
$$

[^5]The Jones matrix related to the horizontal linear dichroism is

$$
\mathbf{J}_{\mathrm{LD}}=e^{-i \eta / 2}\left[\begin{array}{cc}
e^{-\frac{2 \pi}{\lambda} k_{x} l} & 0  \tag{1.48}\\
0 & e^{-\frac{2 \pi}{\lambda} k_{y} l}
\end{array}\right]=e^{-i \eta / 2} e^{-i \kappa / 2}\left[\begin{array}{cc}
e^{-\mathrm{LD} / 2} & 0 \\
0 & e^{\mathrm{LD} / 2}
\end{array}\right] .
$$

where the $\kappa$ term accounts for the isotropic absorption.
The same coordinate rotation procedure that we used to calculate the Jones matrix corresponding to the $45^{\circ}$ linear birefringence can be introduced here to determine the Jones matrix associated to a $45^{\circ}$ linear dichroism:

$$
\mathbf{J}_{\mathrm{LD}^{\prime}}=\mathbf{R}\left(-45^{\circ}\right)\left[\begin{array}{cc}
e^{-\mathrm{LD} / 2} & 0  \tag{1.49}\\
0 & e^{\mathrm{LD} / 2}
\end{array}\right] \mathbf{R}\left(45^{\circ}\right)=\left[\begin{array}{cc}
\cosh \mathrm{LD}^{\prime} / 2 & -\sinh \mathrm{LD}^{\prime} / 2 \\
-\sinh \mathrm{LD}^{\prime} / 2 & \cosh \mathrm{LD}^{\prime} / 2
\end{array}\right]
$$

If $\mathrm{LD}_{\phi}$ is the maximum value of the linear dichroism and it is obtained at an angle $\phi$ from the the x axis then LD and $\mathrm{LD}^{\prime}$ are related to $\mathrm{LD}_{\phi}$ and $\phi$ by:

$$
\begin{align*}
& \mathrm{LD}=\mathrm{LD}_{\phi} \cos 2 \phi  \tag{1.50a}\\
& \mathrm{LD}^{\prime}=\mathrm{LD}_{\phi} \sin 2 \phi \tag{1.50b}
\end{align*}
$$

The angles $\theta$ and $\phi$ appearing in Eqs. (1.45) and (1.50) are not necessarily equal. In practice, however, for a considerable number of situations (i.e. uniaxial crystals, oriented molecules, etc) the principal axes for linear retardation and linear dichroism coincide.

## Circular birefringence

Linearly polarized light that propagates through a medium exhibiting circular birefringence remains linearly polarized as the light progresses, but the direction of the vibration of the electric vector rotates uniformly and continuously with distance. This is why a material with circular birefringence is said to possess optical rotatory power or produce optical rotation.

Circular birefringence occurs due to a difference between the indices of refraction $n_{-}$and $n_{+}$for left and right circularly polarized light waves, respectively, with a typical magnitude of $\left|n_{-} n_{+}\right| \sim 10^{-5}$. Circular birefringence arises from an intrinsic helical structures present in atom or in molecules. The parameter CB that quantifies the circular birefringence is defined as:

$$
\begin{equation*}
\mathrm{CB}=\frac{2 \pi}{\lambda}\left(n_{-}-n_{+}\right) l, \tag{1.51}
\end{equation*}
$$

A circularly birefringent medium can be described by the following Jones matrix [3]:

$$
\mathbf{J}_{\mathrm{CB}}=\left[\begin{array}{cc}
\cos \mathrm{CB} / 2 & \sin \mathrm{CB} / 2  \tag{1.52}\\
-\sin \mathrm{CB} / 2 & \cos \mathrm{CB} / 2
\end{array}\right] .
$$



Figure 1.3.: Rotation of an incident polarization state induced by a circularly birefringent medium.

Comparing this matrix to matrix in Eq. (1.20) is easy to realize that CB is twice the angle of rotation. Therefore the angle of rotation $\rho$ induced by a medium with circular birefringence is $\rho=\mathrm{CB} / 2$. According to our sign criterium, Fig. 1.3 shows a positive optical rotation on linearly polarized radiation induced by a circularly birefringent medium.

## Circular dichroism

In a circularly dichroic medium there is a different absorption for the left an right circularly polarized light. The differences in the extinction coefficient for left, $k_{-}$, and right, $k_{+}$circularly polarized light waves is quantified as

$$
\begin{equation*}
\mathrm{CD}=\frac{2 \pi}{\lambda}\left(k_{-}-k_{+}\right) l . \tag{1.53}
\end{equation*}
$$

The Jones matrix corresponding to CD is

$$
\mathbf{J}_{\mathrm{CD}}=\left[\begin{array}{cc}
\cosh \mathrm{CD} / 2 & i \sinh \mathrm{CD} / 2  \tag{1.54}\\
-i \sinh \mathrm{CD} / 2 & \cosh \mathrm{CD} / 2
\end{array}\right] .
$$

Any medium that cause the differential absorption of circularly polarized light -circular dichroism- will transform a linearly polarized input into elliptically polarized. Usually, a medium containing CB or CD but not $\mathrm{LB}, \mathrm{LD}, \mathrm{LB}^{\prime}, \mathrm{LD}^{\prime}$ is said to be optically active and isotropic. An illustrative example is a sugar solution, although it can change the polarization of incident light (because of its $\mathrm{CB})$ its optical properties are the same in all directions.

Table 1.1 summarizes the eight optical effects that we have studied in this section. The two first ones $(\eta, \kappa)$ are not included in a normalized Mueller-Jones
matrix, but, as it will be shown later, the remaining 6 effects can generate the 6 independent parameters that form a Mueller-Jones matrix. The dispersive effects $\eta, \mathrm{LB}, \mathrm{LB}^{\prime}$ and CB are related to the real part of the complex index of refraction, while the absorptive parameters $\kappa, \mathrm{LD}, \mathrm{LD}^{\prime}$ and CD involve the imaginary part. For later convenience it is useful to go one step further and define a general complex retardation for each birefringence-dichroism pairs:

$$
\begin{align*}
\chi & =\eta-i \kappa  \tag{1.55a}\\
\mathrm{~L} & =\mathrm{LB}-i \mathrm{LD}^{\prime}  \tag{1.55b}\\
\mathrm{L}^{\prime} & =\mathrm{LB}^{\prime}-i \mathrm{LD}^{\prime}  \tag{1.55c}\\
\mathrm{C} & =\mathrm{CB}-i \mathrm{CD} \tag{1.55d}
\end{align*}
$$

Table 1.1.: Symbols used and definitions

| effect | symbol | definition $^{a}$ | Jones matrix |
| :--- | :---: | :---: | :---: |
| isotropic phase re- <br> tardation | $\eta$ | $\frac{2 \pi}{\lambda}\left(n_{x}+n_{y}\right) l$ | $\left[\begin{array}{cc}e^{-i \eta} & 0 \\ 0 & e^{-i \eta}\end{array}\right]$ |
| isotropic ampli- <br> tude absorption | $\kappa$ | $\frac{2 \pi}{\lambda}\left(k_{x}+k_{y}\right) l$ | $\left[\begin{array}{cc}e^{-\kappa} & 0 \\ 0 & e^{-\kappa}\end{array}\right]$ |
| horizontal linear <br> dichroism projec- | LD | $\frac{2 \pi}{\lambda}\left(k_{x}-k_{y}\right) l$ | $\left[\begin{array}{cc}e^{-\mathrm{LD} / 2} & 0 \\ 0 & e^{\mathrm{LD} / 2}\end{array}\right]$ |
| tion |  |  |  |

[^6]
## Chapter 2.

## Molecular and supramolecular optical activity

### 2.1. Historical introduction

Historically the study of chirality in molecules is intimately relied to the chirality in crystals. In fact quartz crystal played an important role in the firsts studies about optical activity that date from the beginning of of the nineteenth century. The first observation is reported to having been done by the French catalan astronomer Arago in 1811 [14] when he observed colours in the sunlight that had passed along the optic axis of a quartz crystal placed between crossed polarizers. Soon after, Jean Baptiste Biot (1812) [15] reported more experiments with quartz and discovered a second form of the crystal which rotated the plane of polarization in the opposite direction.

Biot also discovered that the rotation could also be observed in in organic liquids such as turpentine. At that time, it was already appreciated that the optical activity of fluids must reside in the individual molecules, and may be observed even when the molecules are randomly oriented; whereas that of quartz was a property of the crystal structure and not of the individual molecules, since molten quartz was not optically active.

The first phenomenological theory about optical activity was completed by Fresnel in 1825 [16]. At that time he had discovered circular polarization and he realized that incident linearly polarized light can be regarded as a superposition of the two possible forms of circularly polarized light [right (R) and left (L)]. He correctly attributed optical rotation to the propagation at different speeds in an optically active medium of the left- and right-circularly polarized components of the linearly polarized beam in the medium. Fresnel was able to show the existence of the two circular components and separate them by an ingenious construction of a composite prism consisting of R- and L- quartz: as the two components travel with different velocities in an optical active medium they also refract by different amounts at an oblique interface.

By 1846, Haidinger [17] reported differences in the measured absorptions of left and right circularly polarized light, which is the origin of CD, but until the work of Aimé Cotton in 1895 [18] the phenomenon was not put in relief. The first theoretical molecular model of optical activity was proposed by Drude in 1900. In his model absorbing media contain charged particles or ions, which

## Chapter 2. Molecular and supramolecular optical activity

posses natural periods of vibration. The vibrations of these particles around an equilibrium position are more or less violent if their periods of vibration agree more or less closely with the periods of the incident light. In the Drude model the rotation of the plane of polarization $\rho$ is given by the equation ${ }^{1}$

$$
\begin{equation*}
\rho=\sum_{r} \frac{Q_{r}}{\lambda^{2}-\lambda_{r}^{2}}, \tag{2.1}
\end{equation*}
$$

where $Q_{r}$ is a constant depending on the number of vibrators in unit volume and other constants of the medium and $\lambda_{r}$ is the wavelength corresponding with the natural period of vibration of the molecule. The Drude equation has a great application in the molecular optical activity, however its application in the crystalline optical activity is relatively small

It is worth also mentioning that a thorough account of the experimental and theoretical side of the subject was given in a book by Lowry in 1935 [19] that for years was the main reference on the subject and still is a valuable source of information due to the great amount of experimental measurements of optical activity he reviews. During the XXth century the coupled oscillator model for optical activity introduced by Chandrasekhar in 1956 [20] had special relevance because it was the basis for several of the theoretical approaches to study molecular optical activity. It is based on the conception that the valence electrons of atoms or molecules forming the optically active crystals can be represented by the helically arranged and mutually coupled linear harmonic oscillators. The Chandrasekhar formula involves only two constants k and $\lambda_{0}$ :

$$
\begin{equation*}
\rho=\frac{k \lambda^{2}}{\left(\lambda^{2}-\lambda_{0}^{2}\right)^{2}} . \tag{2.2}
\end{equation*}
$$

One of the earliest applications of optical activity was in sugar industry, where the rotatory power measured on sugar solutions is used as a measure of the quality of the sugar (saccharimetry). Besides this industrial application, in recent years molecular optical activity has arisen as a very important subject of fundamental research in several branches of chemistry and that also has many applications in pharmacology and biochemistry for the study of the molecules of living organisms.

In this Chapter we will descriptively review several aspects about the presence of electronic optical activity in molecules and molecular aggregates. Further we will also introduce the concept of homochirality and then we will briefly discuss the implications that it has in the study of the emergence of life in Earth. An

[^7]additional section will present the basic characteristics of the commercial instruments that are used in laboratories all over the world to measure the optical activity and, there, we will discuss the limitations they have. In the last section we will review some of the most notable published contributions on the measurement of optical activity in anisotropic media and the current state-of-art of the subject. All the concepts discussed in this chapter will be studied from a purely descriptive approach and without introducing many details, as the only objective of this chapter is to set a proper background for a further understanding of the following chapters.

### 2.2. Circular dichroism and circular birefringence in molecules

The optical definitions for circular dichroism (CD) and circular birefringence (CB) has been given in the previous Chapter. A molecule is chiral when the mirror image of the molecular structure cannot be brought to coincide with itself. Here "brought" refers to translations and rotations. A chiral molecule is said to be optical active (or possess optical activity) if it exhibits CD and CB.

The structure of chiral molecules belong to symmetry point groups ${ }^{2}$ that lack the symmetry elements of a mirror plane, a center of inversion, or a rotationreflection axis $n>2 .{ }^{3}$ Other symmetry elements like axis of rotation are allowed. Image and mirror image of a chiral molecule are called enantiomers. As a consequence of the lack of mirror symmetry the enantiomers of a chiral molecule are "non-superposable". In chemistry, the word racemic is used for compounds or mixtures of chiral molecules that have equal amounts of left- and right-handed enantiomers of a chiral molecule. Therefore, a racemic compound despite containing chiral molecules is optically inactive, i. e. shows no CD or CB , because the contributions of both enantiomers, that are present in equal amounts, cancel each other.

The electronic transitions between two different electronic configurations of a molecule induced by electromagnetic radiation of any wavelength is called a chromophoric transition. Usually we understand the chromophore as that region of a molecule where the energy difference between two different localized molecular orbital falls within the range of the uv-visible spectrum. Visible light that hits the chromophore can thus be absorbed by exciting an electron from its ground state into an excited state, and, as a consequence, these chromophoric transitions at the visible range are responsible for the colour of the molecule. To a first approxima-

[^8]tion chromophores themselves are usually not chiral, although there are notable exceptions, like helicenes. In case of achiral chromophores the observed optical activity then arises from interaction between different chromophores within the chiral geometrical framework of the molecule.

A substance can, however, also be optically active due to an asymmetrical arrangement of molecules. In this cases, the molecules themselves need not necessarily to be chiral. Arranging non-chiral molecules on a helix, results, for instance, in an optically active molecule because a helix and its mirror image cannot be superposed since reflection reverses the screw sense. Chirality excludes improper symmetry elements (operations involving reflection and inversion), namely centers of inversion, reflection planes and rotation-reflection axes. A chiral molecule is not necessarily asymmetric as it can have, for example, rotational symmetry. The spatial symmetry restrictions that impose chirality are not only applicable to molecules, but also to crystals and they are mathematically studied with the point group classification and constitute a very important subject in stereochemistry (which studies molecular geometry) and crystallography.

The origin of the optical activity in crystals can be of two types. An optically active crystal can be formed of optically active molecules -molecular origin- or it can result from the crystal structure -crystalline origin-. It is known that in the case of the crystalline origin the crystal is characterized by an helix structure, i.e. the molecules or atoms lie on the helices the axes of which are parallel to the optical axis of the crystal. The best known optically active crystals are those of the space groups of symmetry $D_{3}^{4}$ and $D_{3}^{6}$ the elementary cell of which consist of three atoms or molecules. Typical examples of these crystals are $\alpha$-quartz, tellurium, selenium, benzil or cinnabar. The majority of them are optical active due to the crystalline structure, which means that, for example, if they are fused they lose the optical activity. However, a certain number of crystals, camphor is a well-known example, are constituted by optically active molecules, and thus its optical activity has both contributions: molecular and crystalline.

Although CD and CB are two effects that are interrelated they are usually treated separately. CD is nonzero only in the spectral regions in which the molecule under study has optical transitions. However, CB is a dispersive measurement, and for an accurate study requires a scan over a wide spectrum, while CD is better suited for studies of complex chromophores having closely spaced electronic transitions because of its inherently better resolution. With CD is easier to measure isolated transitions whereas CB measurements usually combine information about many electronic transitions.

CD spectroscopy is used routinely to study biological molecules and the applications of this spectroscopy in chemistry are multiple. We will not go into many details about these applications as they fall out of the scope of this thesis as we are more interested on fundamental research about homochirality (see subsection 2.3). We can mention, for example, that the far-UV CD spectrum of proteins
reveal important characteristics of their secondary structure. For example CD spectra is used to estimate the fraction of a molecule that is in the alpha-helix conformation, in the beta-sheet conformation, in the beta-turn conformation, etc. The tertiary structure of proteins can be studied in the near-UV region ( 250 nm to 350 nm ), for example, the chromophores are sensible to a folded structure in this region.

### 2.2.1. Cotton effect

Optically active substances can present CB and CD simultaneously in the range of frequencies where they have an absorption band. Such phenomenon is usually referred as Cotton-effect. When an absorption band of an isolated electronic transition is scanned in wavelength the typical Cotton-effect profiles displayed in Fig. 2.1 are obtained. A Cotton-effect is called to be positive, when the ORD maximum is found at longer wavelengths than the absorption maximum (case b of Fig. 2.1). Otherwise it is called negative.


Figure 2.1.: Scheme of the characteristic profiles that appear in the Cotton effect. The Cotton effect is called positive if the CB first decreases as the wavelength increases, and negative if CB first increases.

The Cotton effect results from nothing else than the application of the KramersKronig relations to an absorption band. They are relations between the real and
imaginary parts of a response function, and they are widely used in solid-state physics to evaluate the corresponding quantity if only one component is measured. In the subject of optical activity the real part, CB , and the imaginary part, CD , [see Eq. (1.55d)] are related by a Kramers-Kronig transform and, therefore, from a purely theoretical standpoint, provide the same information. However, from a practical standpoint it is highly desirable to have access to both parameters, since the application of a Kramers-Kronig transform theoretically requires the knowledge of the CB or CD response in the complete electromagnetic spectrum and, as a consequence, the transformation between experimental CD and CB has been rarely undertaken. Complete studies about the application of KramersKronig transformation in experimental measurements of CD and CB can be found in Refs. [21,22].

### 2.2.2. Excitons

Changes in the distribution of the electron density in the chromophores are associated with changes in the UV-visible absorption spectrum. The movement of the electron density in going from the ground electronic state to an excited state creates a momentary dipole, that is called a transition dipole. Each transition has both a direction and an intensity that vary according to the nature of the chromophore and the particular excitation. When two or more chromophores are brought into proximity, their electric transition dipoles may interact through dipole-dipole (or exciton) coupling. Therefore if one molecule in a group of close packed molecules is excited, the excitation can be transferred from molecule to molecule in the manner of a moving particle. Such moving excitation is called exciton. Excitons play an important role in determining the optical response of condensed-phase systems.

A Frenkel exciton [23] is basically an intramolecular excited state, in which the excited electron and the vacancy or "hole" migrate together from one unit cell to another unit cell. The Frenkel exciton differs from an electron in the fact that is neutral and carries no current, but it carries energy. The Frenkel exciton describes a delocalized electronic excitation, while the electron remains localized in the same unit cell. Therefore a Frenkel exciton can be considered as a neutral particle that can diffuse from site to site, perhaps moving hundreds of molecules away from its origin (energy is transported without migration of the electric charge). The Wannier excitons, typical of inorganic semiconductors, are different from Frenkel excitons in the fact that the electron and hole are separated by many lattice spacings, and, as a result the electron and hole are not tightly bounded as in the case of Frenkel exitons. The coupling of the optical transitions of the molecules results in the formation of a band of Frenkel exciton states. The concept of exciton can be also applied to the atoms or molecules of a crystalline solid [24], but in this thesis we are only interested in this concept when applied
to a molecular aggregate.
The absorption spectrum of an organic compound can differ in the solid and solution states. The formation of ordered structures usually referred as aggregates is fairly common in concentrated dye solutions and is often difficult to avoid when the molecules are in the solid state. Aggregates are of fundamental interest as model materials to study the nature of excitons in molecular systems of reduced dimensionality often with a chain-like configuration. The optical properties of these molecular aggregates are, to a large extent, governed by exciton delocalizations and other transport and relaxation phenomena. The aggregate formation can be usually well identified by the observation of the UV-visible absorption spectra because aggregation leads to a shifted, broadened, and sometimes split bands. Likewise, the interaction of two or more achiral chromophores that are held in close proximity can also give rise to optical activity. Assuming achiral chromophores there are three recognized methods (see Ref. [25]) of generation of optical activity: (i) dipole coupling mechanism; (ii) $\mu$-m mechanism and (iii) one-electron mechanism.

The dipole coupling requires two chromophoric groups coupled via dipoledipole interaction [26]. Fig. 2.2 displays the different coupling possibilities between the induced dipoles, that depend on the orientation of the aggregates. If the mutual orientation of the dipoles is chiral it results in a couplet of CD bands that correspond (see the oblique case of Fig. 2.2) to in-phase and out-of-phase combinations of uncoupled components. The pair of CD bands are equal in intensity but have opposite signs, and usually result in bisignate spectrum. The $\mu$-m mechanism is similar in concept but in this case the interaction is not between two dipoles but between one dipole and one quadrupole (magnetic moment) and it also requires at least two chirally oriented chromophores. The one-electron mechanism is conceptually different and, in principle, it does not requires a chiral orientation of the chromophores. It describes a mixing of electronic configurations within a single originally non-chiral chromophore under the influence of a chiral perturbation potential.

For an observed chiral transition it is highly likely that it would contain contributions of all three mechanisms. These are only highly simplified models that are used to understand the complicated nature of the excitonic states in molecules.

### 2.2.3. J-aggregates

We devote a special section of this introductory chapter to the description of a particular type of aggregates: the J-aggregates. We do so because several of the experimental studies that will be presented in further chapters are developed on systems with J-aggregation. Furthermore J-aggregates are of fundamental importance for the general understanding of molecular and solid-state physics because they bridge the gap between the physics of single molecules and perfectly


Figure 2.2.: Orientation dependence in exciton coupling between two chromophores (rectangles) and their long-axis induced dipoles (represented by doubleheaded arrows). When the two chromophores lie sufficiently close the excitations are delocalized between the two chromophores and the excited state ( E ) is splitted. The solid arrows connecting the ground (G) and excited (E) states represent allowed transitions, while dashed arrows represent forbidden transitions. The relative orientation of the chromophores are found in the wavelength-shifted absorption spectra -red-shifted for in-line and blue-shifted for parallel orientations-. The oblique orientation, when both transitions are allowed, leads to a broadened absorption band. CD transitions are only allowed when the dipoles have a chiral oblique orientation and results in a bisignate CD curve.
ordered crystals

J-aggregates were independently discovered in 1930's by Günter Scheibe and Edwin E. Jelley $[27,28]$ (from him comes the $J$ letter in their denomination). They discovered that by increasing the concentration of a solution of the dye pseudo-isocyanine the absorption spectrum underwent dramatic changes. In low concentration there was a relatively broad absorption band that upon increasing the concentration was replaced by a red-shifted much narrower absorption band ${ }^{4}$. Both authors attributed the changes to the formation of a molecular aggregate of dye molecules. Since then the aggregation behavior of cyanine dyes has been studied extensively and nowadays they are probably the best-known self-aggregating dyes.


Figure 2.3.: J-aggregate chain made of porphyrin molecules. J-aggregation takes place due to the electrostatic intermolecular interaction between the anionic sulphonate ( R radicals in the figure) and the positive-charged ring.

J-aggregates have been investigated as model material for one-dimensional Frenkel excitons. According to exciton theory the dye molecule is regarded as a point dipole and the excitonic state of the dye aggregate splits into two levels through the interaction of transition dipoles. Exciton theories such as the one shown in Fig. 2.2 are able to predict position of the absorption band of the Jaggregates and the unique feature of the narrowness of this so-called J band is explained by the absence of vibrational broadening, because the J band is shifted to energies where monomer electronic vibrational coupling is absent. Coherent exciton scattering theories play also an important role to fully understand the formation of J-aggregates.

Due to their narrow absorption spectra J-aggregates only absorb photons with a very narrow energy distribution and, for example, this gave them applications in the (analogical) photographic industry as film sensitizer. Applications of these molecules in various devices or for constructing molecular machines are also inves-

[^9]tigated because J-aggregates show interesting linear and nonlinear optical properties. We dedicate special attention to J-aggregates in this work because when they are in solution they can undergo asymmetric transformations that convert non-chiral molecules into chiral fiber-like associations.

### 2.3. Homochirality

Homochirality is a term used to refer to a group of molecules that possess the same sense of chirality. Homochirality is observed in living organisms because the nucleotids are only composed of right handed (D-) carbohydrates (sugars) and the proteins are formed by left handed (L-) amino acids. There are no traces on Earth of life based on D-aminoacids and L-carbohydrates.

Life is considered to be basically a molecular phenomenon and, at the moment, there is no clue why the existing life broke the mirror symmetry and chose these forms of aminoacids and carbohydrates instead of their mirror images. When aminoacids are being synthesized in the laboratory, e.g. via the famous UreyMiller experiment, a racemic mixture of right and left handed molecules is produced (equal amount of L and D molecules). However, it is known that life cannot originate in a racemic mixture because processes that are crucial for life such as replication or regulation of the genetic expression imply chiral supramolecular structures. Therefore the intriguing point of biological homochirality is not only the exclusive presence of L-aminoacids and D-carbohydrates in the biological forms, but also why life is tied to molecular structures and molecular recognition processes that are determined by chirality [29].

Theories for the origin of chirality in biological world can be grouped in two major categories: biotic and abiotic [30]. The biotic theory states that selection of one of the enantiomers took place at a late stage in the biological evolution of living matter. The second scenario proposes that chiral material were formed prior to the appearance of earlier biological polymers. In any of these two possible scenarios the first question to be asked is whether there was a cause that provoked the mirror-symmetry breaking. There have been several "explanations" about how this asymmetry was generated and we can group them in deterministic and stochastic or "chance" theories [30,31]. Deterministic theories defend that the observed mirror symmetry breaking was caused by an earlier external chiral influence (i.e. an asymmetric physical agent) even if this was on a minuscule state. Stochastic or chance -in the sense of randomness- theories [32] state that the mirror symmetry breaking takes place due to a very small random enantiomeric excess.

Among the most cited deterministic mechanisms there is circularly polarized light or the combination of plane polarized light and oriented magnetic field that could induce asymmetric synthesis or asymmetric decomposition of enantiomers.

Astrophysicists have observed that light emitted from interstellar stars is circularly polarized and thus it might have some influence in the generation of nonracemic extra-terrestrial materials that landed on Earth. Nevertheless there is a long-standing controversy as to whether these effects are large enough to have determined the selection of the handedness of life [33]. The stochastic scenarios can happen in far-from-equilibrium systems that undergo phase transitions, and there are several experimental examples of symmetry breaking phenomena related to crystallization or phase-transition phenomena.

Both stochastic and deterministic theories of biomolecular homochirality evolved through the same sequential stages: mirror-symmetry breaking followed by chiral amplification and chiral transmission [30]. The mirror-symmetry breaking imply the creation of a primordial minute enantiomeric imbalance. It is the key process in deterministic mechanism, whereas it is not for stochastic mechanisms, since for them the symmetry breaking occurs spontaneously the crucial process is an efficient mechanism of chiral amplification [30]. The chiral amplification stage has attracted great interest and the attention has been put in autocalytic reactions that amplify the chirality (i. e. the chirality of the reaction products is greater that of the starting materials). In this context Frank proposed a well-known model [34] for the autocatalytic self-replicating process in which a chemical substance catalyzes its own production and acts as an anticatalyst for the production of the opposite enantiomer [31]. This permits the amplification of even an extraordinary small initial enantiomeric excess to high levels. Only a small number of autocatalytic processes are known so far, the most famous one is the reaction discovered by Soai et al. [35].

In this thesis we defend that homochirality emerged in the abiotic stages of evolution. There exists the likelihood that most of the early organic material on Earth was brought in by comets and asteroids, and supporting our hypothesis, there are reports that indicate that aminoacids extracted from some meteorites such as the Murchison meteorite already contain some chiral bias towards Lenantiomers $[36,37]$. Most of the investigations presented in this thesis could be described, as a last resort, as a research on homochirality. Several of the results of this work were motivated by investigation on mechanisms that act in processes that undergo mirror symmetry breaking. In this respect we can anticipate that in the following chapters we will study macroscopic processes such as simple stirring (mechanical force) that can lead to chiral selection in certain aggregates, which are mechanisms that until very recently where not recognized to affect the molecular chirality. However we should warn the reader that the results that will be presented in this work concern only a small and quite specialized part with respect to the complete and multidisciplinary investigation carried on homochirality. Furthermore this thesis is, more generally, intended to contribute to the understanding of the interaction of polarized with chiral anisotropic media, so only occasionally we will refer to the implications that our research has in the
study of the processes of emergence of homochirality.

### 2.4. Standard methods to measure optical activity

CD is now a routine tool in many laboratories and CD instruments are usually referred to as circular dichrometers, dichrographs or CD spectropolarimeters. The essential feature of a CD spectropolarimeter is the implementation of a polarization phase modulation technique that enhances the sensibility of the instrument in the detection of the absorbance signal of a CD spectrum, that typically is orders of magnitude weaker than a normal absorbance spectrum.

In most commercial CD spectrometers, a photoelastic modulator ${ }^{5}$-polarizer pair (in older instruments a Pockels cell) modulates the polarization of the incident light at a typical frequency of 50 kHz . Typically CD spectropolarimeters use a double prism monochromator rather than one based on gratings, because they tend to perform better in the UV. The radiation source in UV-visible CD spectrometers is a high energy ( $150-450 \mathrm{~W}$ ) xenon arc lamp. In some instruments the lamp in water cooled. Sometimes, as a result of the high energy lamps used, the optics of the instruments are purged with nitrogen gas to avoid ozone being created and reacting with the surfaces of the optical elements. Nitrogen purging is also necessary in the sample compartment for measurements running below 200 nm to avoid absorption by ozone of UV radiation.

Among the available commercial designs of dicrographs those from the Japanese manufacturer $\mathrm{Jasco}^{6}$ are probably the ones present in a larger number of laboratories. In this work when we mention in general mention commercial instruments or standard measurements of optical activity we will be implicitly referring measurements obtained with Jasco instruments.

The most used optical arrangement that commercial instruments use for CD measurement is the sequence: polarizer, phase modulator, sample and detector. In some instruments the measurement of CB is also possible through the addition of an "accessory"(a polarizer) between the sample and the detector resulting in the sequence: polarizer, phase modulator, sample, polarizer and detector. We should point out that these optical setups are very similar or even identical to the ones one might find in spectroscopic phase-modulated ellipsometers. In spite of these evident similarities it seems that CD and CB techniques and ellipsometric techniques have progressed in parallel but with little communication between the two communities. Our feeling is that, in general, there is no consciousness about the important similarities between dichrographs and ellipsometers, and this avoids technical exchanges between both communities that otherwise could be very valuable. This intercommunication will become even more justified in

[^10]the future because, from a technical standpoint, our opinion is both communities will share the interest of measuring complete Mueller matrices.

CD spectrometers are usually designed to work with solutions contained in cuvettes. For UV-visible CD, high quality fused quartz that transmits a broad UV-visible wavelength range is required. The default cuvette pathlength is 1 cm , but for samples with large absorbance signal the use of shorter pathlength cells ( $1 \mathrm{~mm}, 0.1 \mathrm{~mm}$ ) is also common. In the 1950 s there were commercial CB spectropolarimers but they are no longer available, and now the CB capability is typically offered as an add-on to a CD spectropolarimeter.

### 2.5. The concern of anisotropic media in chemistry

The problem of measuring optical activity in systems coexisting circular and linear anisotropies is a long-term concern in chemistry. As far back as 1969 Disch an Sverdlik [38] realized that their method to record CD spectra was influenced by linear dichroism an linear birefringence ${ }^{7}$.Due to this influence several authors use the terminology true $C D$ to refer to the circular dichroism of the system under study, which may not correspond to the measured $C D$. The contributions that cause this miscorrespondence are commonly referred as artifacts, or, more specifically, as chiroptical artifacts and may occur due to the presence of linear dichroism and linear birefringence in the medium or due to the non-ideal behaviour of the instrument used to perform the measurement (uncorrected strain of the optical elements for example). In the next chapter we will introduce the theory that justifies the apparition of these artifacts; in this section we will only introduce some approaches that have been proposed to sort this difficulty.

Since the introduction of CD measurement techniques the artifact problem has received a permanent interest from the chemistry communite [39-41]. Several different instruments or measurement techniques have been proposed to sort this problem but, probably, none of them has yet been widely recognized or has transformed into a commercial measurement system. In this section we will critically review some of the most significant contributions published in this field.

One of the most important works on the subject, which is the reference for several other following works, is the excellent review of Schellmann and Jensen published in 1987 [41]. In this work they extensively review every aspect related to the measurement of CD and CB in presence of linear dichroism and linear birefringence. They introduce the Jones approach to describe anisotropic homogeneous media and they suggest instrumental configurations that could be used

[^11]in the measurements. This work compiled the results of several other previous publications by the authors $[42,43]$ and introduced the notation based on the parameters $\mathrm{CD}, \mathrm{CB}, \mathrm{LD}, \mathrm{LB}, \mathrm{LD}^{\prime}$ and $\mathrm{LB}^{\prime}$ to designate the optical effects described in section 1.5 that we also use in this thesis.

Japanese authors are among the ones that have dedicated more attention to the problem from an instrumental point of view. Group of Y. Shindo was pioneer in determining the problems that presented commercial CD spectrometers [39,44, $45]$ and proposed specific modifications for them [46,47]. More recently the group of R. Kuroda (also in collaboration with group of Shindo) has made significant contributions to the measurement of CD in solid samples [48-50] -which tend to show linear birefringence and linear dichroism- and, in collaboration with Jasco, the Japanese manufacturer of CD spectrometers, they have developed prototypes based on modifications of commercial instruments [51,52]. However the proposed measurement methods seem to be only valid for specific types of homogenous solid samples because the require the rotation of the samples. As their measurements continue to be restricted to the determination of few elements of the Mueller matrix, they do not provide a complete characterization of the optical behavior of the sample. In Ref. [53] one can find a recent review, form a chemistry perspective, of the different approaches developed to measure CD in solid state samples.

Several of the most interesting recent approaches to the problem have come from the field of crystallography. In 1983 Kobayashi and Uesu [54] invented an apparatus to study the optical activity of crystals in directions out of the optic axis. This instrument was called HAUP - after "High accuracy universal polarimeter" - and since then similar instruments have been used in a number of laboratories mainly to study crystals [55-58]. W. Kaminsky have recently proposed and developed several innovating measurements for various tensorial properties of crystals including the gyrotropy that gives rise to optical activity [59-63]. One might also find in Ref . [64] a comprehensive review about the measurement of optical activity in crystals.

Current work on the subject is also directed towards circular dichroism and circular birefringence imaging techniques which permit the study of heterogenous samples along their surface. In 1982, Maestre and Katz adapted a commercial Cary spectropolarimeter to a microscope [65] and performed several measurements. However, they faced important artifacts arising from the optical components of the instrument. The group of Bart Kahr has extensively worked on polarimetric imaging techniques [66] applied to crystal optics and, more recently, they have started using Mueller matrix imaging polarimetry coupled with microscope optics to study chiroptical phenomena in crystals [67,68].

## Part II.

## Theory

## Chapter 3.

## Light propagation in anisotropic optical active media

The description of light propagation in anisotropic optical active media ${ }^{1}$ can be highly complex. Maxwell did not treat chiral media and, in comparison with the vast amount of references to Maxwell's equations, one may find in the literature only a limited number or books and papers dealing with the resolution of Maxwell equations in anisotropic optical active media.

Optical activity is a clear manifestation of the molecular or atomic structure of matter. The circular birefringence or the rotation of the polarization's direction per unit thickness of light propagating in an optically-active material scales as the ratio $d / \lambda^{2}$, where $d$ is some typical molecular length and $\lambda$ is the light wavelength. In fact optical activity is due to the inhomogeneous nature of the medium, but as $d$ is usually much smaller than $\lambda$ we can effectively use homogeneous models to describe optical activity.

By using homogenous models we can distinguish two different approaches to study light propagation through anisotropic optically active media. One consist in solving Maxwell equations with adequate tensorial constitutive relations. The calculations corresponding to this method are rather complicated and several elegant matrix formalism have been developed to systematize and simplify the calculations, being the most well-known the Berreman method [69]. Even with these tools, to our knowledge analytic solutions are only available either for anisotropic but non-gyrotropic media [70] or for gyrotropic but isotropic media [71]. No analytic expressions have been found for the rest of cases but they can be treated numerically. The second approach is semi-phenomenological and it was introduced by Jones in the framework of his formalism to represent polarized light. It is based on an infinitesimal representation of the medium using a differential matrix calculus. In this chapter we will briefly introduce the Berreman method, and we will revisit in detail the method by Jones, as most of the results of this work have been studied using this method.

[^12]
### 3.1. Constitutive Equations. Introduction to Berreman matrix formalism

Different authors have been using different forms for the constitutive equations ${ }^{2}$ of chiral media. Here we will present a constitutive equations based on those proposed by Condon [72], who noted that the essential feature of chiral media is that part of the polarization $\mathbf{P}$ that is dependent on $\partial \mathbf{H} / \partial t$ the part of the magnetization $\mathbf{M}$ that is dependent on $\partial \mathbf{E} / \partial t$ :

$$
\begin{align*}
\mathbf{D} & =\varepsilon \mathbf{E}-\mathbf{g} \frac{\partial \mathbf{H}}{\partial t}  \tag{3.1a}\\
\mathbf{B} & =\boldsymbol{\mu} \mathbf{H}+\mathbf{g} \frac{\partial \mathbf{E}}{\partial t} \tag{3.1b}
\end{align*}
$$

where $\mathbf{g}$ is the tensor of gyrotropy. Although the derivation of these relations was heuristic and not particularly rigorous, microscopic models have been presented that support these equations. For monocromatic waves with a $e^{i w t}$ time dependence of the fields they can be rewritten as

$$
\begin{align*}
\mathbf{D} & =\varepsilon \mathbf{E}+\rho \mathbf{H}  \tag{3.2a}\\
\mathbf{B} & =\mu \mathbf{H}+\rho^{\prime} \mathbf{E} \tag{3.2~b}
\end{align*}
$$

in which $\boldsymbol{\rho}=-i \omega \mathbf{g}$ and $\boldsymbol{\rho}^{\prime}=i \omega \mathbf{g}$. Upon the literature one might find different names for these equations, being probably the most adequate Drude-CondonFedorov constitutive relations, as these three authors made contributions to this formulation. For anisotropic media $\boldsymbol{\varepsilon}, \boldsymbol{\mu}$ and $\boldsymbol{\rho}$ are tensors, while for an isotropic optically active medium they simplify to scalar (a pseudoscalar in the case of $\boldsymbol{\rho}$.).

All these complex-valued tensor, e. g. the permittivity, can be expressed in laboratory Cartersian coordinates $(x, y, z)$ as :

$$
\varepsilon=\left[\begin{array}{lll}
\varepsilon_{x x} & \varepsilon_{x y} & \varepsilon_{x z}  \tag{3.3}\\
\varepsilon_{y x} & \varepsilon_{y y} & \varepsilon_{y z} \\
\varepsilon_{z x} & \varepsilon_{z y} & \varepsilon_{z z}
\end{array}\right]
$$

It is interesting to note that for a symmetric permittivity tensor it is possible to perform a coordinate transformation from the laboratory $(x, y, z)$ coordinates to another Cartesian coordinate system $\left(x^{\prime}, y^{\prime}, z^{\prime}\right)$ characteristic of the anisotropic medium (its natural basis), for which the tensor becomes diagonal.

$$
\boldsymbol{\varepsilon}=\mathbf{A}\left[\begin{array}{ccc}
\varepsilon_{x^{\prime}} & 0 & 0  \tag{3.4}\\
0 & \varepsilon_{y^{\prime}} & 0 \\
0 & 0 & \varepsilon_{z^{\prime}}
\end{array}\right] \mathbf{A}^{-1}
$$

[^13]where $\mathbf{A}$ is a unitary rotation matrix that depends on the three Euler angles that can be found somewhere else [7]. This equation has fundamental importance in crystal optics, because, for example, it makes the distinction between uniaxial and biaxial crystals depending on if the components of the diagonalized permittivity tensor are all different (biaxial) or two of them are equal (uniaxial). If all of them are equal we have the isotropic case we described in Chapter 1. It is also important to stress that the $\rho$ tensor may not have the same form as the dielectric tensor. And since we focus our attention in nonmagnetic materials $\mu$ is the identity matrix.

The Eqs. (3.2) can be casted in matrix form:

$$
\left[\begin{array}{l}
\mathbf{D}  \tag{3.5}\\
\mathbf{B}
\end{array}\right]=\mathbf{M}_{o}\left[\begin{array}{c}
\mathbf{E} \\
\mathbf{H}
\end{array}\right], \quad \mathbf{M}_{o}=\left[\begin{array}{cc}
\varepsilon & \rho \\
\rho^{\prime} & \boldsymbol{\mu}
\end{array}\right]
$$

where $\mathbf{M}_{o}$ is a $6 \times 6$ matrix called the optical matrix.
Now we are ready to briefly introduce the Berreman method [69]. It uses transverse components of the electric and magnetic field vectors and thus, in a first term, it is not intuitively related to the experiment and requires some steps of calculations. We will not deal with this subject in detail, we just outline the most significative calculations it involves. It can be described as a re-formulation of Maxwell's equations and it is given by,

$$
\begin{equation*}
\frac{d \boldsymbol{\Psi}}{d z}=-i \boldsymbol{\Delta} \Psi \tag{3.6}
\end{equation*}
$$

where $\boldsymbol{\Psi}^{T}=\left(E_{x}, H_{y}, E_{y},-H_{x}\right)$ and the z-direction is perpendicular to the sample surface. $\boldsymbol{\Delta}$ is a $4 \times 4$ complex matrix that contains all the information about the dielectric functions of the medium. The elements of the matrix $\boldsymbol{\Delta}$ are functions of the elements of the optical matrix $\mathbf{M}_{o}$ and the direction of the wave normal of the incident wave. The explicit relations can be found in Refs [69, 73]. If $\mathbf{M}_{o}$ (and thus $\boldsymbol{\Delta}$ ) is constant independent of $z$ over a continuous interval Eq. (3.6) is integrable and yields:

$$
\begin{equation*}
\Psi(z+h)=\mathbf{L}(h) \Psi(z), \quad \text { where } \quad \mathbf{L}(h)=e^{-i \omega h \Delta} \tag{3.7}
\end{equation*}
$$

$\mathbf{L}(h)$ is the layer matrix that relates the vector $\Psi$ at two different parallel planes, separated by a distance $h$, in a homogeneous anisotropic medium whose fields are excited by an incident plane wave. The case where $\mathbf{M}_{o}$ depends on $z$ can be treated similarly if the medium is divided into layers that are thin enough to make $\mathbf{M}_{o}$ independent of $z$ within each layer

The main difficulty of the Berreman calculus remains in the obtention of the layer matrix, i. e. in the calculation of the exponential of the matrix $\boldsymbol{\Delta}$. An analytical form of the layer matrix has only been found for a limited number of forms of $\boldsymbol{\Delta}$, for the rest of cases the layer matrix can be numerically computed.

Analytic expressions of the layer matrix (sometimes also referred as characteristic matrix or transfer matrix) can be found in the works of Berreman [69], Wöhler [74], Schubert [70], etc.

The Berreman method permits the calculation of the optical properties at oblique incidence and also takes into account multiple reflections at the interfaces. It is also an appropriate and flexible technique to deal with layered systems since it can be seen as a generalization of the Abelès $2 \times 2$ matrix method for layered isotropic media. In spite of all these virtues this matrix method is perhaps not as used as one might think. In a first moment it was mainly used to study the optics of liquid crystals and only lately has gained much attention in the field of generalized ellipsometry. The reason probably is that it is a method of rather difficult application (at least algebra-wise) if one wants to retrieve analytical solutions, and for many problems, specially those involving gyrotropic media, the solutions can only be treated numerically. Also in other fields, a clear example is in crystal optics, there exists "traditional" analytic expressions, sometimes derived phenomenologically, that are useful to study particular problems of crystal optics and that nearly cover every situation that a experimentalist can find.

### 3.2. The lamellar representation

Jones solved a number of years ago the general problem of the polarization description of a medium possessing eight different optical parameters [75]. These parameters correspond to the eight physical measurements one can make on such a system with a given light path; namely: mean absorption, mean refraction, optical rotation, circular dichroism, and two parameters for linear birefringence and two more for linear dichroism. He studied these properties in terms of a "layered"medium interpretation that, by means of a differential equation analysis, leads to a exponential representation of the Jones matrix. Years later R.M.A. Azzam [76] developed and analogous infinitesimal calculus based on Mueller matrices that can be used for the same purposes.

The use of the differential matrix calculus introduced by Jones to describe the continuous propagation of partially polarized light through linear anisotropic media is somewhat less general that the Berreman method. As it will be presented, this method is only described for normal incidence and it does not take into account multiple reflections at the layer interfaces. However, this method is specially useful to study light transmission through a homogeneous anisotropic medium with optical activity, because it is easy to apply and yields analytic results that can be directly related to the experiment. The approach for this method is quite different from the Berreman method: instead of using the constitutive relations between the different electromagnetic elements as basic elements of the calculus, it uses the optical effects that such medium exhibits as the basis


Figure 3.1.: According to the Jones lamellar representation a thin slab of the medium, $\Delta z$, can be considered as the sum eight elemental subslabs, $\Theta_{k}$, each of them corresponding to only one optical property. Together they form the so-called $\mathbf{N}$-matrix. Figure adapted from Ref. [77].
set for the construction of the model.
The Jones matrix $\mathbf{J}$ governing an optical system can be written in exponential form [78]:

$$
\begin{equation*}
\mathbf{J}(\omega, z)=\exp [z \mathbf{N}(\omega)], \tag{3.8}
\end{equation*}
$$

where $\omega$ is the frequency and $z$ is the distance into the medium ( $z=0$ denotes the origin of the scattering medium). The matrix $\mathbf{N}$ is the infinitesimal generator of $\mathbf{J}$, and satisfies:

$$
\begin{equation*}
\mathbf{N}(\omega)=\lim _{z \rightarrow 0} \frac{\mathbf{J}(\omega, z)-\mathbf{I}}{z} \tag{3.9}
\end{equation*}
$$

Note that the matrix $\mathbf{N}$ no longer depends on $z$; also in future equations we will omit everywhere the explicit dependence in $\omega$ and $z$. The $\mathbf{N}$-matrix can be also understood as the operator that gets $d \mathbf{J} / d z$ from $\mathbf{J}$ :

$$
\begin{equation*}
\frac{d \mathbf{J}}{d z}=\mathbf{N J} \tag{3.10}
\end{equation*}
$$

Chapter 3. Light propagation in anisotropic optical active media

According to Jones lamellar representation, a thin slab of a given medium is equivalent to a stack of eight subslabs each possessing one and only one of the eight fundamental optical effects listed in Table 1.1. Each one of these effects on this infinitesimal layer was described by Jones with a matrix $\boldsymbol{\Theta}_{k}$; namely $\boldsymbol{\Theta}_{0}$ for the isotropic refraction, $\boldsymbol{\Theta}_{1}$ for the isotropic absorption, $\boldsymbol{\Theta}_{2}$ for the horizontal linear dichroism, $\boldsymbol{\Theta}_{3}$ for the horizontal linear birefringence, $\boldsymbol{\Theta}_{4}$ for the $45^{\circ}$ linear dichroism, $\boldsymbol{\Theta}_{5}$ for the $45^{\circ}$ linear birefringence, $\boldsymbol{\Theta}_{6}$ for the $45^{\circ}$ circular birefringence and $\boldsymbol{\Theta}_{7}$ for the circular birefringence. Each one of this eight matrices can be calculated applying Eq. (3.9) on the corresponding the Jones matrices given in table 1.1. In the limit of infinitesimal thickness, the trigonometric, hyperbolic and exponential functions of the Jones matrices of table 1.1 can be taken to first order and every $\boldsymbol{\Theta}_{k}$ gets a very simple form. Then the N-matrix can be then calculated by adding all $\Theta_{k}$ matrices [75]:

$$
\begin{equation*}
\mathbf{N}=\sum_{k=0}^{7} \boldsymbol{\Theta}_{k} \tag{3.11}
\end{equation*}
$$

The most general expression of the $\mathbf{N}$ matrix $\mathbf{N}$ for a infinitesimal layer containing all the eight basic optical properties introduced in section 1.5 is:

$$
\mathbf{N}=\frac{1}{2 z}\left(\begin{array}{cc}
-\mathrm{LD}-i \mathrm{LB}-2(i \eta+\kappa) & -\mathrm{LD}^{\prime}-i \mathrm{LB}^{\prime}+\mathrm{CB}-i \mathrm{CD}  \tag{3.12}\\
-\mathrm{LD}^{\prime}-i \mathrm{LB}^{\prime}-\mathrm{CB}+i \mathrm{CD} & \mathrm{LD}+i \mathrm{LB}-2(i \eta+\kappa)
\end{array}\right) .
$$

where we divide by $z$ to drop the dependence on the slab thickness that is implicit in the optical effects as they were defined in Table 1.1. Note also that the notation we use is mostly the one introduced by Schellman, Jensen and Troxell [41, 42], which slightly differs from the notation used by Jones. The notation given in table 1.1 is appropriate to keep the theoretical discussion close to the experiment, because these basic retardances and absorbances correspond to standard experimental quantities which are related to eight measurable physical effects.

Note that, for example, the matrix $\mathbf{N}$ associated with LB commutes with the matrix $\mathbf{N}$ associated with LD but not with the matrix associated to $\mathrm{CB}, \mathrm{CD}$, $\mathrm{LB}^{\prime}$ or $\mathrm{LD}^{\prime}$. The same can also be verified for the others: they commute with their partner of Eqs. (1.55) but not with the ones that appear in the other lines. This means that, for example, a system that has both LD and LB can be simulated by a dichroic sample and a birefringent sample no matter which order they are placed. Conversely, for a system with LD and $\mathrm{LB}^{\prime}$ this is not true. This question will be further discussed in section 4.3.2. It is useful to in the $\mathbf{N}$ matrix the general retardations defined in Eqs. (1.55). With these definitions $\mathbf{N}$ can be rewritten as:

$$
\mathbf{N}=\frac{-i}{2 z}\left(\begin{array}{cc}
\chi+\mathrm{L} & \mathrm{~L}^{\prime}+i \mathrm{C}  \tag{3.13}\\
\mathrm{~L}^{\prime}-i \mathrm{C} & \chi-\mathrm{L}
\end{array}\right)
$$

Using equation (3.13) in (3.8) we obtain:

$$
\mathbf{J}=\exp [-i \mathbf{R}] \quad \text { where } \quad \mathbf{R}=\frac{1}{2}\left(\begin{array}{cc}
\chi+\mathrm{L} & \mathrm{~L}^{\prime}+i \mathrm{C}  \tag{3.14}\\
\mathrm{~L}^{\prime}-i \mathrm{C} & \chi-\mathrm{L}
\end{array}\right)
$$

And if we define a generalized retardance vector $\vec{T} \equiv\left(L, L^{\prime},-C\right)$ we can write $\mathbf{R}$ in a compact form:

$$
\begin{equation*}
\mathbf{R}=\frac{1}{2}\left(\chi \boldsymbol{\sigma}_{0}+\overrightarrow{\boldsymbol{\sigma}} \cdot \overrightarrow{\mathrm{T}}\right) \tag{3.15}
\end{equation*}
$$

where $\boldsymbol{\sigma}_{0}$ is a $2 \times 2$ identity matrix, $\overrightarrow{\boldsymbol{\sigma}}=\left(\boldsymbol{\sigma}_{1}, \boldsymbol{\sigma}_{2}, \boldsymbol{\sigma}_{3}\right)$, and $\boldsymbol{\sigma}_{1}, \boldsymbol{\sigma}_{2}$ and $\boldsymbol{\sigma}_{3}$ are the three Pauli spin matrices:

$$
\begin{array}{ll}
\boldsymbol{\sigma}_{0}=\left[\begin{array}{ll}
1 & 0 \\
0 & 1
\end{array}\right], \quad \boldsymbol{\sigma}_{1}=\left[\begin{array}{cc}
1 & 0 \\
0 & -1
\end{array}\right],  \tag{3.16}\\
\boldsymbol{\sigma}_{2}=\left[\begin{array}{ll}
0 & 1 \\
1 & 0
\end{array}\right], \quad \boldsymbol{\sigma}_{3}=\left[\begin{array}{cc}
0 & -i \\
i & 0
\end{array}\right] .
\end{array}
$$

Using this definition, eq. (3.14) transforms in

$$
\begin{equation*}
\mathbf{J}=\exp [-i \mathbf{R}]=e^{-i \chi / 2} e^{-i(\overrightarrow{\boldsymbol{\sigma}} \cdot \overrightarrow{\mathrm{~T}}) / 2}=e^{-i \chi / 2} e^{-i\left(\overrightarrow{\boldsymbol{\sigma}} \cdot \overrightarrow{\mathrm{~T}}_{R}+\overrightarrow{\boldsymbol{\sigma}} \cdot\left(-i \overrightarrow{\mathrm{~T}}_{D}\right)\right) / 2} \tag{3.17}
\end{equation*}
$$

On the right hand side we have written $\overrightarrow{\mathrm{T}}$ as the sum of two vectors: $\overrightarrow{\mathrm{T}}=$ $\overrightarrow{\mathrm{T}}_{R}-i \overrightarrow{\mathrm{~T}}_{D}$ where $\overrightarrow{\mathrm{T}}_{R} \equiv\left(\mathrm{LB}, \mathrm{LB}^{\prime},-\mathrm{CB}\right)$ contains only dispersive terms and $\overrightarrow{\mathrm{T}}_{D} \equiv$ ( $\mathrm{LD}, \mathrm{LD}^{\prime},-\mathrm{CD}$ ) is formed by absorptive terms.

The Jones matrix $\mathbf{J}$ can also be written in terms of trigonometric functions:

$$
\begin{equation*}
\mathbf{J}=e^{-i \chi / 2} \exp \left(-i \frac{\mathrm{~T}}{2} \frac{\overrightarrow{\mathrm{~T}}}{\mathrm{~T}} \cdot \overrightarrow{\boldsymbol{\sigma}}\right)=e^{-i \chi / 2}\left[\boldsymbol{\sigma}_{0} \cos \frac{\mathrm{~T}}{2}-\frac{i}{\mathrm{~T}} \overrightarrow{\mathrm{~T}} \cdot \overrightarrow{\boldsymbol{\sigma}} \sin \frac{\mathrm{~T}}{2}\right] \tag{3.18}
\end{equation*}
$$

where $\overrightarrow{\mathrm{T}}=\left(\mathrm{L}, \mathrm{L}^{\prime},-\mathrm{C}\right)$ and T is the module of this vector $\left(\mathrm{T}=\sqrt{\mathrm{L}^{\prime 2}+\mathrm{C}^{2}+\mathrm{L}^{2}}\right)$. Using complete matrix notation (3.18) can be rewritten as:

$$
\mathbf{J}=e^{-i \chi / 2}\left(\begin{array}{cc}
\cos \frac{\mathrm{T}}{2}-\frac{i \mathrm{~L}}{\mathrm{~T}} \sin \frac{\mathrm{~T}}{2} & \frac{\left(\mathrm{C}-i \mathrm{~L}^{\prime}\right)}{\mathrm{T}} \sin \frac{\mathrm{~T}}{2}  \tag{3.19}\\
-\frac{\left(\mathrm{C}+i \mathrm{~L}^{\prime}\right)}{\mathrm{T}} \sin \frac{\mathrm{~T}}{2} & \cos \frac{\mathrm{~T}}{2}+\frac{i \mathrm{~L}}{\mathrm{~T}} \sin \frac{\mathrm{~T}}{2}
\end{array}\right) .
$$

The infinitesimal method reviewed in this section uses the the eight optical effects given in Table 1.1 as a basis set for constructing the properties of any complex optical element. In contrast, the Berreman method presented before uses the dielectric and gyration tensors of the medium as a basis set. Although both methods are constructed from a different perspective it should be possible to correlate both basis sets, and, in fact, this was attempted by Jones in last paper of his series [79]. Future work should be directed towards revising this paper, that otherwise has remained quite overlooked, to strengthen the connections between both methods.

Chapter 3. Light propagation in anisotropic optical active media

### 3.2.1. Calculation of the Mueller-Jones matrix

Our purpose here is to find the Mueller-Jones matrix corresponding to the Jones matrix of Eq. (3.19). This Mueller matrix can be directly found applying the transformation relationship of Eq. (1.29) to our Jones matrix. Alternatively, we can also use calculate the elements of the Mueller matrix, $m_{i j}$, one by one using the following expression:

$$
\begin{equation*}
m_{i j}=\frac{1}{2} \operatorname{Tr}\left(\sigma_{i} \mathbf{J} \sigma_{j} \mathbf{J}^{\dagger}\right) \tag{3.20}
\end{equation*}
$$

where $\mathbf{J}^{\dagger}$ is the conjugate transpose of $\mathbf{J}$. With any of the two possible approaches after some lengthly algebra [80] we find:

$$
\begin{align*}
& \mathbf{M}=e^{-2 \kappa}\left[\begin{array}{rr}
X & -U \mathrm{LD}-V \mathrm{BB} \\
+(W / 2)\left(\overrightarrow{\mathrm{T}^{*}} \cdot \overrightarrow{\mathrm{~T}}\right) & +W\left(\mathrm{CBLD}^{\prime}-\mathrm{LB}^{\prime} \mathrm{CD}\right) \\
-U \mathrm{LD}-V \mathrm{LB} & X+W\left(\mathrm{LD}^{2}+\mathrm{LB}^{2}\right) \\
-W\left(\mathrm{CBLD}^{\prime}-\mathrm{LB}^{\prime} \mathrm{CD}\right) & -(W / 2)\left(\overrightarrow{\mathrm{T}}^{*} \cdot \overrightarrow{\mathrm{~T}}\right) \\
-U \mathrm{LD}^{\prime}-V \mathrm{LB}^{\prime} & -U \mathrm{CB}+V \mathrm{CD} \\
-W\left(\mathrm{LBCD}^{\prime}-\mathrm{CBLD}\right) & +W\left(\mathrm{LBLB}^{\prime}+\mathrm{LDLD}^{\prime}\right) \\
U \mathrm{CD}^{\prime}+V \mathrm{CB} & -U L B^{\prime}+V \mathrm{LD}^{\prime} \\
-W\left(\mathrm{LBLD}^{\prime}-\mathrm{LB}^{\prime} \mathrm{LD}\right) & -W\left(\mathrm{CBLB}^{\prime}+\mathrm{CDLD}\right)
\end{array}\right. \\
& \begin{array}{rr}
-U \mathrm{LD}^{\prime}-V \mathrm{LB}^{\prime} & U \mathrm{CD}+V \mathrm{CB} \\
+W(\mathrm{LBCD}-\mathrm{CBLD}) & +W\left(\mathrm{LBLD}^{\prime}-\mathrm{LB}^{\prime} \mathrm{LD}\right)
\end{array} \\
& U C B-V C D \quad U \mathrm{LB}^{\prime}-V \mathrm{LD}^{\prime} \\
& +W\left(\mathrm{LBLB}^{\prime}+\mathrm{LDLD}^{\prime}\right) \quad-W(\mathrm{CBLB}+\mathrm{CDLD}) \\
& X+W\left(\mathrm{LB}^{\prime 2}+\mathrm{LD}^{\prime 2}\right) \quad-U \mathrm{LB}+V \mathrm{LD}  \tag{3.21}\\
& -(W / 2)\left(\overrightarrow{\mathrm{T}}^{*} \cdot \overrightarrow{\mathrm{~T}}\right) \quad-W\left(\mathrm{CBLB}^{\prime}+\mathrm{CDLD}^{\prime}\right) \\
& U L B-V L D \quad X+W\left(\mathrm{CD}^{2}+\mathrm{CB}^{2}\right) \\
& \left.-W\left(\mathrm{CBLB}^{\prime}+\mathrm{CDLD}^{\prime}\right) \quad-(W / 2)\left(\overrightarrow{\mathrm{T}}^{*} \cdot \overrightarrow{\mathrm{~T}}\right) \quad\right]
\end{align*}
$$

where

$$
\begin{gather*}
U=\left(\mathrm{T}^{i} \sinh \mathrm{~T}^{i}+\mathrm{T}^{r} \sin \mathrm{~T}^{r}\right) / \mathrm{TT}^{*} \\
V=\left(\mathrm{T}^{i} \sin \mathrm{~T}^{r}-\mathrm{T}^{r} \sinh \mathrm{~T}^{i}\right) / \mathrm{TT}^{*} \\
W=\left(\cosh \mathrm{T}^{i}-\cos \mathrm{T}^{r}\right) / \mathrm{TT}^{*}  \tag{3.22}\\
X=\left(\cosh \mathrm{T}^{i}+\cos \mathrm{T}^{r}\right) / 2
\end{gather*}
$$

where $\mathrm{T}^{r}$ is the real part of T and $\mathrm{T}^{i}$ is the imaginary part of $\mathrm{T}\left(\mathrm{T}=\sqrt{\mathrm{L}^{\prime 2}+\mathrm{C}^{2}+\mathrm{L}^{2}}\right)$. This Mueller matrix was first given by Jensen, Schellman and Troxell in 1978 [42],
and it was rewritten in a more convenient way by Shellman and Jensen in 1987 [41] using the same $U, V, W$ and $X$ functions that we have introduced. However, note that the order of the elements of the Mueller matrix given in [41] differs from this one, because a different order of the Stokes parameters has been used.


Figure 3.2.: Values of $U, V$ and $W$ as a function of $T^{r}$ (the real part of total complex retardance $\mathrm{T}=\sqrt{\mathrm{L}^{\prime 2}+\mathrm{C}^{2}+\mathrm{L}^{2}}$ ). The imaginary part, $\mathrm{T}^{i}$, has been fixed to 0.2 rad. In the limit of small anisotropies, we obtain small values of $\left|\mathrm{T}^{r}\right|$ (tipically $<0.5 \mathrm{rad}$ ) and $U, W$ and $V$ can be respectively approximated as $1,1 / 2$ and 0 .

The matrix appearing in eq. (3.21) is rather complex, but it can be easily simplified assuming small values for the anisotropic effects [81], as often happens for thin specimens. In this case the parameters $U, V W$ and $X$ of eq. (3.22) tend, respectively, to $1,0,1 / 2$ and 1 , as it is shown in Fig. 3.2, and the following simplification can be rather useful:

Chapter 3. Light propagation in anisotropic optical active media

$$
\begin{align*}
& \mathbf{M}=e^{-2 \kappa}\left[\begin{array}{cc}
1 & -\mathrm{LD} \\
+\frac{1}{2}\left(\mathrm{LD}^{\prime 2}+\mathrm{LD}^{2}\right) & \\
-\mathrm{LD} & +\frac{1}{2}\left(\mathrm{LD}^{2}-\mathrm{LB}^{\prime 2}\right) \\
-\mathrm{LD}^{\prime} & +\frac{1}{2}\left(\mathrm{LBLB}^{\prime}+\mathrm{LDLD}^{\prime}\right) \\
& \\
\mathrm{CD}^{\prime} & -\mathrm{LB}^{\prime}
\end{array}\right. \\
& -\mathrm{LD}^{\prime} \\
& \begin{array}{c}
\mathrm{CD} \\
+\frac{1}{2}\left(\mathrm{LBLD}^{\prime}-\mathrm{LB}^{\prime} \mathrm{LD}\right)
\end{array} \\
& \text { CB } \\
& +\frac{1}{2}\left(\text { LBLB }^{\prime}+\text { LDLD }^{\prime}\right)  \tag{3.23}\\
& \begin{array}{c}
\stackrel{1}{2}\left(\mathrm{LD}^{\prime 2}-\mathrm{LB}^{2}\right)
\end{array} \\
& \text { LB } \\
& 1 \\
& \left.-\frac{1}{2}\left(\mathrm{LB}^{2}+\mathrm{LB}^{\prime 2}\right) \quad\right]
\end{align*}
$$

This matrix indicates that for small values of the anisotropic effects the elements of the antidiagonal $\left(m_{03}, m_{12}, m_{21}\right.$ and $\left.m_{30}\right)$ are the most important for the study of optical activity: $\left(m_{03}+m_{30}\right) / 2$ can be a measure of CD and CB can be obtained from $\left(m_{12}-m_{21}\right) / 2$. In this particular situation the linear effects can be directly "read" from the elements of the Mueller matrix (i.e. $-m_{01}$ is LD, $-m_{02}$ is $\mathrm{LD}^{\prime}$, etc) .

Commercial CD spectropolarimeters take the CD values directly from a Mueller matrix element (either $m_{03}$ or $m_{30}$ ). However, from the presented analysis is easy to realize that 3 conditions must be fulfilled for this values to be "real" measurements of CD:

- the light beam must go through an homogeneous medium,
- all the optical effects (birefringences and dichroism) must be small enough to permit the simplification that results in matrix of Eq. (3.23),
- the factor $\frac{1}{2}\left(\mathrm{LBLD}^{\prime}-\mathrm{LB}^{\prime} \mathrm{LD}\right)$ must vanish.

These condition seem very restrictive, but fortunately for most of the standard samples that are measured in chemistry, biology, pharmacy, etc the conditions are fulfilled. This is the reason why commercial instruments usually work well.

The mathematical characterization of the general Mueller matrix of a homogeneous non-depolarizing media has an important interest for experimentalist because it allows the physical interpretation of the information that provides the sixteen elements of the Mueller matrix. Two interesting Mueller matrices can be straightforward derived from Eq. (3.21). Imposing $\mathrm{CB}=\mathrm{LB}=\mathrm{LB}^{\prime}=0$ we obtain the Mueller matrix of a general diattenuator, $\mathbf{M}_{D}$, i.e. the Mueller matrix of an optical element that shows no birefringence or retardation. With the condition $\mathrm{CD}=\mathrm{LD}=\mathrm{LD}^{\prime}=0$ the Mueller matrix of a general retarder, $\mathbf{M}_{R}$, an optical element that has no dichroism and diattenuation ${ }^{3}$ :

$$
\mathbf{M}_{R}=\left[\begin{array}{cccc}
1 & 0 & 0 & 0  \tag{3.24}\\
0 & \cos \mathrm{~T}_{R}+\mathrm{LB}^{2} \alpha & \mathrm{LBLB}^{\prime} \alpha+\mathrm{CB} \beta & -\mathrm{LBCB} \alpha+\mathrm{LB}^{\prime} \beta \\
0 & \mathrm{LBLB}^{\prime} \alpha-\mathrm{CB} \beta & \cos \mathrm{~T}_{R}+\mathrm{LB}^{\prime 2} \alpha & -\mathrm{LB}^{\prime} \mathrm{CB} \alpha-\mathrm{LB} \beta \\
0 & -\mathrm{LBCB}^{2}-\mathrm{LB}^{\prime} \beta & -\mathrm{LB}^{\prime} \mathrm{CB} \alpha+\mathrm{LB} \beta & \cos \mathrm{~T}_{R}+\mathrm{CB}^{2} \alpha
\end{array}\right],
$$

with $\alpha=\left(1-\cos \mathrm{T}_{R}\right) / \mathrm{T}_{R}^{2}, \beta=\sin \mathrm{T}_{R} / \mathrm{T}_{R}$ and $\mathrm{T}_{R}=\sqrt{\mathrm{LB}^{2}+\mathrm{LB}^{\prime 2}+\mathrm{CB}^{2}}$.

$$
\mathbf{M}_{D}=e^{-2 k}\left[\begin{array}{cccc}
\cosh \mathrm{T}_{D} & -\mathrm{LD} \nu & -\mathrm{LD}^{\prime} \nu & \mathrm{CD} \nu  \tag{3.25}\\
-\mathrm{LD} \nu & 1+\mathrm{LD}^{2} \mu & \mathrm{LDLD}^{\prime} \mu & -\mathrm{LDCD} \mu \\
-\mathrm{LD}^{\prime} \nu & \mathrm{LDLD}^{\prime} \mu & 1+\mathrm{LD}^{\prime 2} \mu & -\mathrm{LD}^{\prime} \mathrm{CD} \mu \\
\mathrm{CD} \nu & -\mathrm{LDCD}^{2} & -\mathrm{LD}^{\prime} \mathrm{CD} \mu & 1+\mathrm{CD}^{2} \mu
\end{array}\right]
$$

with $\mu=\left(\cosh \mathrm{T}_{D}-1\right) / \mathrm{T}_{D}^{2}, \nu=\sinh \mathrm{T}_{D} / \mathrm{T}_{D}$ and $\mathrm{T}_{D}=\sqrt{\mathrm{LD}^{2}+\mathrm{LD}^{\prime 2}+\mathrm{CD}^{2}}$.

### 3.2.2. The effect of the interfaces

The infinitesimal matrix method described in 3.2 does not take into account the effect of the interfaces. When studying light transmission through a certain homogeneous material the most common situation is having two well defined interfaces: one delimiting the incident medium (usually air) to the material and the other one between the material an the exiting medium (also usually air). If multiple reflections are excluded, the transmission Mueller matrix of the complete system is then given by:

$$
\begin{equation*}
\mathbf{M}=\mathbf{M}_{I 1} \mathbf{M}_{S} \mathbf{M}_{I 0} \tag{3.26}
\end{equation*}
$$

[^14]where $\mathbf{M}_{I 0}$ and $\mathbf{M}_{I 1}$ are, respectively, the Mueller matrices corresponding to the incident and exiting interfaces. $\mathbf{M}_{S}$ is the Mueller-Jones matrix of the "bulk" material given in Eq. (3.21), and constructed from the thickness-dependent parameters given in Table 1.1.

At normal-incidence, $\mathbf{M}_{I 0}$ and $\mathbf{M}_{I 1}$ can only have a certain contribution if the measured sample is anisotropic, otherwise the matrices $\mathbf{M}_{I 0}$ and $\mathbf{M}_{I 1}$ are the identity matrix. If the sample under study is nonabsorbent (i.e. if LD, LD' and CD in $\mathbf{M}_{S}$ vanish) then $\mathbf{M}_{I 0}$ and $\mathbf{M}_{I 1}$ describe the difference polarization transmission at the interface, i.e. a diattenuation, and, in this case, the interface matrices do not include any retardation. In the case where $\mathbf{M}_{S}$ contains any anisotropic absorption then there appears a phase shift at the interface that translates in Mueller matrices $\mathbf{M}_{I 0}$ and $\mathbf{M}_{I 1}$ including also retardation terms.

The interface diattenuation for two orthogonal polarization states, e.g. $x$ and $y$, as it appears in the Mueller matrix, can be calculated as:

$$
\begin{equation*}
D_{y x}^{I}=\frac{t_{y} t_{y}^{*}-t_{x} t_{x}^{*}}{t_{y} t_{y}^{*}+t_{x} t_{x}^{*}}, \tag{3.27}
\end{equation*}
$$

where the superscript $I$ refers to the interface and $t_{y}$ and $t_{x}$ are the complex transmission coefficients, that, at normal incidence, are:

$$
\begin{equation*}
t_{k}=\frac{2 n_{0}}{n_{0}+n_{k}-i k_{k}}, \quad k=x, y \tag{3.28}
\end{equation*}
$$

where $n_{0}$ is the refractive index in the incident medium and $n_{k}-i k_{k}$ is the complex refractive index of the sample for light polarized along the $k$ direction.

A subtlety that needs to be noted is that, for an interface diattenuation, the real-valued terms $t_{k} t_{k}^{*}$ used in Eq.(3.27) are not directly the intensity transmission coefficients $T_{x}$ and $T_{y}$ for light polarized along the $x$ or $y$, used in other definitions of the global diattenuation [82]. These transmission coefficients are

$$
\begin{equation*}
T_{k}=t_{k} t_{k}^{*} \frac{n_{k}}{n_{0}}=1-R_{k} . \quad(\text { normal incidence }) \tag{3.29}
\end{equation*}
$$

where the factor $n_{k} / n_{0}$ is due to the change of medium. The reason why this transmittance coefficient, $T_{k}$, should not be used here is that the diattenuation occurring at one interface depends only on the amplitude of the transmitted fields as a function of the incident polarization, and the index of refraction is not included in the definitions of the Jones vectors or in the Stokes parameters. In case we were looking at the diattenuation of the reflected light, the reflection coefficients $R_{k}$ could be used for the calculation of the diattenuation [83].

The retardation for the transmitted beam upon the interface for two orthogonal polarization states, e.g. $x$ and $y$, is

$$
\begin{equation*}
\Delta_{y x}=\arg \left(t_{y}\right)-\arg \left(t_{x}\right), \tag{3.30}
\end{equation*}
$$

which, at normal incidence, is

$$
\begin{equation*}
\Delta_{y x}=\operatorname{atan}\left[\frac{k_{y}\left(n_{0}+n_{x}\right)-k_{x}\left(n_{0}+n_{y}\right)}{\left(n_{0}+n_{x}\right)\left(n_{0}+n_{y}\right)+k_{x} k_{y}}\right], \tag{3.31}
\end{equation*}
$$

demonstrating that for $k_{x}=k_{y}=0$ no retardation appears in the interface.
At normal incidence the effect of the interfaces will be, in general, very small and only will be noticeable for samples with large anisotropy. For example, in a crystal with huge birefringence such as calcite, the transmission Mueller matrices at the interfaces are:

$$
\begin{gather*}
\mathbf{M}_{I 0}=\left[\begin{array}{cccc}
1 & 0.0668 & 0 & 0 \\
0.0668 & 1 & 0 & 0 \\
0 & 0 & 0.9978 & 0 \\
0 & 0 & 0 & 0.9978
\end{array}\right], \\
\mathbf{M}_{I 1}=\left[\begin{array}{cccc}
1 & -0.0426 & 0 & 0 \\
-0.0426 & 1 & 0 & 0 \\
0 & 0 & 0.9991 & 0 \\
0 & 0 & 0 & 0.9991
\end{array}\right], \tag{3.32}
\end{gather*}
$$

where we have assumed that the optic axis is parallel to the y axis of the laboratory frame ( $n_{y}=n_{e}=1.486, n_{x}=n_{o}=1.658$ ). Only when studying samples with large anisotropy does the effect of the interfaces need to be considered in detail. Otherwise the Mueller interface matrices may be taken as the identity matrix without introducing significant errors. In the normal-incidence transmission experiments performed in this thesis the effect of the interfaces has not been considered.

### 3.2.3. The controversy about Jones birefringence ( $\mathrm{LB}^{\prime}$ ) and Jones dichroism ( $\mathrm{LD}^{\prime}$ )

Certain of the contents of this section are probably redundant considering the introduction to the optical effects we made in section 1.5.2. However we consider that they are important enough to be studied here in more detail as they have led to confusions in a considerable number of publications.

In 1983 Graham and Raab published a paper [84] stating that in Jones 1948 work [75] a new kind of linear birefringence, together with its corresponding dichroism, had been postulated, and proposed the names Jones birefringence and Jones dichroism for such effects. They argued that the linear birefringence and the linear dichroism parallel with the bisectors of the coordinate axes $x, y$, that in this work we have respectively named as $\mathrm{LB}^{\prime}$ and $\mathrm{LD}^{\prime}$ (see Eq. (1.41) and Eq. (1.47) ), were new types of optical effects that were unknown up to the date. In their work they predicted that Jones birefringence can occur naturally in
certain uniaxial and biaxial non-magnetic crystals, where it should accompany the "familiar" linear birefringence. They also predicted the presence of Jones birefringence in liquids subjected to parallel electric and magnetic fields. Later, in Ref. [85] they found, by using a multipole approach to the electromagnetic effects, that Jones birefringence would be typically four orders of magnitude smaller than the usual linear birefringence.

To our knowledge no experimental observations of the Jones birefringence in crystals have been reported. According to [86], Jones birefringence occurring naturally in crystals is probably too weak to be measurable. However, the first experimental observation of Jones birefringence in systems with parallel electric and magnetic fields was published in 2000 [87], and, in 2003, the first experimental observation of Jones dichroism was reported [88]. Since then, the interest on the subject of Jones birefringence and Jones dichroism seems to increase [89-94] and recent books, most notably the well-known book about optical activity by Barron [95], review the subject. Much like Ref. [84], most of these publications state that these supposed new effects were deduced by Jones, and even, in Ref. [87], it is said that their experimental observation constitutes the final validation of the Jones formalism in polarization optics.

We argue that in his 1948 paper Jones did not introduce neither deduce any intriguing new optical phenomenon. We think that linear birefringence and linear dichroism parallels with the bisectors of the coordinate axes that Jones used are not new optical effects of difficult experimental observation, and that the controversy about these effects stems from a misinterpretation of the coordinated system used in the original Jones publication, misinterpretation that probably started with Ref. [84].

Jones calculus can be developed for an arbitrary basis, although the most usual choice is a laboratory Cartesian coordinate system in which light propagates along the positive $z$ axis as the one we have been using in this chapter. Rather than a new type of birefringence, $\mathrm{LB}^{\prime}$ is only a measure of the part of the linear birefringence which is parallel to the bisector of the coordinate axes and it does not supposes any new finding. We suspect that the confusion must have arisen with the incorrect assumption that Jones used a coordinate system based on the optical axis of the optical element under study. As we said when we introduced the optical effects, it is important to stress that, in general, neither LB nor $\mathrm{LB}^{\prime}$, as defined by Jones, are equivalent to the usual definition of linear birefringence given in crystal optics:

$$
\begin{equation*}
\delta=\frac{2 \pi}{\lambda}\left(n_{e}-n_{o}\right) l, \tag{3.33}
\end{equation*}
$$

because this definition is given for the natural basis of the birefringent element, i. e. a coordinate system based on the crystallographic (ordinary and extraordinary) axes of the optical element. If the optic axis lies in the xy plane (it may be not always the case) the relation between $\delta$ and LB and $\mathrm{LB}^{\prime}$ clearly shows that LB
and $\mathrm{LB}^{\prime}$ can be understood as measurements of the "projected" of birefringence

$$
\begin{equation*}
\mathrm{LB}=\frac{\delta}{\cos }(2 \theta), \quad \mathrm{LB}^{\prime}=\frac{\delta}{\sin }(2 \theta) \tag{3.34}
\end{equation*}
$$

where $\theta$ is the angle shown in Fig. 3.3. For an arbitrary orientation of the optic axis, the correspondence between $\delta$ and LB and $\mathrm{LB}^{\prime}$ is more complex and involves the complete set of Euler angles.


Figure 3.3.: For an optical element with the optic axis lying in the xy plane, the laboratory coordinates $x$ and $y$ (reference basis) are related to the crystallographic coordinates (natural basis) by a simple rotation.

To our knowledge there are no further and solid arguments proving the existence of a supposed new type of birefringence or dichroism independent from the "usual" one. Theoretical works on this subject are not aimed to demonstrate the Jones birefringence and Jones dichroism as new phenomena independent from the standard linear birefringence; instead, they assume that they exist by citing Jones work, and focus their attention on identifying systems in which these effects may be possible.

According to our thesis [96], experimental observations of that part of the linear birefringence or the linear dichroism parallel to the bisector of the laboratory coordinate axes can be easily done with polarimetric techniques, as it is shown, for example, in [97], and it does not involve any special difficulty. Probably, most of the reported experimental results on Jones birefringence or Jones dichroism could be interpreted as measurements of "projected" birefringence or dichroism,
and, therefore, their physical meaning would be given by their correspondence, in terms of Euler angles, to the birefringence or dichroism of the natural basis. We also use these lines to advise against the use of the terms of "Jones birefringence" and "Jones dichroism" as we consider that they were proposed following a misinterpretation of Jones' work.

### 3.2.4. Twisted crystal

We have used the lamellar representation to study in detail the important special case in which the $\mathbf{N}$-matrices are independent of $z$. There is a another special dependence of $\mathbf{N}$ upon $z$ that permits a simple solution: the case of a uniformly twisted crystal in which light is incident along the helical axis. This problem was already solved by Jones [75] and it is specially suitable for the the study of the cholesteric phase of liquid crystals. We will dedicate our attention to study this problem, since it constitutes a simple example of how a material constituted by achiral elements can give rise to optical activity.

Let $\mathbf{N}_{0}$ be the $\mathbf{N}$-matrix of the untwisted crystal. For simplicity we consider that each thin slab does not has own CB and CD and the untwisted $\mathbf{N}_{0}$ is:

$$
\mathbf{N}_{0}=\frac{1}{2 z}\left(\begin{array}{cc}
-\mathrm{LD}-i \mathrm{LB}-2(i \eta+\kappa) & 0  \tag{3.35}\\
0 & \mathrm{LD}+i \mathrm{LB}-2(i \eta+\kappa)
\end{array}\right)
$$

Then, according to Eq. (1.21) the $\mathbf{N}$-matrix corresponding to a twisted crystal at a distance $z$ is given by

$$
\begin{equation*}
\mathbf{N}=\mathbf{R}(-a z) \mathbf{N}_{0} \mathbf{R}(a z) \tag{3.36}
\end{equation*}
$$

where $a$ is the angular twist per unit of thickness (e.g. radians per meter). Fig. 3.4 shows that the progressive and uniform rotation of the infinitesimal slices give yields an helical structure. The $z$ axis of this structure coincides with the direction of light propagation. The pitch $P$ of the helix (the width of one complete helix turn) is related to $a$ by

$$
\begin{equation*}
a=\frac{2 \pi}{P} . \tag{3.37}
\end{equation*}
$$

Eq. (3.10) can now be written as

$$
\begin{equation*}
\frac{d \mathbf{J}}{d z}=\mathbf{R}(-a z) N_{0} \mathbf{R}(a z) \mathbf{J}, \tag{3.38}
\end{equation*}
$$

that after introducing the substitution

$$
\begin{equation*}
\mathbf{J}^{\prime} \equiv \mathbf{R}(a z) \mathbf{J} \tag{3.39}
\end{equation*}
$$

becomes

$$
\begin{equation*}
\frac{d \mathbf{J}^{\prime}}{d z}=\left[\mathbf{N}_{0}-\mathbf{R}(a z) \frac{d \mathbf{R}(-a z)}{d z}\right] \mathbf{J}^{\prime}=\mathbf{N}^{\prime} \mathbf{J} \tag{3.40}
\end{equation*}
$$



Figure 3.4.: Scheme of the lamellar representation for a homogeneously twisted crystal. Red marks indicate the direction of the axis of birefringence for each one of the infinitesimal slices. The pitch $P$ indicates the distance at which the axis has made a complete turn.
where $\mathbf{N}^{\prime}=\mathbf{N}_{0}-\mathbf{R}(a z) \frac{d \mathbf{R}(-a z)}{d z}$. After deriving the rotator matrix we find that $\mathbf{N}^{\prime}$ can also be written in the following form:

$$
\begin{equation*}
\mathbf{N}^{\prime}=\mathbf{N}_{0}+a \mathbf{R}(\pi / 2) \tag{3.41}
\end{equation*}
$$

The solution of Eq. (3.40) may be written

$$
\begin{equation*}
\mathbf{J}^{\prime}=\exp \left[\mathbf{N}^{\prime} z\right] \tag{3.42}
\end{equation*}
$$

and by the use of (3.39) this equation becomes

$$
\begin{equation*}
\mathbf{J}=\mathbf{R}(-a z) \exp \left[\mathbf{N}^{\prime} z\right]=\mathbf{R}(-a z) \exp \left(\mathbf{N}_{0}+a \mathbf{R}(\pi / 2)\right) z \tag{3.43}
\end{equation*}
$$

The Jones matrix $\mathbf{J}$ of the twisted crystal is equal is equal to $\mathbf{R}(-a z)$ multiplied by the Jones matrix corresponding to an homogeneous medium with a N-matrix equal to $\mathbf{N}_{0}+a \mathbf{R}(\pi / 2)$.

Eq (3.43) can be easily written in terms of a product of Mueller-Jones matrices. The rotation matrix $\mathbf{R}$ for the Mueller formalism has been given in Eq. (1.27) and the Mueller-Jones matrix corresponding to an homogeneous medium with a N -matrix equal to $\mathbf{N}_{0}+a \mathbf{R}(\pi / 2)$ is easily derivable from the general MuellerJones matrix of a homogeneous medium given in Eq. (3.21). To make things even easier we will consider that the $\mathbf{N}_{0}$ matrix does not contain absorption effects (i.e. LD and $\kappa$ terms of (3.35) vanish). In this case we can write the Mueller-Jones

Chapter 3. Light propagation in anisotropic optical active media
matrix for a twisted crystal as the following product of matrices:
$\mathbf{M}=\left[\begin{array}{cccc}1 & 0 & 0 & 0 \\ 0 & \cos 2 a z & -\sin 2 a z & 0 \\ 0 & \sin 2 a z & \cos 2 a z & 0 \\ 0 & 0 & 0 & 1\end{array}\right]\left[\begin{array}{cccc}1 & 0 & 0 & 0 \\ 0 & \cos \mathrm{~T}_{L}+\mathrm{LB}^{2} \alpha & 2 a z \beta & -\mathrm{LB} 2 a z \alpha \\ 0 & -2 a z \beta & \cos \mathrm{~T}_{L} & -\mathrm{LB} \beta \\ 0 & -2 a z \mathrm{LB} \alpha & \mathrm{LB} \beta & \cos \mathrm{T}_{L}+(2 a z)^{2} \alpha\end{array}\right]$
with $\alpha=\left(1-\cos \mathrm{T}_{L}\right) / \mathrm{T}_{L}^{2}, \beta=\sin \mathrm{T}_{L} / \mathrm{T}_{L}$ and $\mathrm{T}_{L}=\sqrt{\mathrm{LB}^{2}+(2 a z)^{2}}$.
Note that that if we impose that isotropy of the medium (i. e. we take $\mathrm{LB}=0$ ) the Mueller matrix on Eq. (3.44) simplifies to an identity matrix (the second matrix factor becomes the inverse of the first factor). This is a expected result, since an the polarization of light that passes though an isotropic medium, no matter how it is oriented, is not altered.

If the pitch is not too large the quantity $2 a z$ will be much greater than LB $(2 a z \gg \mathrm{LB})$. When this happens we can approximate $\mathrm{T}_{L} \simeq 2 a z$ and the Mueller matrix becomes:

$$
\mathbf{M}=\left[\begin{array}{cccc}
1 & 0 & 0 & 0  \tag{3.45}\\
0 & 1+\cos 2 a z \frac{\mathrm{LB}^{2}}{(2 a z)^{2}}(1-\cos 2 a z) & 0 & \frac{\mathrm{LB}}{2 a z}(1-\cos 2 a z) \\
0 & \mathrm{LB}^{2} \sin 2 a z \frac{1-\cos 2 a z}{(2 a z)^{2}} & 1 & -\frac{\mathrm{LB}}{2 a z} \sin 2 a z \\
0 & -\frac{\mathrm{LB}}{2 a z}(1-\cos 2 a z) & \frac{\mathrm{LB}}{2 a z} \sin 2 a z & 1
\end{array}\right] .
$$

The N-matrix for each one of the infinitesimal layers considered in this model did not include optical effects related to optical activity (CB and CD). However the Jones or the Mueller-Jones matrix of the complete twisted crystal will appear to be that of a homogeneous medium exhibiting optical activity. For example, given a general retarder (Eq. (3.24)), the CB is proportional to the difference of Mueller matrix elements $m_{12}$ and $m_{21}$ and LB is proportional to $m_{23}-m_{32}$ :

$$
\begin{align*}
& \mathrm{CB}=\frac{\mathrm{T}_{R}}{\sin \mathrm{~T}_{R}}\left(m_{12}-m_{21}\right) / 2  \tag{3.46a}\\
& \mathrm{LB}=\frac{\mathrm{T}_{R}}{\sin \mathrm{~T}_{R}}\left(m_{32}-m_{23}\right) / 2 \tag{3.46b}
\end{align*}
$$

where $\mathrm{T}_{R}$ is the total retardance of the optical element. We can write:

$$
\begin{equation*}
\mathrm{CB}=\mathrm{LB} \frac{m_{32}-m_{23}}{m_{12}-m_{21}}, \tag{3.47}
\end{equation*}
$$

using this equation with the elements of the matrix shown in Eq. (3.45) we can find the approximate circular birefringence induced by the twist of the crystal:

$$
\begin{equation*}
\mathrm{CB}=-\frac{\mathrm{LB}^{2}}{4 a z}(1-\cos 2 a z) \tag{3.48}
\end{equation*}
$$

and using the definitions $\mathrm{LB}=2 \pi \delta n z / \lambda$ [Eq. (1.40)] and $a=2 \pi / P$ [Eq. (3.37)], the circular birefringence is

$$
\begin{equation*}
\mathrm{CB}=-\frac{\pi(\delta n)^{2} P z}{2 \lambda^{2}}(1-\cos 2 a z) . \tag{3.49}
\end{equation*}
$$

The negative sign indicates that the sense of the rotation is opposite to that of the helical twist of the structure. This solution is quite interesting because it shows that even a simple phenomenological models are able to reproduce the typical $\sim 1 / \lambda^{2}$ dependence for optical rotation which usually observed experimentally. Note also that in this case the pitch, $P$, plays the role of the parameter $d$ (a typical molecular length) we mentioned at the beginning of this chapter. This solution describes, for example, the propagation of light along the helical axis of a cholesteric liquid crystal but only for wavelengths much smaller than the pitch. Otherwise we should use a more complex model to consider the reflection and interference effects that now we have neglected. The same calculus can be performed by directly multiplying uniformly twisted Jones matrices and without introducing the $\mathbf{N}$ matrix (it is shown in Ref. [98] ). It yields the same results but involves more cumbersome algebra.

As a final remark is worth to recall here that from the point of view of the Jones or the Mueller-Jones matrices an optical rotation of, let's say, $20^{\circ}$ is indistinguishable from an optical rotation of $380^{\circ}$ (also from $740^{\circ}, 1100^{\circ}$, etc). This is the reason why Eq. (3.49) contains the term $(1-\cos 2 a z)$. There is always a $2 \pi$ indetermination associated to the Jones/Mueller matrix of a retarder and this facts needs to be considered if we deal with samples that are expected to exhibit a retardance greater than $2 \pi$ radians. This indetermination also appears when measuring linear birefringence although it is somewhat less evident.

## Chapter 4.

## Inversion and decomposition of Mueller matrices

In this chapter we will focus our attention in the obtainment of the relevant polarization-dependent optical effects from a given experimental Mueller matrix. The first method we will present is the analytic inversion of the more general Jones or Mueller-Jones matrices obtained in the previous chapter. This inversion is restricted to the case of light propagation through an homogeneous medium with arbitrary absorptive and refractive anisotropies, but it is one of the most common situations one can find when studying complex materials with polarimetric techniques.

However, the problem of the physical interpretation of a measured Mueller matrix can be also studied with an algebraic approach based on decompositions of the Mueller matrix. Constructing a physical model for a Jones or a Mueller matrix is only feasible in a limited number of optical systems (some of them have been shown in the previous chapter). Moreover, even if a certain model for a system with optical anisotropy is available, it will probably give a description based on the Jones matrix, and, because of the phenomenon of depolarization, it may do not directly correspond to experimental Mueller matrix, i.e. it can be not possible to easily extract a Jones matrix from an experimental Mueller matrix. If used with care, decompositions methods can be used to to extract useful information from a Mueller matrix without the need of a an physical model.

### 4.1. Analytic inversion of the Mueller-Jones polarization matrices for homogeneous media

The purpose of this section is present an analytic solution to the inversion problem of a Jones or a Mueller-Jones matrix corresponding to a homogeneous media with combined arbitrary absorptive and refractive anisotropies. For many simpler situations, for example media presenting only linear birefringent anisotropy or media that exhibits only circular dichroism, the inversion problem is almost trivial [see Eq. (3.24) and Eq. (3.25)]. However, when combined amplitude and phase anisotropy are present [Eq. (3.21)] the situation is more complex because the optical effects are coupled together and, to our knowledge, no rigorous equa-

Chapter 4. Inversion and decomposition of Mueller matrices
tions had been published to make possible an analytic inversion in this more general case.

For convenience we will express the elements of the Jones matrix in polar form:

$$
\mathbf{J}=\left(\begin{array}{ll}
j_{00} & j_{01}  \tag{4.1}\\
j_{10} & j_{11}
\end{array}\right)=e^{i \theta_{00}}\left(\begin{array}{cc}
r_{00} & r_{01} e^{i\left(\theta_{01}-\theta_{00}\right)} \\
r_{10} e^{i\left(\theta_{10}-\theta_{00}\right)} & r_{11} e^{i\left(\theta_{11}-\theta_{00}\right)}
\end{array}\right) .
$$

The Jones matrix elements, as given in Eq. (4.1), corresponding to a given Mueller-Jones matrix can be calculated using the following equations [7]:

$$
\begin{gather*}
r_{00}=\left[\left(m_{00}+m_{01}+m_{10}+m_{11}\right) / 2\right]^{1 / 2},  \tag{4.2}\\
r_{01}=\left[\left(m_{00}-m_{01}+m_{10}-m_{11}\right) / 2\right]^{1 / 2},  \tag{4.3}\\
r_{10}=\left[\left(m_{00}+m_{01}-m_{10}-m_{11}\right) / 2\right]^{1 / 2},  \tag{4.4}\\
r_{11}=\left[\left(m_{00}-m_{01}-m_{10}+m_{11}\right) / 2\right]^{1 / 2},  \tag{4.5}\\
e^{i\left(\theta_{01}-\theta_{00}\right)}=\frac{m_{02}+m_{12}-i\left(m_{03}+m_{13}\right)}{\left[\left(m_{00}+m_{10}\right)^{2}-\left(m_{01}+m_{11}\right)^{2}\right]^{1 / 2}},  \tag{4.6}\\
e^{i\left(\theta_{10}-\theta_{00}\right)}=\frac{m_{20}+m_{21}+i\left(m_{30}+m_{31}\right)}{\left[\left(m_{00}+m_{01}\right)^{2}-\left(m_{10}+m_{11}\right)^{2}\right]^{1 / 2}},  \tag{4.7}\\
e^{i\left(\theta_{11}-\theta_{00}\right)}=\frac{m_{22}+m_{33}+i\left(m_{32}-m_{23}\right)}{\left[\left(m_{00}+m_{11}\right)^{2}-\left(m_{10}+m_{01}\right)^{2}\right]^{1 / 2}} . \tag{4.8}
\end{gather*}
$$

For any Mueller-Jones matrix the Eqs. (4.2)-(4.8) can be used to calculate $r_{00}$, $r_{01}, r_{10}, r_{11}, e^{i\left(\theta_{01}-\theta_{00}\right)}, e^{i\left(\theta_{11}-\theta_{00}\right)}$ and $e^{i\left(\theta_{11}-\theta_{00}\right)}$. This factors can be identified with the parameterized general Jones matrix in Eq. (3.19):

$$
\left(\begin{array}{cc}
\cos \frac{\mathrm{T}}{2}-\frac{i \mathrm{~L}}{\mathrm{~T}} \sin \frac{\mathrm{~T}}{2} & \frac{\left(\mathrm{C}-i \mathrm{~L}^{\prime}\right)}{\mathrm{T}} \sin \frac{\mathrm{~T}}{2}  \tag{4.9}\\
-\frac{\left(\mathrm{C}+i \mathrm{~L}^{\prime}\right)}{\mathrm{T}} \sin \frac{\mathrm{~T}}{2} & \cos \frac{\mathrm{~T}}{2}+\frac{i \mathrm{~L}}{\mathrm{~T}} \sin \frac{\mathrm{~T}}{2}
\end{array}\right)=K\left(\begin{array}{cc}
r_{00} & r_{01} e^{i\left(\theta_{01}-\theta_{00}\right)} \\
r_{10} e^{i\left(\theta_{10}-\theta_{00}\right)} & r_{11} e^{i\left(\theta_{11}-\theta_{00}\right)}
\end{array}\right),
$$

where K is a complex constant that can be determined combining the matrix elements of Eq. (4.9) and using the identity $\cos ^{2} \frac{T}{2}+\sin ^{2} \frac{T}{2}=1$ :

$$
\begin{equation*}
K=\left[r_{00} r_{11} e^{i\left(\theta_{11}-\theta_{00}\right)}-r_{01} r_{10} e^{i\left(\theta_{01}-\theta_{00}\right)} e^{i\left(\theta_{10}-\theta_{00}\right)}\right]^{-1 / 2} \tag{4.10}
\end{equation*}
$$

Once K is known, the determination of $\mathrm{LB}, \mathrm{LB}$, $\mathrm{CB}, \mathrm{LD}, \mathrm{LD}$ ' and CD from Eq. (4.9) becomes straightforward [99]:

$$
\begin{gather*}
\mathrm{LB}=\operatorname{Re}\left[i \Omega\left(r_{00}-r_{11} e^{i\left(\theta_{11}-\theta_{00}\right)}\right)\right]  \tag{4.11}\\
\mathrm{LB}^{\prime}=\operatorname{Re}\left[i \Omega\left(r_{01} e^{i\left(\theta_{01}-\theta_{00}\right)}+r_{10} e^{i\left(\theta_{10}-\theta_{00}\right)}\right)\right] \tag{4.12}
\end{gather*}
$$

$$
\begin{gather*}
\mathrm{CB}=\operatorname{Re}\left[\Omega\left(r_{01} e^{i\left(\theta_{01}-\theta_{00}\right)}-r_{10} e^{i\left(\theta_{10}-\theta_{00}\right)}\right)\right],  \tag{4.13}\\
\mathrm{LD}=-\operatorname{Im}\left[i \Omega\left(r_{00}-r_{11} e^{i\left(\theta_{11}-\theta_{00}\right)}\right)\right],  \tag{4.14}\\
\mathrm{LD}^{\prime}=-\operatorname{Im}\left[i \Omega\left(r_{01} e^{i\left(\theta_{01}-\theta_{00}\right)}+r_{10} e^{i\left(\theta_{10}-\theta_{00}\right)}\right)\right],  \tag{4.15}\\
\mathrm{CD}=-\operatorname{Im}\left[\Omega\left(r_{01} e^{i\left(\theta_{01}-\theta_{00}\right)}-r_{10} e^{i\left(\theta_{10}-\theta_{00}\right)}\right)\right], \tag{4.16}
\end{gather*}
$$

where $\Omega=\mathrm{T} K /[2 \sin (\mathrm{~T} / 2)], \mathrm{T}=2 \cos ^{-1}\left[K\left(r_{00}+r_{11} e^{i\left(\theta_{11}-\theta_{00}\right)}\right) / 2\right]$ and the symbols Re and Im respectively denote the real and imaginary parts.

### 4.2. Inversion of a experimental Mueller matrix

Nevertheless a polarimetric experiment can never provide an exact Mueller-Jones matrix because of sample depolarization effects and measurement uncertainties. In order to apply the analytic inversion method described in the Eqs. (4.11)(4.16) to analyze an experimental Mueller matrix we use the approach based on the sum decomposition introduced by Cloude [100] to obtain a nondepolarizing Mueller (i.e. a Mueller-Jones) matrix estimation for a depolarizing experimental matrix. This sum decomposition is based in the coherency matrix associated to a Mueller matrix. The coherency matrix is a $4 \times 4$ positive semidefinite Hermitian matrix that is used for eigen-analysis (i.e. the calculation of the corresponding eigenvalues and eigenvectors) because it yields information into the different polarimetric contributions that a depolarizing Mueller matrix can contain. The elements $h_{i j}$ of the coherency matrix $\mathbf{H}$ can be derived from the corresponding Mueller matrix as follows [101]:

$$
\begin{array}{ll}
h_{00}=\left(m_{00}+m_{11}+m_{22}+m_{33}\right) / 4, & h_{01}=\left(m_{01}+m_{10}-i m_{23}+i m_{32}\right) / 4, \\
h_{02}=\left(m_{02}+m_{20}+i m_{13}-i m_{31}\right) / 4, & h_{03}=\left(m_{03}-i m_{12}+i m_{21}+m_{30}\right) / 4, \\
h_{10}=\left(m_{01}+m_{10}+i m_{23}-i m_{32}\right) / 4, & h_{11}=\left(m_{00}+m_{11}-m_{22}-m_{33}\right) / 4, \\
h_{12}=\left(i m_{03}+m_{12}+m_{21}-i m_{30}\right) / 4, & h_{13}=\left(-i m_{02}+i m_{20}+m_{13}+m_{31}\right) / 4, \\
h_{20}=\left(m_{02}+m_{20}-i m_{13}+i m_{31}\right) / 4, & h_{21}=\left(-i m_{03}+m_{12}+m_{21}+i m_{30}\right) / 4, \\
h_{22}=\left(m_{00}-m_{11}+m_{22}-m_{33}\right) / 4, & h_{23}=\left(i m_{01}-i m_{10}+m_{23}+m_{32}\right) / 4, \\
h_{30}=\left(m_{03}+i m_{12}-i m_{21}+m_{30}\right) / 4, & h_{31}=\left(i m_{02}-i m_{20}+m_{13}+m_{31}\right) / 4, \\
h_{32}=\left(-i m_{01}+i m_{10}+m_{23}+m_{32}\right) / 4, & h_{33}=\left(m_{00}-m_{11}-m_{22}+m_{33}\right) / 4 .
\end{array}
$$

This matrix is positive semidefinite Hermitian and, hence, has always four real non-negative eigenvalues $\lambda_{i}$. Given the eigenvalues $\lambda_{i}$ any physically realizable Mueller matrix $\mathbf{M}$ can be written as ${ }^{1}$ :

$$
\begin{equation*}
\mathbf{M}=\lambda_{0} \mathbf{M}_{J}^{0}+\lambda_{1} \mathbf{M}_{J}^{1}+\lambda_{2} \mathbf{M}_{J}^{2}+\lambda_{3} \mathbf{M}_{J}^{3} \tag{4.17}
\end{equation*}
$$

[^15]where $\mathbf{M}_{J}^{k}$ are Mueller-Jones matrices that are derived from the following Jones matrices by using Eq. (1.29):
\[

$$
\begin{align*}
j_{00}^{(k)} & =\Psi_{0}^{(k)}+\Psi_{1}^{(k)}, & j_{01}^{(k)} & =\Psi_{2}^{(k)}-i \Psi_{3}^{(k)}  \tag{4.18}\\
j_{10}^{(k)} & =\Psi_{2}^{(k)}+i \Psi_{3}^{(k)}, & j_{11}^{(k)} & =\Psi_{0}^{(k)}-i \Psi_{2}^{(k)} \tag{4.19}
\end{align*}
$$
\]

where $\Psi^{k}=\left(\begin{array}{llll}\Psi_{0} & \Psi_{1} & \Psi_{2} & \Psi_{3}\end{array}\right)_{k}^{T}$ is the $k$ th eigenvector of the coherency matrix H.

If $\mathbf{M}$ is a Mueller-Jones matrix one has $\lambda_{0} \neq \lambda_{1}=\lambda_{2}=\lambda_{3}=0$. For a experimentally determined $\mathbf{M}$, if depolarization is not very intense, usually $\lambda_{0} \gg$ $\lambda_{1}, \lambda_{2}, \lambda_{3}$ and then the summand $\lambda_{0} \mathbf{M}_{J 0}$ can be considered as the nondepolarizing estimate for the measured $\mathbf{M}$.

In this section we have shown that the polarization effects of a homogeneous medium contained in a Mueller-Jones matrix can be easily revealed by means of simple analytic equations. In contrast to the matrix product decomposition methods that we will describe in the following section, this inversion offers exact results for Mueller-Jones matrices. In the case of experimentally determined Mueller matrices the Eqs. (4.11) to (4.16) can also be used if a nondepolarizing estimate of the experimental Mueller matrix $\left(\lambda_{0} \mathbf{M}_{J 0}\right)$ is found with anticipation. In general we can say that the reliability of the results offered by this method when applied experimental matrices will largely depend on their degree of depolarization: for homogeneous media exhibiting little depolarization this is the best option for the interpretation of Mueller matrices.

### 4.3. Decomposition of Mueller matrices

In this section we switch to a different philosophy: in the absence of a physical model describing and explaining the interaction of an electromagnetic wave with a sample, it may be still possible to phenomenologically interpret a experimental Mueller matrix usually by decomposing it into simpler components having a simpler physical interpretation. In this section we review various decompositions of non-depolarizing and depolarizing Mueller matrices.

### 4.3.1. Polar and Lu-Chipman decompositions

One approach to model homogeneous anisotropic media is based on the polar decomposition theorem, which states that any complex matrix $\mathbf{A}$ can be represented by a product

$$
\begin{equation*}
\mathbf{A}=\mathbf{U P} \quad \text { or } \quad \mathbf{A}=\mathbf{P}^{\prime} \mathbf{U}, \tag{4.20}
\end{equation*}
$$

where $\mathbf{P}$ and $\mathbf{P}^{\prime}$ are Hermitian matrices and $\mathbf{U}$ is a unitary matrix. Polar decomposition has been applied in the polarization theory to represent arbitrary
optical systems either in Jones $[102,103]$ or Mueller space [82, 104]. The Hermitian matrix appearing in the polar decomposition is associated to the amplitude anisotropy while the unitary matrix is associated to the phase anisotropy. The polar decomposition applied to a non-depolarizing Mueller matrix is:

$$
\begin{equation*}
\mathbf{M}=\mathbf{M}_{D 1} \mathbf{M}_{R}=\mathbf{M}_{R} \mathbf{M}_{D 2} \tag{4.21}
\end{equation*}
$$

where $\mathbf{M}_{D i}$ and $\mathbf{M}_{R}$ are respectively the matrices shown in Eq. (3.25) and (3.24). Because of the non-commutativity of the matrix product the two diattenuators $\mathbf{M}_{D 1}$ and $\mathbf{M}_{D 2}$ are not identical and are related through

$$
\begin{equation*}
\mathbf{M}_{D 2}=\mathbf{M}_{R}^{T} \mathbf{M}_{D 1} \mathbf{M}_{R} \quad \text { and } \quad \mathbf{M}_{D 1}=\mathbf{M}_{R} \mathbf{M}_{D 2} \mathbf{M}_{R}^{T} \tag{4.22}
\end{equation*}
$$

The algorithm to calculate the polar decomposition is presented in the following section in the more general depolarizing case.

## Depolarizing case, Lu-Chipman decomposition

The Lu-Chipman decomposition [82] is a natural generalization of the polar decomposition to the case of depolarizing Mueller matrices. In this case an arbitrary depolarizing Mueller matrix $\mathbf{M}$ is decomposed into the product of a diattenuator, a retarder and a depolarizer:

$$
\begin{equation*}
\mathbf{M}=\mathbf{M}_{\Delta} \mathbf{M}_{R} \mathbf{M}_{D} \tag{4.23}
\end{equation*}
$$

$\mathbf{M}_{\Delta}$ is the Mueller matrix of a depolarizer with polarizance ${ }^{2}$, which can be expressed as:

$$
\mathbf{M}_{\Delta}=\left[\begin{array}{cc}
1 & \overrightarrow{0}^{T}  \tag{4.24}\\
\overrightarrow{P_{\Delta}} & \mathbf{m}_{\Delta}
\end{array}\right]
$$

in which $\mathbf{m}_{\boldsymbol{\Delta}}$ is a $3 \times 3$ symmetric matrix, $\overrightarrow{0}$ denotes the three-element zero vector, and $\overrightarrow{P_{\Delta}}$ is the so-called polarizance vector of the depolarizer which is formed by Mueller matrix elements $m_{10}, m_{20}$ and $m_{30}$ of the depolarizer.

Here we describe briefly the algorithm to perform the polar decomposition on a experimental Mueller matrix. To be consistent with the published algorithms on this decomposition and because it permits a more compact development we will use here the same notation used by Lu and Chipman (that otherwise is common in polarization optics) which is based in the diattenuating, $\vec{D}=\left(D_{H}, D_{45}, D_{c}\right)$, and retardance, $\vec{R}=\left(R_{H}, R_{45}, R_{c}\right)$ vectors. There is a simple equivalence between our usual notation and that of Lu and Chipman and it is given in Table 4.1.

[^16]Chapter 4. Inversion and decomposition of Mueller matrices

Table 4.1.: Relation to the notation of Lu-Chipman

| Retardances | Absorbances |
| :--- | :--- |
| $\mathrm{LB}=-R_{H}$ | $\mathrm{LD}=-\left(\mathrm{T}_{D} / D\right) D_{H}$ |
| $\mathrm{LB}^{\prime}=-R_{45}$ | $\mathrm{LD}^{\prime}=-\left(\mathrm{T}_{D} / D\right) D_{45}$ |
| $\mathrm{CB}=R_{c}$ | $\mathrm{CD}=\left(\mathrm{T}_{D} / D\right) D_{c}$ |
| $\overrightarrow{\mathrm{~T}}_{R}=\left(\mathrm{LB}, \mathrm{LB}^{\prime},-\mathrm{CB}\right)=-\vec{R}$ | $\overrightarrow{\mathrm{~T}}_{D}=\left(\mathrm{LD}, \mathrm{LD}^{\prime},-\mathrm{CD}\right)=-\left(\mathrm{T}_{D} / D\right) \vec{D}$ |
| $\mathrm{~T}_{R}=R$ | $\mathrm{~T}_{D}=\operatorname{arctanh} D$ |

Any experimental Mueller normalized matrix can be identified with the following matrix:

$$
\mathbf{M}=\left[\begin{array}{cc}
1 & \vec{D}^{T}  \tag{4.25}\\
\vec{P} & \mathbf{m}
\end{array}\right]
$$

where the $m_{01}, m_{02}$ and $m_{03}$ form the diattenuation vector, $\vec{D}, \vec{P}$ is the so-called polarizance vector which is formed by $m_{10}, m_{20}$ and $m_{30}$ and $\mathbf{m}$ is $3 \times 3$ matrix obtained by striking out the first row and the first column of $\mathbf{M}$. Using the relations given in Table 4.1 we use the diattenuation vector to find the values of $\mathrm{LD}, \mathrm{LD}^{\prime}$ and CD and then to construct $\mathbf{M}_{D}$ according to Eq. (3.25). If the matrix $\mathbf{M}_{D}$ is not singular (i.e. it is invertible) we can define a new matrix $\mathbf{M}^{\prime}$ :

$$
\begin{equation*}
\mathbf{M}^{\prime} \equiv \mathbf{M M}_{D}^{-1}=\mathbf{M}_{\Delta} \mathbf{M}_{R} \tag{4.26}
\end{equation*}
$$

$\mathbf{M}^{\prime}$ has no diattenuation and contains both retardance and depolarization. $\mathbf{M}^{\prime}$ is of the form:

$$
\mathbf{M}^{\prime}=\left[\begin{array}{cc}
1 & \overrightarrow{0}^{T}  \tag{4.27}\\
\vec{P}_{\Delta} & \mathbf{m}^{\prime}
\end{array}\right]
$$

in which $\vec{P}_{\Delta}$ is the polarizance vector of the depolarizer and $\mathbf{m}^{\prime}$ is a $3 \times 3$ submatrix of $\mathbf{M}^{\prime} . \vec{P}_{\Delta}$ is:

$$
\begin{equation*}
\vec{P}_{\Delta}=\frac{\vec{P}-\mathbf{m} \vec{D}}{1-D^{2}} \tag{4.28}
\end{equation*}
$$

The submatrix $\mathbf{m}_{\Delta}$ appearing in Eq. (4.24) can be now obtained by

$$
\begin{align*}
\mathbf{m}_{\Delta}= & \pm\left[\mathbf{m}^{\prime}\left(\mathbf{m}^{\prime}\right)^{T}+\left(\sqrt{\lambda_{0} \lambda_{1}}+\sqrt{\lambda_{1} \lambda_{2}}+\sqrt{\lambda_{2} \lambda_{0}}\right) \mathbf{I}\right]^{-1}  \tag{4.29}\\
& \times\left[\left(\sqrt{\lambda_{0}}+\sqrt{\lambda_{1}}+\sqrt{\lambda_{2}}\right) \mathbf{m}^{\prime}\left(\mathbf{m}^{\prime}\right)^{T}+\sqrt{\lambda_{0} \lambda_{1} \lambda_{2}} \mathbf{I}\right] . \tag{4.30}
\end{align*}
$$

where $\lambda_{0}, \lambda_{1}, \lambda_{2}$ are the eigenvalues of $\mathbf{m}^{\prime}\left(\mathbf{m}^{\prime}\right)^{T}$. The minus sign is applied if the determinant of $\mathbf{m}^{\prime}$ is negative, otherwise, the plus sign is applied. Thus, $\mathbf{M}_{\Delta}$
can be determined through Eqs. (4.28) and (4.29). Then $\mathbf{M}_{R}$ is obtained by

$$
\begin{equation*}
\mathbf{M}_{R}=\mathbf{M}_{\Delta}^{-1} \mathbf{M}^{\prime} \tag{4.31}
\end{equation*}
$$

The Lu-Chipman estimation for $\mathrm{LB}, \mathrm{LD}, \mathrm{LB}^{\prime}, \mathrm{LD}^{\prime}, \mathrm{CD}$ and CB can be obtained using the following relations:

$$
\begin{array}{r}
\mathrm{LD}=-\mathrm{m}_{D_{01}} \operatorname{arctanh}(\mathrm{D}) / \mathrm{D}, \\
\mathrm{LD}^{\prime}=-\mathrm{m}_{D_{02}} \operatorname{arctanh}(D) / D, \\
\mathrm{CD}=\mathrm{m}_{D_{03}} \operatorname{arctanh}(D) / D, \\
\mathrm{LB}=\left(\mathrm{m}_{R_{32}}-\mathrm{m}_{R_{23}}\right) R / 2 \sin R, \\
\mathrm{LB}^{\prime}=\left(\mathrm{m}_{R_{13}}-\mathrm{m}_{R_{31}}\right) R / 2 \sin R, \\
\mathrm{CB}=\left(\mathrm{m}_{R_{12}}-\mathrm{m}_{R_{21}}\right) R / 2 \sin R, \tag{4.32f}
\end{array}
$$

where

$$
\begin{gather*}
R=\arccos \left(\operatorname{tr}\left(\mathbf{M}_{R}\right) / 2-1\right),  \tag{4.33}\\
D=\left(\mathrm{m}_{D_{01}}^{2}+\mathrm{m}_{D_{02}}^{2}+\mathrm{m}_{D_{03}}^{2}\right)^{1 / 2}, \tag{4.34}
\end{gather*}
$$

and the terms $\mathrm{m}_{R_{i j}}$ and $\mathrm{m}_{D_{i j}}(i, j=0, . ., 3)$ respectively indicate matrix elements of $\mathrm{i}^{\text {th }}$ row and $\mathrm{j}^{\text {th }}$ column of $\mathbf{M}_{R}$ and $\mathbf{M}_{D}$.

## Noncommutativity

Before we have seen that the factor of the polar decomposition did not commute [see Eq. (4.22)]. In the case of the Lu-Chipman decomposition something similar happens, the three factors of the Lu-Chipman decomposition give six possible variations of the decomposition that depend on the order in which the factors are multiplied [105]:

$$
\begin{align*}
& \mathbf{M}=\mathbf{M}_{\Delta} \mathbf{M}_{R} \mathbf{M}_{D},  \tag{4.35a}\\
& \mathbf{M}=\mathbf{M}_{\Delta 1} \mathbf{M}_{D 1} \mathbf{M}_{R 1},  \tag{4.35b}\\
& \mathbf{M}=\mathbf{M}_{R 2} \mathbf{M}_{\Delta 2} \mathbf{M}_{D 2},  \tag{4.35c}\\
& \mathbf{M}=\mathbf{M}_{R 3} \mathbf{M}_{R 3} \mathbf{M}_{\Delta 3},  \tag{4.35d}\\
& \mathbf{M}=\mathbf{M}_{D 4} \mathbf{M}_{R 4} \mathbf{M}_{\Delta 4},  \tag{4.35e}\\
& \mathbf{M}=\mathbf{M}_{D 5} \mathbf{M}_{\Delta 5} \mathbf{M}_{R 5}, \tag{4.35f}
\end{align*}
$$

These six equations can be separated in two families, depending on if the depolarizer factor is behind the diattenuating factor(Eqs. (4.35a), (4.35b) and (4.35c))sometimes referred as "forward" family- or in front (Eqs. (4.35d) (4.35e) and $(4.35 f)$ ) of the diattenuating factor -"reverse" family-. A detailed discussion on the decomposition families and how to obtain them is available in [105]. It is
important to stress that although there are some particular coincidences (for example $\mathbf{M}_{R}=\mathbf{M}_{R 1}$, see [105] for more examples) the factors $\mathbf{M}_{R i}, \mathbf{M}_{D i}$ and $\mathbf{M}_{\Delta i}$ are different depending on the order in which they are multiplied. Also the depolarizer factor on the reverse family is that of a depolarizer with diattenuation, while in the forward family is a depolarizer with polarizance (Eq. (4.24)).

Due to the noncommutativity of the polar or the Lu-Chipman decompositions, in the most general case, the matrices resulting from these decompositions have lost their physical interpretability, in the sense that the diattenuation properties of the Hermitian matrix and the retardation characteristics of the unitary matrix do not correspond to the diattenuation and retardation of the original Jones or Mueller matrix. This has been already demonstrated in Ref. [106] by Savenkov.

### 4.3.2. Pseudopolar decomposition

The work on this decomposition started when we noted that the optical effects (CD, CB, LD, etc) obtained form the factors of polar decomposition of a MuellerJones matrix were not equivalent to those obtained from the analytic inversion introduced in this chapter. This is an obvious fact if one attends to the noncommutativity of the matrix factors involved in polar or in the Lu-Chipman decomposition. However we found that this question was forgotten or not sufficiently stressed in some experimental investigations [107-111] and the Lu-Chipman is erroneously taken as a "universal" method to obtain the optical effects of any optical system.

The pseudopolar decomposition remains in close relationship with the polar or the Lu-Chipman decompositions, but, by offering a treatment of the noncommutivity of retarder and diattenuating factors of the decomposition, it is intended to maintain the physical interpretability of the factors of the decomposition. The noncommutative properties of the matrices that describe the optical properties of a system were already noted by Jones [75]. He wrote: "as first attempt, one might try to find a simple way of factoring the matrix $M$ of the crystal into the product of a finite number of simple M-matrices, each of which would represent a simple crystal property, such as circular dichroism, linear birefringence, or isotropic absorption. This effort fails, because the constants which specify the component matrices depend on the order in which the matrices are multiplied".

The theoretical development of the pseudopolar decomposition is partly based on the decomposition of an optical element into infinitesimal sublayers that we already introduced in the previous chapter. At this point we can recuperate Eq. (3.17), that gives the exponential version of Jones matrix of homogenous optical element having both diattenuating and retarding properties:

$$
\begin{equation*}
\mathbf{J}=\exp [-i \mathbf{R}] e^{-i \chi / 2} e^{-i\left[\vec{\sigma} \cdot \overrightarrow{\mathrm{~T}}_{R}+\vec{\sigma} \cdot\left(-i \overrightarrow{\mathrm{~T}}_{D}\right)\right] / 2} \tag{4.36}
\end{equation*}
$$

Note that $\vec{\sigma} \cdot \overrightarrow{\mathrm{T}}_{R}$ and $\vec{\sigma} \cdot \overrightarrow{\mathrm{T}}_{D}$ define two non-commutative matrices $\left(\left[\vec{\sigma} \cdot \overrightarrow{\mathrm{T}}_{R}, \vec{\sigma} \cdot \overrightarrow{\mathrm{~T}}_{D}\right] \neq\right.$

0 ) and that the exponential of their sum cannot be found from the product of their individual exponentials. At this point it is interesting to introduce the so called Zassenhaus formula, which states that the exponential of the sum of two non-commutative operators $X$ and $Y$ can be obtained as an infinite product of exponentials of the operators and their commutators:

$$
\begin{equation*}
e^{t(X+Y)}=e^{t X} e^{t Y} e^{-\frac{t^{2}}{2}[X, Y]} e^{\frac{t^{3}}{6}(2[Y,[X, Y]]+[X,[X, Y]])} e^{t^{4} \ldots} \ldots \tag{4.37}
\end{equation*}
$$

and further terms are given by a recursion relation $[112,113]$. This is an infinite and convergent series [114] and the Zassenhaus exponents can be obtained in terms of nested commutators with growing complexity. Here we will consider only the first and second Zassenhaus exponents because, although more correction terms could be considered, they are enough for most of the applications in polarimetry.

Comparing equation (4.37) with (3.17) we can use the Zassenhaus formula making the identifications $X=\vec{\sigma} \cdot \overrightarrow{\mathrm{T}}_{R}, Y=-i \vec{\sigma} \cdot \overrightarrow{\mathrm{~T}}_{D}$ and $t=-i / 2$. We can write $\mathbf{J}$ as follows:

$$
\begin{equation*}
\mathbf{J} \cong \mathbf{J}_{R} \mathbf{J}_{D} \mathbf{J}_{1 C} \mathbf{J}_{2 C}, \tag{4.38}
\end{equation*}
$$

where

$$
\begin{align*}
& \mathbf{J}_{R}=e^{-i \eta} \exp \left(-i \frac{\mathrm{~T}_{R}}{2} \frac{\overrightarrow{\mathrm{~T}}_{R}}{\mathrm{~T}_{R}} \cdot \vec{\sigma}\right)=e^{-i \eta}\left[\sigma_{0} \cos \frac{\mathrm{~T}_{R}}{2}-\frac{i}{\mathrm{~T}_{R}} \overrightarrow{\mathrm{~T}}_{R} \cdot \vec{\sigma} \sin \frac{\mathrm{~T}_{R}}{2}\right], \\
& \mathbf{J}_{D}=e^{-k} \exp \left(-\frac{\mathrm{T}_{D}}{2} \frac{\overrightarrow{\mathrm{~T}}_{D}}{\mathrm{~T}_{D}} \cdot \vec{\sigma}\right)=e^{-k}\left[\sigma_{0} \cosh \frac{\mathrm{~T}_{D}}{2}-\frac{1}{\mathrm{~T}_{D}} \overrightarrow{\mathrm{~T}}_{D} \cdot \vec{\sigma} \sinh \frac{\mathrm{~T}_{D}}{2}\right], \\
& \mathbf{J}_{1 C}=\exp \left(\frac{\mathrm{A}}{8}\left[\frac{\overrightarrow{\mathrm{~A}}}{\mathrm{~A}} \cdot \vec{\sigma}\right]\right)=\sigma_{0} \cosh \frac{\mathrm{~A}}{8}+\frac{1}{\mathrm{~A}} \overrightarrow{\mathrm{~A}} \cdot \vec{\sigma} \sinh \frac{\mathrm{~A}}{8},  \tag{4.39}\\
& \mathbf{J}_{2 C}=\exp \left(i \frac{\mathrm{~B}}{48}\left[\frac{\overrightarrow{\mathrm{~B}}}{\mathrm{~B}} \cdot \vec{\sigma}\right]\right)=\sigma_{0} \cos \frac{\mathrm{~B}}{48}+\frac{i}{\mathrm{~B}} \overrightarrow{\mathrm{~B}} \cdot \vec{\sigma} \sin \frac{\mathrm{~B}}{48}
\end{align*}
$$

and $\overrightarrow{\mathrm{A}}=\left(\mathrm{A}_{1}, \mathrm{~A}_{2}, \mathrm{~A}_{3}\right)$ and $\overrightarrow{\mathrm{B}}=\left(\mathrm{B}_{1}, \mathrm{~B}_{2}, \mathrm{~B}_{3}\right)$. The expressions for $\mathrm{A}_{i}, \mathrm{~B}_{i}, \mathrm{~A}$, $\mathrm{B}, \mathrm{T}_{R}$ and $\mathrm{T}_{D}$ are given in table 4.2. In Eq. (4.39) we have used common algebra properties related to Pauli matrices to write the exponential in terms of trigonometric and hyperbolic functions. Also in table 4.2 the matrices $\mathbf{J}_{R}, \mathbf{J}_{D}$, $\mathbf{J}_{1 C}, \mathbf{J}_{2 C}$ are written explicitly without making use of the Pauli notation.

We will christen the decomposition of Eq. (4.38) as the pseudo-polar decomposition. Like in the polar-decomposition, $\mathbf{J}_{R}$ is an unitary matrix that changes only the phases of the components of the electric field and $\mathbf{J}_{D}$ is a Hermitian matrix that changes only the amplitudes of the components of the electric field vector. $\mathbf{J}_{1 C}$, which is also Hermitian, and $\mathbf{J}_{2 C}$ are respectively the first and second correction matrices.

More terms of the expansion series of Eq. (4.37) could be considered, and this would give us more correction matrices (we could name them $\mathbf{J}_{3 C}, \mathbf{J}_{4 C}$, etc). However, note that the elements of the first correction matrix are quadratic with the anisotropies, while the elements of the second correction matrix are cubic.

Chapter 4. Inversion and decomposition of Mueller matrices

Table 4.2.: Factorized Jones Matrix, $\mathbf{J} \cong \mathbf{J}_{R} \mathbf{J}_{D} \mathbf{J}_{1 C} \mathbf{J}_{2 C}$


Third and higher order correction matrix would respectively have elements with fourth and higher order dependence in anisotropies. Provided that for most of the experimentally-described systems the basic building blocks for the correction terms (the elements $A_{1}, A_{2}$ and $A_{3}$ defined in table 4.2) are much smaller than one, the correction matrices rapidly approach to the identity matrix.

It is important to stress again that all the linear and circular anisotropies involved in this factorization are the real anisotropies of the system. Conversely, the linear and circular birefringence and dichroism involved in the polar decomposition are mathematical entities that, in general, do not correspond with the anisotropies of the sample. Moreover, the pseudo-polar decomposition verifies:

$$
\begin{equation*}
\mathbf{J} \cong \mathbf{J}_{R} \mathbf{J}_{D} \mathbf{J}_{1 C} \mathbf{J}_{2 C} \cong \mathbf{J}_{D} \mathbf{J}_{R} \mathbf{J}_{1 C}^{-1} \mathbf{J}_{2 C}^{-1} \tag{4.40}
\end{equation*}
$$

This property can be verified from Eq. (4.37): permuting $X$ with $Y$ does not alter the first two terms of the Zassenhaus formula, but the fact that $[X, Y]=-[Y, X]$
modifies the exponent sign of all the following factors.
In the way the matrices $\mathbf{J}_{R}$ and $\mathbf{J}_{D}$ are build (see Eqs. (4.39)), it can be clearly seen that the pseudo-polar decomposition also satisfies the following property:

$$
\begin{equation*}
\overrightarrow{\mathrm{T}}_{D}(\mathbf{J})=\overrightarrow{\mathrm{T}}_{D}\left(\mathbf{J}_{D}\right) \quad \text { and } \quad \overrightarrow{\mathrm{T}}_{R}(\mathbf{J})=\overrightarrow{\mathrm{T}}_{R}\left(\mathbf{J}_{R}\right) \tag{4.41}
\end{equation*}
$$

which means that the diattenuation vectors of $\mathbf{J}_{D}$ and the retardation vector of $\mathbf{J}_{R}$ correspond with those of $\mathbf{J}$. This indicates that the dichroic and birefringent physical parameters of original matrix $\mathbf{J}$ (Eq. + (3.18)) are respectively preserved and separated in factors $\mathbf{J}_{D}$ and $\mathbf{J}_{R}$ (eqs. (4.39)).

There are some cases in which both first and second corrections do not have any contribution, which means that $\mathbf{J}_{1 C}=\mathbf{J}_{2 C}=\sigma_{0}$. From the matrices and the definitions given in table 4.2, we find that the conditions for this to happen are:

$$
\begin{align*}
& \mathrm{CBLD}^{\prime}-\mathrm{LB}^{\prime} \mathrm{CD}=0, \\
& \mathrm{CDLB}-\mathrm{LDCB}=0,  \tag{4.42}\\
& \mathrm{LBLD}^{\prime}-\mathrm{LB}^{\prime} \mathrm{LD}=0 .
\end{align*}
$$

Leaving aside the mathematical restrictions we can derive from this set of equations and recalling the physical meaning of the parameters involved in these equations, we can easily realize that there are two common types of anisotropic optical media that satisfy the three conditions of Eqs. (4.42). The first one corresponds to a circular birefringent and dichroic media $\mathrm{CD}, \mathrm{CB} \neq 0$ without linear anisotropies $\mathrm{LB}=\mathrm{LD}=\mathrm{LD}^{\prime}=\mathrm{LB}^{\prime}=0$. Typically, solutions of chiral molecules constitute a good example for this kind of media although, in some cases, they have associated linear anisotropies too. The second one corresponds to a nonchiral high symmetry sample in which the principal axes for linear retardation and for linear dichroism coincide $\mathrm{LD} / \mathrm{LB}=\mathrm{LD}^{\prime} / \mathrm{LB}^{\prime}$. Uniaxial crystals and flowor field-oriented molecules are common optical media in which these conditions might be found. For these two media we can certainly write $\mathbf{J}=\mathbf{J}_{R} \mathbf{J}_{D}=\mathbf{J}_{D} \mathbf{J}_{R}$ and, therefore, the pseudo-polar decompositions transforms into the polar decomposition. This demonstrates that, in these media, the polar decomposition can be applied without losing the physical significance of the involved anisotropies. The media in which Eqs. (4.42) hold are media without non-commuting optical properties and, in those kind of media, polar decomposition can be safely applied.

We can derive the Mueller-Jones matrices corresponding to the Jones matrices given in table 4.2 by using Eq. (1.29) or Eq. (3.20). The Mueller-Jones matrices $\mathbf{M}_{R}$ and $\mathbf{M}_{D}$ associated to $\mathbf{J}_{R}$ and $\mathbf{J}_{M}$ have been already given in Eq. (3.24) and Eq. (3.25) respectively. The Mueller-Jones matrix, $\mathbf{M}_{1 C}$, corresponding to the

Chapter 4. Inversion and decomposition of Mueller matrices
correction term result into a symmetric matrix:

$$
\mathbf{M}_{1 C}=\left[\begin{array}{cccc}
\cosh ^{2} \frac{\mathrm{~A}}{8}+\sinh ^{2} \frac{\mathrm{~A}}{8} & \mathrm{~A}_{1} \gamma & \mathrm{~A}_{2} \gamma & \mathrm{~A}_{3} \gamma  \tag{4.43}\\
\mathrm{~A}_{1} \gamma & 1+2 \mathrm{~A}_{1}^{2} \delta & 2 \mathrm{~A}_{1} \mathrm{~A}_{2} \delta & 2 \mathrm{~A}_{1} \mathrm{~A}_{3} \delta \\
\mathrm{~A}_{2} \gamma & 2 \mathrm{~A}_{1} \mathrm{~A}_{2} \delta & 1+2 \mathrm{~A}_{2}^{2} \delta & 2 \mathrm{~A}_{2} \mathrm{~A}_{3} \delta \\
\mathrm{~A}_{3} \gamma & 2 \mathrm{~A}_{1} \mathrm{~A}_{3} \delta & 2 \mathrm{~A}_{2} \mathrm{~A}_{3} \delta & 1+2 \mathrm{~A}_{3}^{2} \delta
\end{array}\right]
$$

with $\gamma=\frac{1}{A} \sinh \frac{\mathrm{~A}}{4}$ and $\delta=\frac{1}{\mathrm{~A}^{2}} \sinh ^{2} \frac{A}{8}$. The calculation of the Mueller matrix associated to the second correction term, $\mathbf{M}_{2 C}$, requires some more algebra than the matrices above, since $\vec{B}$ is a complex vector. After some algebra we find

$$
\left.\mathbf{M}_{2 C}=\left[\begin{array}{cc}
\rho & -\varepsilon \mathrm{B}_{1}^{i}+\zeta \mathrm{B}_{1}^{r} \\
+(\xi / 2)\left(\mathrm{B}^{*} \cdot \overrightarrow{\mathrm{~B}}\right) & +\xi\left(\mathrm{B}_{3}^{r} \mathrm{~B}_{2}^{i}-\mathrm{B}_{2}^{r} \mathrm{~B}_{3}^{i}\right) \\
-\varepsilon \mathrm{B}_{1}^{i}+\zeta \mathrm{B}_{1}^{r} & \rho+\xi\left(\left(-\mathrm{B}_{1}^{i}\right)^{2}+\left(\mathrm{B}_{1}^{r}\right)^{2}\right) \\
-\xi\left(\mathrm{B}_{3}^{r} \mathrm{~B}_{2}^{i}-\mathrm{B}_{2}^{r} \mathrm{~B}_{3}^{i}\right) & -(\xi / 2)\left(\mathrm{B}^{*} \cdot \overrightarrow{\mathrm{~B}}\right) \\
-\varepsilon \mathrm{B}_{2}^{i}+\zeta \mathrm{B}_{2}^{r} & -\varepsilon \mathrm{B}_{3}^{r}-\zeta \mathrm{B}_{3}^{i}  \tag{4.44}\\
-\xi\left(\mathrm{B}_{1}^{r} \mathrm{~B}_{3}^{i}-\mathrm{B}_{3}^{r} \mathrm{~B}_{1}^{i}\right) & +\xi\left(\mathrm{B}_{1}^{r} \mathrm{~B}_{2}^{r}+\mathrm{B}_{1}^{i} \mathrm{~B}_{2}^{i}\right) \\
-\varepsilon \mathrm{B}_{3}^{i}+\zeta \mathrm{B}_{3}^{r} & \varepsilon \mathrm{~B}_{2}^{r}+\zeta \mathrm{B}_{2}^{i} \\
-\xi\left(-\mathrm{B}_{1}^{r} \mathrm{~B}_{2}^{i}+\mathrm{B}_{2}^{r} \mathrm{~B}_{1}^{i}\right) & -\xi\left(-\mathrm{B}_{3}^{r} \mathrm{~B}_{1}^{r}-\mathrm{B}_{3}^{i} \mathrm{~B}_{1}^{i}\right)
\end{array}\right] \begin{array}{cc}
-\varepsilon \mathrm{B}_{2}^{i}+\zeta \mathrm{B}_{2}^{r} & -\varepsilon \mathrm{B}_{3}^{i}+\zeta \mathrm{B}_{3}^{r} \\
+\xi\left(\mathrm{B}_{1}^{r} \mathrm{~B}_{3}^{i}-\mathrm{B}_{3}^{r} \mathrm{~B}_{1}^{i}\right) & +\xi\left(-\mathrm{B}_{1}^{r} \mathrm{~B}_{2}^{i}+\mathrm{B}_{2}^{r} \mathrm{~B}_{1}^{i}\right) \\
\varepsilon \mathrm{B}_{3}^{r}+\zeta \mathrm{B}_{3}^{i} & -\varepsilon \mathrm{B}_{2}^{r}-\zeta \mathrm{B}_{2}^{i} \\
+\xi\left(\mathrm{B}_{1}^{r} \mathrm{~B}_{2}^{r}+\mathrm{B}_{1}^{i} \mathrm{~B}_{2}^{i}\right) & -\xi\left(-\mathrm{B}_{3}^{r} \mathrm{~B}_{1}^{r}-\mathrm{B}_{3}^{i} \mathrm{~B}_{1}^{i}\right) \\
\rho+\xi\left(\left(\mathrm{B}_{2}^{r}\right)^{2}+\left(-\mathrm{B}_{2}^{i}\right)^{2}\right) & \varepsilon \mathrm{B}_{1}^{r}+\zeta \mathrm{B}_{1}^{i} \\
-(\xi / 2)\left(\mathrm{B}^{*} \cdot \overrightarrow{\mathrm{~B}}\right) & -\xi\left(-\mathrm{B}_{3}^{r} \mathrm{~B}_{2}^{r}-\mathrm{B}_{3}^{i} \mathrm{~B}_{2}^{i}\right) \\
-\varepsilon\left(-\mathrm{B}_{1}^{r} \mathrm{~B}_{2}^{r}-\zeta \mathrm{B}_{1}^{i} \mathrm{~B}_{2}^{i}\right) & \rho+\xi\left(\left(\mathrm{B}_{3}^{i}\right)^{2}+\left(-\mathrm{B}_{3}^{r}\right)^{2}\right) \\
& -(\xi / 2)\left(\overrightarrow{\mathrm{B}}^{*} \cdot \overrightarrow{\mathrm{~B}}\right)
\end{array}\right]
$$

where

$$
\begin{gather*}
\varepsilon=\left(\mathrm{B}^{i} \sinh \frac{\mathrm{~B}^{i}}{24}+\mathrm{B}^{r} \sin \frac{\mathrm{~B}^{r}}{24}\right) / \mathrm{BB}^{*}, \\
\zeta=\left(\mathrm{B}^{i} \sin \frac{\mathrm{~B}^{r}}{24}-\mathrm{B}^{r} \sinh \frac{\mathrm{~B}^{2}}{24}\right) / \mathrm{BB}^{*},  \tag{4.45}\\
\xi=\left(\cosh \frac{\mathrm{B}^{i}}{24}-\cos \frac{\mathrm{B}^{r}}{24}\right) / \mathrm{BB}^{*}, \\
\rho=\left(\cosh \frac{\mathrm{B}^{i}}{24}+\cos \frac{\mathrm{B}^{r}}{24}\right) / 2
\end{gather*}
$$

and the superscript $r$ refers to the real part and the superscript $i$ refers to the imaginary part.

The pseudo-polar decomposition for nondepolarizing Mueller can be written as follows:

$$
\begin{equation*}
\mathbf{M} \cong \mathbf{M}_{R} \mathbf{M}_{D} \mathbf{M}_{1 C} \mathbf{M}_{2 C} \tag{4.46}
\end{equation*}
$$

where $\mathbf{M}$ is the general Mueller matrix given in Eq. (3.21).

## Application to experimental matrices

Until now we have introduced the pseudo-polar decomposition and we have shown how to build up the factors from the physical effects defined in table 1.1. The purpose of this section is to briefly examine how to find the factors of the decomposition (this is $\mathbf{J}_{R}, \mathbf{J}_{D}, \mathbf{J}_{1 C}$ or $\mathbf{J}_{2 C}$ or $\mathbf{M}_{R}, \mathbf{M}_{D}, \mathbf{M}_{1 C}$ and $\mathbf{M}_{2 C}$ ) from one input matrix $\mathbf{J}$ o $\mathbf{M}$. In this case we will focus our attention only in the Mueller matrices, which are the ones that normally are determined experimentally.

The procedure to obtain the pseudopolar decomposition on a given experimental Mueller matrix is based in a process of filtering of the experimental Mueller matrix and in a recursive application of the the algorithm to obtain the LuChipman decomposition that we have presented in section 4.3.1. In most cases the polar decomposition will offer a reasonable estimation of the physical parameters involved in the decomposition although, as discussed before, it may be not accurate if there are noncommuting optical properties. Therefore we can use the Lu-Chipman decomposition to find a first estimate of $\mathrm{LB}, \mathrm{LD}, \mathrm{LB}^{\prime}, \mathrm{LD}^{\prime}, \mathrm{CD}$ and CB.

Eqs. (4.43) and (4.44) describe the way to calculate $\mathbf{M}_{1 C}$ and to $\mathbf{M}_{2 C}$ from the calculated optical properties, so that an estimation of these two correction matrices can be found. We will denote this first approximation as $\mathbf{M}_{1 C}^{(0)} \mathbf{M}_{2 C}^{(0)}$. We recall that these two matrices account for the noncommutative optical properties of the medium and, as shown in section 2, in the cases where the non-commutative optical properties vanish (Eqs. (4.42)) the polar and pseudo-polar decompositions coincide. As long as we are able to find these two correction matrices, we will use them to filter the experimental Mueller matrix in a way that the polar decomposition of the filtered matrix matches the pseudo-polar decomposition of the experimental matrix.

Once $\mathbf{M}_{1 C}^{(0)}$ and $\mathbf{M}_{2 C}^{(0)}$ are calculated we can define a new matrix, $\mathbf{M}_{e}^{\prime(1)}$, that, in some sense, will be the experimental matrix but corrected, for the non-commutative optical properties:

$$
\begin{equation*}
\mathbf{M}_{e}^{\prime(1)}=\mathbf{M}_{e}\left(\mathbf{M}_{2 C}^{(0)}\right)^{-1}\left(\mathbf{M}_{1 C}^{(0)}\right)^{-1} \tag{4.47}
\end{equation*}
$$

Note that the idea to make this definition comes from the pseudo-polar decomposition as it is given in Eq. (4.46), where the $\mathbf{M}_{1 C}^{(0)}$ and $\mathbf{M}_{2 C}^{(0)}$ factors have been moved to the left part of the equation. Certainly, we can apply again the LuChipman decomposition $\mathbf{M}_{e}^{(1)}$ and we will be able to calculate some new values for the physical effects. Now these values will be somehow more accurate than in
the previous step since our corrected experimental Mueller matrix, $\mathbf{M}_{e}^{\prime(1)}$, would already have less non-commutative optical properties than the original. Again, after finding the physical parameters and using equations (4.43) and (4.44) we can calculate a new version of the correction matrices $\mathbf{M}_{1 C}^{(1)}$ and $\mathbf{M}_{2 C}^{(1)}$ which, again, can be used to filter the experimental matrix. Clearly this process can be generalized and, at each step, we can find a better correction for the original experimental Mueller matrix:

$$
\begin{equation*}
\mathbf{M}_{e}^{\prime(i)}=\mathbf{M}_{e}\left(\mathbf{M}_{2 C}^{(i-1)}\right)^{-1}\left(\mathbf{M}_{1 C}^{(i-1)}\right)^{-1} \tag{4.48}
\end{equation*}
$$

in which each time we apply one Lu-Chipman decomposition to filter the experimental Mueller matrix the superscript ( $i$ ) increases in one unit.

At each step we will have a better correction for the experimental Mueller matrix, which implies that the polar decomposition will produce more accurate results. We will arrive clearly to the situation in which the polar decomposition factors $\widetilde{\mathbf{M}}_{R}$ and $\widetilde{\mathbf{M}}_{D}$ of the corrected experimental matrix will be mostly the same that the factors $\mathbf{M}_{R}$ and $\mathbf{M}_{D}$ of the pseudo-polar decomposition of the experimental matrix. After a certain number of steps $(n)$ the condition $\| \mathbf{M}_{e}-$ $\mathbf{M}_{\Delta}^{(n)} \widetilde{\mathbf{M}}_{R}^{(n)} \widetilde{\mathbf{M}}_{D}^{(n)} \mathbf{M}_{1 C}^{(n)} \mathbf{M}_{2 C}^{(n)} \| \simeq 0$, where the bars denote the Frobenius norm ${ }^{3}$, will fulfill and the product $\mathbf{M}_{\Delta}^{(n)} \widetilde{\mathbf{M}}_{R}^{(n)} \widetilde{\mathbf{M}}_{D}^{(n)} \mathbf{M}_{1 C}^{(n)} \mathbf{M}_{2 C}^{(n)}$ could be interpreted as the pseudo-polar decomposition of $\mathbf{M}_{e}$.

As an example of the application of the polar decomposition and the pseudopolar decomposition, we have inverted two Mueller-Jones matrices that were generated using some representative values of CD, CB, LD, LB, LD' and LB'. The values for these effects obtained from the polar and pseudopolar decompositions are displayed in Table 4.3. The table also includes the values for CD, CB, LD, LB, LD' and LB' obtained using the inversion method described in the previous chapter. The analytic inversion offers an exact solution to the problem, i. e. we obtain again the same parameters used for the generation, while the results on the others methods, specially for the polar decomposition, largely depend on the magnitude of the effects and/or on their noncommutativity. In Case 1 all the optical effects are small and the noncommutative effects (that always are second and superior order effects) can be omitted, thus obtaining a satisfactory result by any of the three methods. It is interesting to note that here the pseudopolar decomposition although it is an approximate method is able to reproduce exact results with an accuracy up to the forth decimal. Case 2 represents a situation in which much bigger effects are present and, in such a way, that they generate significant noncommutativity between the diattenuating and retarding effects. Here the pseudopolar decomposition offers a much better approximation to the exact

[^17]Table 4.3.: Comparison of methods to analyze homogeneous Mueller-Jones matrices

${ }^{a}$ The analytic inversion gives the original parameters used for the generation of the MuellerJones matrices.
${ }^{b}$ Pseudopolar decomposition have been calculated with two correction terms and 100 iterations.
result provided by the analytic inversion than the polar decomposition, that fails to find accurate results specially for CD.

Fig. 4.1 shows the values of the polarization fraction ( $\beta$, see Eq. (1.33)), CD and CB obtained from experimental Mueller matrices for a water solution of Jaggregates of the pseudocyanine dye in a 0.1 mm path length cuvette. The only difference between case $a$ and case $b$ is the amount of aggregate that was present in the cuvette. The samples can be well described as a homogeneous anisotropic medium that contained linear birefringence and dichroism, due to the oriented elongated aggregates, and as well as circular dichroism and birefringence, due to the chirality of the involved molecules. The Mueller matrices were spectroscopically measured with the instrument described in chapters 5 and 6 and, later, studied by the different methods we have described in this chapter.

Chapter 4. Inversion and decomposition of Mueller matrices


Figure 4.1.: For an optical element with the optic axis lying in the xy plane, the laboratory coordinates $x$ and $y$ (reference basis) are related to the crystallographic coordinates (natural basis) by a simple rotation.

The top panels of Fig. 4.1 indicate that for the studied samples there exists a band of with depolarization around 570 nm that is caused by light scattering by aggregates of electronically interacting chromophores. At these wavelengths the depolarization is important for case a $(\beta \sim 0.5)$, while it is much less significative for case $b(\beta \sim 0.8)$. The CD and CB panels of the figure show a comparison of the results offered by the inversion and decomposition methods of the experimental Mueller matrices. In general the three methods give coincident results out of the area of depolarization but at the polarization band they offer different solutions. For case $a$ the results of the analytic inversion seem to be not really accurate, which can be attributed to the fact that the process of finding a nondepolarizing estimate for Mueller matrices containing a strong depolarization does not offer good results because there is not a single nondepolarizing matrix that can be
related to the experimental matrix. In contrast, for case $b$ we see that the product decompositions do not handle the CB as well as the inversion: the analytic inversion method gives CB values that are more Kramers-Kronig consistent with a CD bisignated band. We can conclude in this example we would use the pseudopolar decomposition (pseudopolar and polar decomposition often yields similar results, but we prefer those of the pseudopolar for being more accurate) method to study case $a$, and the analytic inversion method to analyze case $b$. There is not a universal criteria for choosing one method or other to analyze experimental measurements. Often the best approach is to test them all. In the majority cases they will yield to very similar results, and in the case of different results it will be possible to use similar arguments to the ones we have used with this example to chose one method over the other.

### 4.3.3. Symmetric decomposition

The symmetric decomposition was recently proposed by Ossikovsky [115] and it constitutes one of the most interesting approaches to study physical systems that introduce significant depolarization. The decomposition takes the form:

$$
\begin{equation*}
\mathbf{M}=\mathbf{M}_{D 2} \mathbf{M}_{R 2} \mathbf{M}_{\Delta d} \mathbf{M}_{R 1}^{T} \mathbf{M}_{D 1} \tag{4.49}
\end{equation*}
$$

The main particularity of this decomposition is that depolarizer factor is placed "in the middle" of a symmetric optical sequence and that it is diagonal:

$$
\mathbf{M}_{\Delta d}=\left[\begin{array}{cccc}
d_{0} & 0 & 0 & 0  \tag{4.50}\\
0 & d_{1} & 0 & 0 \\
0 & 0 & d_{2} & 0 \\
0 & 0 & 0 & d_{3}
\end{array}\right]
$$

This term contrasts with the depolarizer factor that was found in the Lu-Chipman decomposition (see Eq. (4.24), which was nondiagonal because it corresponded to a depolarizer with polarizance. It is known both from theory and experiment that a great number of depolarizing media are better characterized by a diagonal depolarizers.

We will not describe in this section the algorithm to apply the symmetric decomposition. Although the algorithm is relatively simple its application to experimental matrices contain some subtleties that require discussion. The details can be found in Refs. $[115,116]$.

If the depolarizer of the symmetric decomposition is "close" to the identity matrix (i.e. the depolarization is small) then there may be no clear separation between the factors placed before and after the depolarization factor of Eq. (4.49). If this happens probably the factors of the decomposition lose the physical significance. Therefore this decomposition is, in practice, more intended to the study
of Mueller matrices corresponding to strongly depolarizing media, and, for example, can be very useful when trying to localize the various effects of a sample that for example is composed of two well-differentiated parts.

The algorithms of all the product decompositions described in this chapter have been incorporated in a computer program that acts as a complement to the software developed for the instrument described in the following part of the work . This program takes series of measured Mueller matrix (either from spectroscopic measurements or space resolved measurement) and calculates the type of decomposition that the user selects. Usually these decompositions are useful to obtain a first (and sometimes definitive) interpretation of experimental measurements.

## Part III.

## Experimental sets

## Chapter 5.

## Two-modulator generalized ellipsometer

The experimental device used to develop the present work has been a twomodulator generalized ellipsometer (2-MGE). This apparatus was customly build in a long process that started in 2006 and reached the status of development that will be detailed in this part of the work.

The 2-MGE was introduced by G. E. Jellison and F. A. Modine in 1997 with two papers: one dealing with the theory behind the analysis of the timedependent detected signal [117] and the other focused on the experimental description and the calibration of the instrument [118]. Many of the contents of this chapter are based on these papers, and we have maintained the same notation used there, so, together with the patent [119], they can be an appropriate complement for the interested reader. Besides some subtle differences in the experimental configuration between our instrument and that of Jellison an Modine, in this work we have tried to provide a more step-by-step approach to the calculations required to interpret the detected waveform.

### 5.1. Introduction

The 2-MGE is an instrument that measures the change of light polarization upon interacting with a sample. The 2 -MGE can operate in either reflection or transmission. In reflection, it acts as a generalized ellipsometer, measuring the standard ellipsometry parameters, as well as the cross-polarization parameters. In transmission, it measures all parameters associated with a general diattenuation and retarder. All the elements of the transmission or reflection Mueller matrix of a sample are accessible to this apparatus, but not in single measurement, and different measurement need to be combined.

The 2-MGE is based in the use of photoelastic modulators (PEMs). For years these phase modulators have been used in ellipsometry, allowing high sensitivity and high acquisition rate that are required for spectral measurements, in situ applications, and fast processes monitoring. The first report of a spectroscopic ellipsometer that used a PEM was published by Jasperson and Schnatterly in 1970 [120], but phase modulated ellipsometry did not grew more popular until '80s and '90s due to the availability of fast computers. Drévillon and coworkers were pioneers in performing a digital Fourier analysis on the intensity waveform
[121, 122]. Canillas et al. developed a phase-modulated ellipsometer using a Fourier transform spectrometer to work in the infrared [123, 124]. In 1997 G. E. Jellison and F. Modine published in a series of two papers the first implementation of an ellipsometer that used two free running PEMs of different frequencies and introduced the acronym 2-MGE for such instrument [117,118].

Instruments based on four PEMs of different frequencies have also been suggested [125]. Such polarimeter would be capable of measuring all the 16 elements of a Mueller matrix in a single measurement and without any moving parts. However, to our knowledge, no experimental realizations of this setup have been developed to the moment.

### 5.2. The photoelastic modulator

The working principle of a modern photoelastic modulator (PEM) is based on the optical birefringence induced by periodical stress. A PEM is composed of an optically transparent material with a high elasto-optic efficiency and low mechanical dissipation (e.g. fused quartz) and of an oscillating piezoelectric transducer. The light beam passes through the optical element which is mechanically stressed by a periodic strain variation (typically at $20-80 \mathrm{kHz}$ ) by the transducer. This oscillating stress causes changes in the the refractive index of the optical element, exhibiting a birefringence proportional to the strain. Therefore, the optical element acts as a dynamic wave plate so, if a polarizer is fixed before the PEM, where the azimuthal angle of the polarizer is not aligned with the modulation axis of the PEM, dynamically elliptically polarized light is generated with the ellipticity changing at the frequency of the PEM.

The modern PEM was invented in 1969 by James Kemp [126], then manufactured by a private company in the USA (Hinds Instruments, Inc., Hillsboro, Oregon, USA). That design has since been dominant in almost all PEM applications, yet the construction and performance have changed little. A second type PEM was invented by Canit and Badoz of France in 1983 [127]. The Kemp design, utilizes a - 18.5 degree X-cut quartz crystal as the excitation source, which delivers a nearly pure longitudinal wave. The Canit-Badoz design, uses a thin slab of piezoelectric ceramic transducer that is adhered to the optical element. This design is less vulnerable to temperature changes but causes considerable unwanted vibrations and acoustic reflections.

The photoelastic modulators used in our 2-MGE were manufactured by Hinds Instruments and consist of a piece of precisely-cut crystalline quartz (the piezoelectric transducer) mechanically coupled to a piece of optically-isotropic fused quartz (the optical element). An ac voltage is applied between the front and back faces of the crystalline quartz component in order to drive it at its resonant frequency. The light beam passes through the central region of the fused quartz
bar and undergoes a retardation that is periodic at the resonant frequency of the crystal quartz bar. This frequency is dependent only on the size and shape of crystalline and fused quartz elements. The two PEMs of our instrument have frequencies $\sim 50 k H z$ and $\sim 60 k H z$. As a PEM is a resonant device and its operating frequency is very stable although it is somewhat dependent on temperature. The amplitude of modulation is usually set with an external dc voltage, and the corresponding phase retardation is dependent on the wavelength of light and can be easily calibrated.

As a result of its design a PEM has unique optical features, such as high modulation purity and efficiency, broad spectral range, high power handling capability, large acceptance angle, large useful aperture and high retardation stability [128]. These features make the PEM the best choice for polarization modulation in applications that require high sensitivity, because it offers a modulation quality unsurpassed by other types of electro-optic modulators or liquid crystal modulators.

A PEM may exhibit a small static strain in the optical element that is independent of dynamic retardation. If it is collinear with the modulation axis of the optical element it is possible to calibrate it and to correct the data for its effect. In this case the time-dependent retardation of a PEM is normally expressed as

$$
\begin{equation*}
\delta(t)=A \sin (\omega t+\phi)+\delta_{0} \tag{5.1}
\end{equation*}
$$

where $A$ is the amplitude of modulation, $2 \pi \omega$ is the frequency of the modulator, $\phi$ is the phase of the modulator, and $\delta_{0}$ is the static retardation. For a more complete description about PEMs and their use in ellipsometry see for example [129] and the chapter about polarization modulation ellipsometry by Jellison and Modine in [130].

### 5.3. Experimental configuration

The 2-MGE, understood as a whole from apparatus, is composed of several different elements. In order to review the characteristics of all of them and their position and functionality in the instrument we will distribute them in two big groups: optical components and electronic hardware.

### 5.3.1. Optical components

This group of components comprises all the elements that affect or modify the light beam from its generation until its detection. Fig. 5.1 shows and scheme of all optical elements as well as a qualitative ray trace of the light beam as it goes through them. Table 5.1 lists all the optical elements displayed in Fig. 5.1, with their respective brand name and model, and other relevant characteristics.

Table 5.1.: Enumeration of light related components

| Element | Brand, model | Details |
| :--- | :--- | :--- |
| Light Source | Oriel | Xe 75 W short arc Lamp |
| Optical fiber 0 | Oriel LLS308 | $800 \mu \mathrm{~m}$ quartz core, solarization re- <br> sistant |
| Focusing optics | Horiba Jobin- | Working distance $\sim 50 \mathrm{~mm}$. Avail- <br> Yvon MicroSpot <br> able spots: $\sim 1500, ~ 200,100 ~$ m |
| Polarizers | Fichou | $\mathrm{MgF}_{2}$ Rochon polarizer. Suitable <br> for UV |
|  | Hinds Instru- | Fused Si, $\nu_{0} \sim 50 \mathrm{kHz}$ |

In Fig. 5.1 the light source is a 75 -watt xenon arc lamp that is focused on the end of a quartz fiber optic using a condenser lens. The other end of the fiber optic is attached to the light entrance of the focusing optics module which is composed of two mirrors and focuses light at a 50 cm from the module. The spot diameter can be tuned by changing the diameter of a pinhole mask that is placed centered at the end of the fiber. Three different pinhole masks are available with diameters $1 \mathrm{~mm}, 0.100 \mathrm{~mm}$ and 0.050 mm , that respectively translate in approximate spot sizes of $1.5 \mathrm{~mm}, 0.200 \mathrm{~mm}$ and 0.100 mm .

Light exiting the focusing module passes through the PSG, which consists of a polarizer-PEM pair. The polarizer is attached to the PEM by using a precision manual rotator and is oriented at $\theta_{b 0}=45^{\circ}$ with respect to the longitudinal axis of the PEM. This PEM has a resonant frequency of 49986 Hz . The complete PSG is attached to a precision automatic step-by-step rotator.


Figure 5.1.: Schematic drawing of the optical elements involved in the 2-MGE. Light beam goes from left to right.

The sample is placed at the focusing point thats falls between the PSG and the PSA. The PSA consist of another PEM-polarizer pair. The nominal frequency of operation of this PEM is 60 kHz . As in the PSG, here the polarizer is also mounted on a precision manual rotator and oriented at $\theta_{b 1}=45^{\circ}$ with respect to the longitudinal axis of this PEM. The PSA is attached to another precision automatic step-by-step rotator. At the end of the PSA there is a lens that focuses the light onto the core of a optical fiber with a core diameter of $400 \mu \mathrm{~m}$.

The light from the optical fiber passes through an optical filter (to eliminate second-order effects) before entering the fiber optics coupling accessory of the monochromator. This accessory uses a pair of mirrors to image the light coming from the fiber to the entrance slit of the monochromator. The light intensity is detected at the output slit by using a photomultiplier tube (PMT).

### 5.3.2. Electronic hardware

A considerable number of electronic hardware pieces are required to run a 2 MGE. They permit the appropriate transformation of the detected light into a digital signal and the automatization of the measurement process. Table 5.2 enumerates the fundamental electronic elements of the 2-MGE, and some of they main features are also displayed. In Fig. 5.2 there is a schematic representation of connections among the different elements. With the exception of the positioning
and alignment of the sample the rest of functions are automatic and can be controlled with a personal computer.

Table 5.2.: Enumberation of electronic hardware

| Element | Brand, model | Details |
| :---: | :---: | :--- |
| Rotator Units | Newport UE31PP (UR80) | Custom controller |
| Linear translation units | UTS50CC | SMC100CC controller, 0.1 |
|  |  | $\mu m$ precision |
| PEM 0 control | PEM-90 controller | Remote voltage control |
| PEM 1 control | PEM-100 controller | RS232 control |
| Filter wheel | Thorlabs FW102B | USB control |
| Preamplifier | Hammamatsu C7319 | gain of 10 5 |
| PMT control circuit | CRIC | Custom design, uses SDS |
|  |  | HV source |
| Trigger circuit |  | Custom design, selectable |
|  |  | gate, 0.5 ${ }^{\circ}$ used |
| Waveform digitizer | Spectrum MI3130 | 2 channel 12 bit A/D with |
| Personal Computer | Custom PC | 25 MS/s |
|  |  | Core2Duo, 1gb RAM, |
| Oscilloscope | Tektronix | 4 Windows XP |

In our the 2-MGE the motion control elements consist of two precision stepper rotation units that azimuthally rotate the whole PSG and PSA and two precision linear translation units that allow to conveniently position the sample in the xy plain. These four motion elements are computer controlled by the software package written for the 2-MGE.

The signal from the PMT is in the form of a photocurrent, and is first converted to a voltage by using a current-mode preamplifier. The signal coming from the preamplifier is fed to a feedback control circuit for the PMT. This feedback circuit has been specifically developed for our 2-MGE and its objective is to dynamically regulate the high voltage delivered to the dynode chain of the PMT. The circuit is designed so that the dc part of the signal coming from the preamplifier is kept at a constant value, that can be chosen by the user (normally we keep dc at 0.5 V ). Thus high voltage given by the PMT feedback circuit will automatically change by increasing (or decreasing) depending on whether the number or photons arriving to the PMT decreases (or increases). The adaptative gain provided by this circuit is essential to have a high dynamic range during a spectroscopic measurement


Figure 5.2.: Schematic drawing of the 2-MGE and associated crontrol circuitry.
which, otherwise, would be limited and the 12-bit resolution of our digitizer. A circuit of this type is described at [131]. The voltage applied to the PMT is permanently visible by means of a LCD display installed in the same circuit enclosure; this feature is useful for the user of the 2-MGE for alignment (usually the best alignment is achieved when this value is minimized) or even to recognize situations when no light arrives to the detector.

The voltage waveform from the preamplifier is digitized by a high-speed 12 bits digitizer installed as a supplementary board in the control computer. Several parameters of the digitization can be controlled, being the most significant the acquisition time, that is usually kept at $0.5 \mu s$ (corresponds to a samplerate of 2000000 samples in a second), and the length of the captured waveform that is keep at 16384. The waveform capture is initialized by a trigger pulse that comes from a coincidence circuit that generates a trigger event each time the monitored reference outputs of the PEM's are in phase. A second channel of the digitizer is used to measure the voltage applied to the PMT (it has been previously attenuated by a factor 1000 so that no longer is a HV), this measure is not a requirement for a $2-\mathrm{MGE}$, but for some single-wavelengths measurement is useful to distinguish sample area's with different transmissivity.

Each PEM is a resonant device, in which only the amplitude of modulation can be controlled by electronics. The frequency and phase are own characteristics of PEM and cannot be externally adjusted. The coincidence circuit allows to find the phase coincidences between the reference signals coming from the PEM's. As the period for the two modulators are $T_{0} \simeq 20 \mu s$ and $T_{1} \simeq 16.67 \mu s$. After five complete cycles of PEM 0, and after six complete cycles of PEM 1 (repeat time of $T \simeq 100 \mu s$ ) we found we found that these two modulators come very close to repeating. The triggering circuit has an adjustable gate time that determines the accuracy of the phase coincidence: the smaller the gate time, the fewer trigger events, but more accurate the phase. For standard measurements the gate time of our coincidence circuit is adjusted so that the monitored reference outputs are within $0^{\circ} \pm 0.5^{\circ}$.

The entire instrument is controlled through a personal computer, which interact with each one of electronic hardware components through the computer bus. An external 4-channel oscilloscope is also used to allow the 2-MGE user to visualize the voltage waveform and the reference signals that come from the PEMs but it is the only electronic hardware that is not connected to the computer bus and it has no contribution in the measurement.

### 5.4. Determination of the intensity with the Stokes-Mueller representation

The most convenient way to describe the polarization effects of optical elements of the 2-MGE is the Stokes-Mueller calculus introduced in section 1.4. We will use the Stokes-Mueller representation to describe the state of polarization of the light beam as it progressively traverses all the polarization-changing optical elements of a 2-MGE.

In a 2-MGE the PSG is composed of a polarizer-PEM pair. The Mueller matrix of a modulator $\mathbf{M}_{m}$ is

$$
\mathbf{M}_{m}=\left(\begin{array}{cccc}
1 & 0 & 0 & 0  \tag{5.2}\\
0 & 1 & 0 & 0 \\
0 & 0 & \cos (\delta) & \sin (\delta) \\
0 & 0 & -\sin (\delta) & \cos (\delta)
\end{array}\right)=\left(\begin{array}{cccc}
1 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 \\
0 & 0 & Y_{\delta} & X_{\delta} \\
0 & 0 & -X_{\delta} & Y_{\delta}
\end{array}\right),
$$

where $\delta$ is given by Eq. (5.1) and

$$
\begin{align*}
& Y_{\delta} \equiv \cos (\delta)  \tag{5.3a}\\
& X_{\delta} \equiv \sin (\delta) \tag{5.3b}
\end{align*}
$$

The Mueller matrix for a polarizer, $\mathbf{M}_{p}$, is

$$
\mathbf{M}_{p}=\left(\begin{array}{llll}
1 & 1 & 0 & 0  \tag{5.4}\\
1 & 1 & 0 & 0 \\
0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0
\end{array}\right)
$$

Considering that the input light is unpolarized, and that the azimuthal angles of the polarizer and PEM are $\theta_{p}$ and $\theta_{m}$, the Stokes vector for the light beam emerging from the PSG is

$$
\mathbf{S}_{P S G}=\mathbf{R}\left(-\theta_{m 0}\right) \mathbf{M}_{m 0} \mathbf{R}\left(\theta_{m 0}\right) \mathbf{R}\left(-\theta_{p 0}\right) \mathbf{M}_{p 0} \mathbf{R}\left(\theta_{p 0}\right)\left(\begin{array}{l}
1  \tag{5.5}\\
0 \\
0 \\
0
\end{array}\right)
$$

where $\mathbf{R}\left(\theta_{x}\right)$ is the rotation matrix defined in Eq. (1.27):

$$
\mathbf{R}\left(\theta_{x}\right)=\left(\begin{array}{cccc}
1 & 0 & 0 & 0  \tag{5.6}\\
0 & \cos \left(2 \theta_{x}\right) & \sin \left(2 \theta_{x}\right) & 0 \\
0 & -\sin \left(2 \theta_{x}\right) & \cos \left(2 \theta_{x}\right) & 0 \\
0 & 0 & 0 & 1
\end{array}\right)=\left(\begin{array}{cccc}
1 & 0 & 0 & 0 \\
0 & C_{x} & S_{x} & 0 \\
0 & -S_{x} & C_{x} & 0 \\
0 & 0 & 0 & 1
\end{array}\right)
$$

where the shorthand notation $\cos \left(2 \theta_{x}\right) \equiv C_{x}$ and $\sin \left(2 \theta_{x}\right) \equiv S_{x}$ has been used. We recall that $\theta$ is positive when the rotation is counterclockwise looking against the direction of propagation the light beam.

Multiplying out Eq. (5.5) the Stokes vector for the PSG is obtained:

$$
\mathbf{S}_{P S G}=\left(\begin{array}{c}
1  \tag{5.7}\\
C_{m 0} C_{b 0}+S_{m 0} S_{b 0} Y_{0 \delta} \\
S_{m 0} C_{b 0}-C_{m 0} S_{b 0} Y_{0 \delta} \\
S_{b 0} X_{0 \delta}
\end{array}\right)
$$

Where the definition $\theta_{b 0} \equiv \theta_{m 0}-\theta_{p 0}$ has been used. In a 2 -MGE the polarizerPEM pair is configured such that $\theta_{b 0}$ is $\pm 45^{\circ}$ (in our instrument we have chosen $45^{\circ}$ ). However, in practice, even using precise polarizer holders there is always some small angular misplacement that we represent by $\varepsilon_{b 0}$. Therefore, in our case, we have $\theta_{b 0}=45^{\circ}+\varepsilon_{b 0}$, which leads to $S_{b 0}=1$ and $C_{b 0}=-2 \varepsilon_{b 0}$, and Eq. (5.7) transforms into

$$
\mathbf{S}_{P S G}=\left(\begin{array}{c}
1  \tag{5.8}\\
-2 C_{m 0} \varepsilon_{b 0}+S_{m 0} Y_{0 \delta} \\
-\left(2 S_{m 0} \varepsilon_{b 0}+C_{m 0} Y_{0 \delta}\right) \\
X_{0 \delta}
\end{array}\right)
$$

In the case of the PSA we can perform an analog calculation. In the PSA light first passes through the PEM and then through the polarizer:

$$
\mathbf{S}_{P S A}^{T}=\left(\begin{array}{cccc}
1 & 0 & 0 & 0 \tag{5.9}
\end{array}\right) \mathbf{R}\left(-\theta_{p 1}\right) \mathbf{M}_{p 1} \mathbf{R}\left(\theta_{p 1}\right) \mathbf{R}\left(-\theta_{m 1}\right) \mathbf{M}_{m 1} \mathbf{R}\left(\theta_{m 1}\right)
$$

Again, taking the definition $\theta_{b 1} \equiv \theta_{m 1}-\theta_{p 1}$ and considering that we orientate the polarizer such that $\theta_{b 1}=45^{\circ}+\varepsilon_{b 1}$, the matrix multiplication yields the following transposed Stokes vector:

$$
\mathbf{S}_{P S A}^{T}=\left(\begin{array}{cccc}
1 & -2 \varepsilon_{b 1} \mathrm{C}_{m 1}+Y_{1 \delta} S_{m 1} & -2 \varepsilon_{b 1} S_{m 1}-Y_{1 \delta} C_{m 1} & -X_{1 \delta} \tag{5.10}
\end{array}\right) .
$$

The intensity of the time-dependent light beam for the 2-MGE is thus given by:

$$
\begin{equation*}
I(t)=\mathbf{S}_{P S A}^{T} \mathbf{M} \mathbf{S}_{P S G} \tag{5.11}
\end{equation*}
$$

where $\mathbf{M}$ is the transmission or reflection Mueller matrix of the sample given in Eq. (1.26). The result of this multiplication can be parameterized as follows:

$$
\begin{align*}
I(t)= & I_{d c}+I_{X 0} X_{0 \delta}+I_{Y 0} Y_{0 \delta}+I_{X 1} X_{1 \delta}+I_{X 0 X 1} X_{0 \delta} X_{1 \delta} \\
& +I_{X 0 Y 1} X_{0 \delta} Y_{1 \delta}+I_{Y 0 X 1} Y_{0 \delta} X_{1 \delta}+I_{Y 0 Y 1} Y_{0 \delta} Y_{1 \delta}, \tag{5.12}
\end{align*}
$$

where the time dependence of the intensity is embedded in the $X_{i \delta}$ and $Y_{i \delta}$ functions, which are given in Eqs. (5.3).

To simplify expressions we temporally assume that $\varepsilon_{b 0}=\varepsilon_{b 1}=0$ (they will be later reconsidered in the calibration section), we obtain that the eight coefficients in Eq. (5.12) are functions of the elements of the sample Mueller matrix and are given by, 1

$$
\begin{align*}
& I_{X 0}=m_{03} \\
& I_{Y 0}=-C_{m 0} m_{02}+S_{m 0} m_{01} \\
& I_{X 1}=-m_{30} \\
& I_{Y 1}=-C_{m 1} m_{20}+S_{m 1} m_{10}  \tag{5.13}\\
& I_{X 0 X 1}=-m_{33} \\
& I_{X 0 Y 1}=S_{m 1} m_{13}-C_{m 1} m_{23}, \\
& I_{Y 0 X 1}=-S_{m 0} m_{31}+C_{m 0} m_{32}, \\
& I_{Y 0 Y 1}=C_{m 0} C_{m 1} m_{22}-S_{m 0} C_{m 1} m_{21}-C_{m 0} S_{m 1} m_{12}+S_{m 0} S_{m 1} m_{11} .
\end{align*}
$$

These 8 terms are usually normalized by the $I_{d c}$ term to eliminate fluctuations of the incident light intensity, and the dependence of the sample transmissivity or reflectivity.

Using Eqs. (5.13) it is possible to define four cases, determined by the azimuthal angles of the PEMs to the sample ( $\theta_{m 0}$ and $\theta_{m 1}$ ) where different elements of the sample Mueller matrix are measured. Schematically, this cases can be represented
by:
case 1: $\theta_{m 0}= \pm 45^{\circ} ; \theta_{m 1}= \pm 45^{\circ}$

$$
\mathbf{M}=\left(\begin{array}{cccc}
1 & \sigma_{0 s} I_{Y 0} & \bullet & I_{X 0}  \tag{5.14a}\\
\sigma_{1 s} I_{Y 1} & \sigma_{0 s} \sigma_{1 s} I_{Y 0 Y 1} & \bullet & \sigma_{1 s} I_{X 0 Y 1} \\
\bullet & \bullet & \bullet & \bullet \\
-I_{X 1} & -\sigma_{0 s} I_{Y 0 X 1} & \bullet & -I_{X 0 X 1}
\end{array}\right)
$$

case 2: $\theta_{m 0}= \pm 45^{\circ} ; \theta_{m 1}=0^{\circ}, 90^{\circ}$

$$
\mathbf{M}=\left(\begin{array}{cccc}
1 & \sigma_{0 s} I_{Y 0} & \bullet & I_{X 0}  \tag{5.14b}\\
\bullet & \bullet & \bullet & \bullet \\
-\sigma_{1 c} I_{Y 1} & -\sigma_{0 s} \sigma_{1 c} I_{Y 0 Y 1} & \bullet & -\sigma_{1 c} I_{X 0 Y 1} \\
-I_{X 1} & -\sigma_{0 s} I_{Y 0 X 1} & \bullet & -I_{X 0 X 1}
\end{array}\right)
$$

case 3: $\theta_{m 0}=0^{\circ}, 90^{\circ} ; \theta_{m 1}= \pm 45^{\circ}$

$$
\mathbf{M}=\left(\begin{array}{cccc}
1 & \bullet & -\sigma_{0 c} I_{Y 0} & I_{X 0}  \tag{5.14c}\\
\sigma_{1 s} I_{Y 1} & \bullet & -\sigma_{0 c} \sigma_{1 s} I_{Y 0 Y 1} & \sigma_{1 s} I_{X 0 Y 1} \\
\bullet & \bullet & \bullet & \bullet \\
-I_{X 1} & \bullet & \sigma_{0 c} I_{Y 0 X 1} & -I_{X 0 X 1}
\end{array}\right)
$$

case 4: $\theta_{m 0}=0^{\circ}, 90^{\circ} ; \theta_{m 1}=0^{\circ}, 90^{\circ}$

$$
\mathbf{M}=\left(\begin{array}{cccc}
1 & \bullet & -\sigma_{0 c} I_{Y 0} & I_{X 0}  \tag{5.14d}\\
\bullet & \bullet & \bullet & \bullet \\
-\sigma_{1 c} I_{Y 1} & \bullet & \sigma_{0 c} \sigma_{1 c} I_{Y 0 Y 1} & -\sigma_{1 c} I_{X 0 Y 1} \\
-I_{X 1} & \bullet & \sigma_{0 c} I_{Y 0 X 1} & -I_{X 0 X 1}
\end{array}\right)
$$

where the signs terms are defined by

$$
\begin{array}{ll}
\sigma_{0 s}=\operatorname{sign}\left[\sin \left(2 \theta_{m 0}\right)\right], & \sigma_{1 s}=\operatorname{sign}\left[\sin \left(2 \theta_{m 1}\right)\right]  \tag{5.15}\\
\sigma_{0 c}=\operatorname{sign}\left[\cos \left(2 \theta_{m 0}\right)\right], & \sigma_{1 c}=\operatorname{sign}\left[\cos \left(2 \theta_{m 1}\right)\right]
\end{array}
$$

In every case the sample Mueller matrix elements that cannot be measured are signified by bullets $(\bullet)$. With these 4 configurations all the elements of the normalized sample Mueller matrix can be measured, and most of elements are more than once (for example elements $m_{03}, m_{30}$ and $m_{33}$ are measured in every case).

### 5.5. Data measurement

In the previous section it was demonstrated that the light intensity detected by the PMT is a function of time and can be represented by Eq. (5.12). The purpose
of this section is to show how the coefficients of that equation, namely $I_{X 0}, I_{Y 0}$, etc can be calculated from the experimental recorded $I(t)$

If the static retardation $\delta_{0}$ and $\delta_{1}$ of the PEMs, introduced in Eq. (5.1), are small $\left(\delta_{0}, \delta_{1} \ll 1\right)$ they can be treated to first order ${ }^{1}$ and Eqs. (5.3) can be rewritten as follows:

$$
\begin{align*}
& X_{0 \delta}=\sin \left[A_{0} \sin \left(\omega_{0} t+\phi_{0}\right)+\delta_{0}\right] \simeq X_{0}+\delta_{0} Y_{0},  \tag{5.16a}\\
& Y_{0 \delta}=\cos \left[A_{0} \sin \left(\omega_{0} t+\phi_{0}\right)+\delta_{0}\right] \simeq Y_{0}-\delta_{0} X_{0},  \tag{5.16b}\\
& X_{1 \delta}=\sin \left[A_{1} \sin \left(\omega_{1} t+\phi_{1}\right)+\delta_{1}\right] \simeq X_{1}+\delta_{1} Y_{1},  \tag{5.16c}\\
& Y_{1 \delta}=\cos \left[A_{1} \sin \left(\omega_{1} t+\phi_{1}\right)+\delta_{1}\right] \simeq Y_{1}-\delta_{1} X_{1}, \tag{5.16d}
\end{align*}
$$

where to short notation we have made the equivalences $X_{0} \equiv \sin \left[A_{0} \sin \left(\omega_{0} t+\phi_{0}\right)\right]$, $Y_{0} \equiv \cos \left[A_{1} \sin \left(\omega_{1} t+\phi_{1}\right)\right], X_{1} \equiv \sin \left[A_{1} \sin \left(\omega_{1} t+\phi_{1}\right)\right]$ and $Y_{1} \equiv \cos \left[A_{1} \cos \left(\omega_{1} t+\phi_{1}\right)\right]$. If the static retardations are temporally assumed to vanish $\left(\delta_{0}=\delta_{1}=0\right)$, Eq. (5.12) can be rewritten as

$$
\begin{align*}
I(t)= & I_{d c}+I_{X 0} X_{0}+I_{Y 0} Y_{0}+I_{X 1} X_{1}+I_{X 0 X 1} X_{0} X_{1} \\
& +I_{X 0 Y 1} X_{0} Y_{1}+I_{Y 0 X 1} Y_{0} X_{1}+I_{Y 0 Y 1} Y_{0} Y_{1}, \tag{5.17}
\end{align*}
$$

which can be also displayed in vector form:

$$
\begin{equation*}
I(t)=\mathbf{B}(t)^{T} \mathbf{I} \tag{5.18}
\end{equation*}
$$

where the basis vector $\mathbf{B}(t)$ and the intensity coefficient vector $\mathbf{I}$ are given by

$$
\begin{gather*}
\mathbf{B}(t)^{T}=\left(\begin{array}{ccccccccc}
1 & X_{0} & Y_{0} & X_{1} & Y_{1} & X_{0} X_{1} & X_{0} Y_{1} & Y_{0} X_{1} & Y_{0} Y_{1}
\end{array}\right),  \tag{5.19}\\
\mathbf{I}^{T}=\left(\begin{array}{ccccccccc}
I_{d c} & I_{X 0} & I_{Y 0} & I_{X 1} & I_{Y 1} & I_{X 0 X 1} & I_{X 0 Y 1} & I_{Y 0 X 1} & I_{Y 0 Y 1}
\end{array}\right) . \tag{5.20}
\end{gather*}
$$

The basis functions $X_{0}, Y_{0}, X_{1}$ and $Y_{1}$ are related to the common Fourier basis functions using an infinite series including integer Bessel functions:

$$
\begin{align*}
& X_{0}=2 \sum_{k=1}^{\infty} J_{2 k-1}\left(A_{0}\right) \sin \left[(2 k-1) \omega_{0} t\right],  \tag{5.21a}\\
& Y_{0}=J_{0}\left(A_{0}\right)+2 \sum_{k=1}^{\infty} J_{2 k}\left(A_{0}\right) \cos \left[2 k \omega_{0} t\right],  \tag{5.21b}\\
& X_{1}=2 \sum_{k=1}^{\infty} J_{2 k-1}\left(A_{1}\right) \sin \left[(2 k-1) \omega_{1} t\right], \tag{5.21c}
\end{align*}
$$

[^18]\[

$$
\begin{equation*}
Y_{1}=J_{0}\left(A_{1}\right)+2 \sum_{k=1}^{\infty} J_{2 k}\left(A_{1}\right) \cos \left[2 k \omega_{1} t\right] \tag{5.21d}
\end{equation*}
$$

\]

For many ellipsometric applications, $A_{i}$ is chosen to be 2.4048 radians, which gives $J_{0}(A)=0, J_{1}(A)=0.5192, J_{2}(A)=0.4318, J_{3}(A)=0.1990, J_{4}(A)=$ $0.0647, J_{5}(A)=0.0164$, etc. For this value of $A$, the Fourier expansion of the $X$ and $Y$ basis functions have no dc terms and the series converges rapidly. Also, for this value, the $J_{1}(A)$ and $J_{2}(A)$ are within $15 \%$ their maximum values

If Eqs. (5.21) are introduced in the time-dependent intensity of Eq. (5.17) we get the following expanded equation for the intensity of the light beam:

$$
\begin{align*}
& I(t)=I_{d c}+ \\
& +I_{X 0}\left[2 J_{1}\left(A_{0}\right) \sin \left(\omega_{0} t+\phi_{0}\right)+2 J_{3}\left(A_{0}\right) \sin \left(3 \omega_{0} t+3 \phi_{0}\right)+\ldots\right]+ \\
& +I_{Y 0}\left[J_{0}\left(A_{0}\right)+2 J_{2}\left(A_{0}\right) \cos \left(2 \omega_{0} t+2 \phi_{0}\right)+2 J_{4}\left(A_{0}\right) \cos \left(4 \omega_{0} t+4 \phi_{0}\right)+\ldots .\right]+ \\
& +I_{X 1}\left[2 J_{1}\left(A_{1}\right) \sin \left(\omega_{1} t+\phi_{1}\right)+2 J_{3}\left(A_{1}\right) \sin \left(3 \omega_{1} t+3 \phi_{1}\right)+\ldots\right]+ \\
& +I_{Y 1}\left[J_{0}\left(A_{1}\right)+2 J_{2}\left(A_{1}\right) \cos \left(2 \omega_{1} t+2 \phi_{1}\right)+2 J_{4}\left(A_{1}\right) \cos \left(4 \omega_{1} t+4 \phi_{1}\right)+\ldots .\right]+ \\
& +I_{X 0 X 1}\left[2 J_{1}\left(A_{0}\right) \sin \left(\omega_{0} t+\phi_{0}\right)+2 J_{3}\left(A_{0}\right) \sin \left(3 \omega_{0} t+3 \phi_{0}\right)+\ldots\right] \times \\
& \times\left[2 J_{1}\left(A_{1}\right) \sin \left(\omega_{1} t+\phi_{1}\right)+2 J_{3}\left(A_{1}\right) \sin \left(3 \omega_{1} t+3 \phi_{1}\right)+\ldots\right]+ \\
& +I_{X 0 Y 1}\left[2 J_{1}\left(A_{0}\right) \sin \left(\omega_{0} t+\phi_{0}\right)+2 J_{3}\left(A_{0}\right) \sin \left(3 \omega_{0} t+3 \phi_{0}\right)+\ldots\right] \times \\
& \times\left[J_{0}\left(A_{1}\right)+2 J_{2}\left(A_{1}\right) \cos \left(2 \omega_{1} t+2 \phi_{1}\right)+2 J_{4}\left(A_{1}\right) \cos \left(4 \omega_{1} t+4 \phi_{1}\right)+\ldots\right]+ \\
& +I_{Y 0 X 1}\left[J_{0}\left(A_{0}\right)+2 J_{2}\left(A_{0}\right) \cos \left(2 \omega_{0} t+2 \phi_{0}\right)+2 J_{4}\left(A_{0}\right) \cos \left(4 \omega_{0} t+4 \phi_{0}\right)+\ldots .\right] \times \\
& \times\left[2 J_{1}\left(A_{1}\right) \sin \left(\omega_{1} t+\phi_{1}\right)+2 J_{3}\left(A_{1}\right) \sin \left(3 \omega_{1} t+3 \phi_{1}\right)+\ldots\right]+ \\
& +I_{Y 0 Y 1}\left[J_{0}\left(A_{0}\right)+2 J_{2}\left(A_{0}\right) \cos \left(2 \omega_{0} t+2 \phi_{0}\right)+2 J_{4}\left(A_{0}\right) \cos \left(4 \omega_{0} t+4 \phi_{0}\right)+\ldots .\right] \times \\
& \times\left[J_{0}\left(A_{1}\right)+2 J_{2}\left(A_{1}\right) \cos \left(2 \omega_{1} t+2 \phi_{1}\right)+2 J_{4}\left(A_{1}\right) \cos \left(4 \omega_{1} t+4 \phi_{1}\right)+\ldots .\right] . \tag{5.22}
\end{align*}
$$

The first four basis functions of the time-dependent intensity of Eq. (5.22) are easily determined from the frequency components of the Fourier expressions given in Eqs. (5.21), but the last four are product functions of that equations. Thank to algebra properties of the sin and cos functions ${ }^{2}$ the product terms accompanying the factors $I_{X 0 X 1}, I_{X 0 Y 1}, I_{Y 0 X 1}$ and $I_{Y 0 Y 1}$ can be rewritten as frequency components at the sum and difference frequencies of the two modulators. Also, we can now assume that the amplitude of the Bessel function have been already adjusted so that $J_{0}\left(A_{0}\right)=J_{0}\left(A_{1}\right)=0$ (i.e. $A_{0}=A_{1}=2.4048 \mathrm{rad}$ ).

[^19]Chapter 5. Two-modulator generalized ellipsometer

The time-dependent intensity can be rewritten as:

$$
\begin{align*}
& I(t)=I_{d c}+ \\
&+I_{X 0} 2\left[J_{1}\left(A_{0}\right) \sin \left(\omega_{0} t+\phi_{0}\right)+J_{3}\left(A_{0}\right) \sin \left(3 \omega_{0} t+3 \phi_{0}\right)+\ldots\right]+ \\
&+I_{Y 0} 2\left[J_{2}\left(A_{0}\right) \cos \left(2 \omega_{0} t+2 \phi_{0}\right)+J_{4}\left(A_{0}\right) \cos \left(4 \omega_{0} t+4 \phi_{0}\right)+\ldots .\right]+ \\
&+I_{X 1} 2\left[J_{1}\left(A_{1}\right) \sin \left(\omega_{1} t+\phi_{1}\right)+J_{3}\left(A_{1}\right) \sin \left(3 \omega_{1} t+3 \phi_{1}\right)+\ldots\right]+ \\
&+I_{Y 1} 2\left[J_{2}\left(A_{1}\right) \cos \left(2 \omega_{1} t+2 \phi_{1}\right)+J_{4}\left(A_{1}\right) \cos \left(4 \omega_{1} t+4 \phi_{1}\right)+\ldots .\right]+ \\
&+I_{X 0 X 1} 2\left[J_{1}\left(A_{0}\right) J_{1}\left(A_{1}\right)\left[\cos \left[\left(\omega_{0}-\omega_{1}\right) t+\phi_{0}-\phi_{1}\right]-\cos \left[\left(\omega_{0}+\omega_{1}\right) t+\phi_{0}+\phi_{1}\right]\right]+\right. \\
& J_{1}\left(A_{0}\right) J_{3}\left(A_{1}\right)\left[\cos \left[\left(\omega_{0}-3 \omega_{1}\right) t+\phi_{0}-3 \phi_{1}\right]-\cos \left[\left(\omega_{0}+3 \omega_{1}\right) t+\phi_{0}+3 \phi_{1}\right]\right]+ \\
& J_{3}\left(A_{0}\right) J_{1}\left(A_{1}\right)\left[\cos \left[\left(3 \omega_{0}-\omega_{1}\right) t+3 \phi_{0}-\phi_{1}\right]-\cos \left[\left(3 \omega_{0}+\omega_{1}\right) t+3 \phi_{0}+\phi_{1}\right]\right]+ \\
&\left.J_{3}\left(A_{0}\right) J_{3}\left(A_{1}\right)\left[\cos \left[\left(3 \omega_{0}-3 \omega_{1}\right) t+3 \phi_{0}-3 \phi_{1}\right]-\cos \left[\left(3 \omega_{0}+3 \omega_{1}\right) t+3 \phi_{0}+3 \phi_{1}\right]\right]+\ldots\right]+ \\
&+I_{X 0 Y 1} 2\left[J_{1}\left(A_{0}\right) J_{2}\left(A_{1}\right)\left[\sin \left[\left(\omega_{0}-2 \omega_{1}\right) t+\phi_{0}-2 \phi_{1}\right]+\sin \left[\left(\omega_{0}+2 \omega_{1}\right) t+\phi_{0}+2 \phi_{1}\right]\right]+\right. \\
& J_{1}\left(A_{0}\right) J_{4}\left(A_{1}\right)\left[\sin \left[\left(\omega_{0}-4 \omega_{1}\right) t+\phi_{0}-4 \phi_{1}\right]+\sin \left[\left(\omega_{0}+4 \omega_{1}\right) t+\phi_{0}+4 \phi_{1}\right]\right]+ \\
& J_{3}\left(A_{0}\right) J_{2}\left(A_{1}\right)\left[\sin \left[\left(3 \omega_{0}-2 \omega_{1}\right) t+3 \phi_{0}-2 \phi_{1}\right]+\sin \left[\left(3 \omega_{0}+2 \omega_{1}\right) t+3 \phi_{0}+2 \phi_{1}\right]\right]+ \\
&\left.\quad J_{3}\left(A_{0}\right) J_{4}\left(A_{1}\right)\left[\sin \left[\left(3 \omega_{0}-4 \omega_{1}\right) t+3 \phi_{0}-4 \phi_{1}\right]+\sin \left[\left(3 \omega_{0}+4 \omega_{1}\right) t+3 \phi_{0}+4 \phi_{1}\right]\right]+\ldots\right]+ \\
&+I_{Y 0 X 1} 2\left[J_{1}\left(A_{0}\right) J_{2}\left(A_{1}\right)\left[\sin \left[\left(\omega_{1}-2 \omega_{0}\right) t+\phi_{1}-2 \phi_{0}\right]+\sin \left[\left(\omega_{1}+2 \omega_{0}\right) t+\phi_{1}+2 \phi_{0}\right]\right]+\right. \\
& J_{2}\left(A_{0}\right) J_{3}\left(A_{1}\right)\left[\sin \left[\left(3 \omega_{1}-2 \omega_{0}\right) t+3 \phi_{1}-2 \phi_{0}\right]+\sin \left[\left(3 \omega_{1}+2 \omega_{0}\right) t+3 \phi_{1}+2 \phi_{0}\right]\right]+ \\
& J_{4}\left(A_{0}\right) J_{1}\left(A_{1}\right)\left[\sin \left[\left(\omega_{1}-4 \omega_{0}\right) t+\phi_{1}-4 \phi_{0}\right]+\sin \left[\left(\omega_{1}+4 \omega_{0}\right) t+\phi_{1}+4 \phi_{0}\right]\right]+ \\
&\left.J_{4}\left(A_{0}\right) J_{3}\left(A_{1}\right)\left[\sin \left[\left(3 \omega_{1}-4 \omega_{0}\right) t+3 \phi_{1}-4 \phi_{0}\right]+\sin \left[\left(3 \omega_{1}+4 \omega_{0}\right) t+3 \phi_{1}+4 \phi_{0}\right]\right]+\ldots\right]+ \\
&+I_{Y 0 Y 1} 2\left[J_{2}\left(A_{0}\right) J_{2}\left(A_{1}\right)\left[\cos \left[\left(2 \omega_{0}-2 \omega_{1}\right) t+2 \phi_{0}-2 \phi_{1}\right]+\cos \left[\left(2 \omega_{0}+2 \omega_{1}\right) t+2 \phi_{0}+2 \phi_{1}\right]\right]+\right. \\
& J_{2}\left(A_{0}\right) J_{4}\left(A_{1}\right)\left[\cos \left[\left(2 \omega_{0}-4 \omega_{1}\right) t+2 \phi_{0}-4 \phi_{1}\right]+\cos \left[\left(2 \omega_{0}+4 \omega_{1}\right) t+2 \phi_{0}+4 \phi_{1}\right]\right]+ \\
& J_{4}\left(A_{0}\right) J_{2}\left(A_{1}\right)\left[\cos \left[\left(4 \omega_{0}-2 \omega_{1}\right) t+4 \phi_{0}-2 \phi_{1}\right]+\cos \left[\left(4 \omega_{0}+2 \omega_{1}\right) t+4 \phi_{0}+2 \phi_{1}\right]\right]+ \\
&\left.J_{4}\left(A_{0}\right) J_{4}\left(A_{1}\right)\left[\cos \left[\left(4 \omega_{0}-4 \omega_{1}\right) t+4 \phi_{0}-4 \phi_{1}\right]+\cos \left[\left(4 \omega_{0}+4 \omega_{1}\right) t+4 \phi_{0}+4 \phi_{1}\right]\right]+\ldots\right] . \tag{5.23}
\end{align*}
$$

Eq. (5.23) can be generalized to

$$
\begin{equation*}
I(t)=I_{d c}+2 \sum_{k=1}^{\infty} R_{k}\left[\alpha_{k} \cos \left(\Omega_{k} t\right)+\beta_{k} \sin \left(\Omega_{k} t\right)\right] \tag{5.24}
\end{equation*}
$$

where the firsts 31 values for $R_{k}$ and $\Omega_{k}$ are given in Table 5.3. There are an infinite number of terms to the sum in Eq. (5.24), but the higher-frequency terms (that are proportional to higher-index integer Bessel functions) are small if $A_{0}$ and $A_{1}$ are set to 2.4048 . However, note from Table 5.3, that still there table many components with frequencies lower than 240 kHz that have significant amplitudes. This makes the detected time-dependent waveform very complicated.

It is possible to write Eq.(5.24) in vector notation:

$$
\begin{equation*}
I(t)=\mathbf{C}^{T}(t) \mathbf{A} \tag{5.25}
\end{equation*}
$$

| $k$ | $\Omega_{k}$ | Frequency ${ }^{a}$ (kHz) | $R_{k}$ | $J_{x}\left(A_{0}\right) J_{y}\left(A_{1}\right)^{b}$ |
| :---: | :---: | :---: | :---: | :---: |
| 1 | $\omega_{0}-\omega_{1}$ | 10 | $I_{X 0 X_{1}} J_{1}\left(A_{0}\right) J_{1}\left(A_{1}\right)$ | 0.269 |
| 2 | $2\left(\omega_{0}-\omega_{1}\right)$ | 20 | $I_{Y O Y 1} J_{2}\left(A_{0}\right) J_{2}\left(A_{1}\right)$ | 0.187 |
| 3 | $4 \omega_{0}-3 \omega_{1}$ |  | $I_{Y 0 X 1} J_{4}\left(A_{0}\right) J_{3}\left(A_{1}\right)$ | 0.013 |
| 4 | $3 \omega_{0}-2 \omega_{1}$ | 30 | $I_{X 0 Y 1} J_{3}\left(A_{0}\right) J_{2}\left(A_{1}\right)$ | 0.084 |
| 5 | $3\left(\omega_{0}-\omega_{1}\right)$ |  | $I_{X 0 X 1} J_{3}\left(A_{0}\right) J_{3}\left(A_{1}\right)$ | 0.040 |
| 6 | $2 \omega_{0}-\omega_{1}$ | 40 | $I_{Y O X 1} J_{2}\left(A_{0}\right) J_{1}\left(A_{1}\right)$ | 0.224 |
| 7 | $4\left(\omega_{0}-\omega_{1}\right)$ |  | $I_{Y O Y 1} J_{4}\left(A_{0}\right) J_{4}\left(A_{1}\right)$ | 0.004 |
| 8 | $\omega_{0}$ | 50 | $I_{X 0} J_{1}\left(A_{0}\right)$ | 0.519 |
| 9 | $\omega_{1}$ | 60 | $I_{X 1} J_{1}\left(A_{1}\right)$ | 0.519 |
| 10 | $\omega_{0}-2 \omega_{1}$ | 70 | $I_{X 0 Y 1} J_{1}\left(A_{0}\right) J_{2}\left(A_{1}\right)$ | 0.224 |
| 11 | $2 \omega_{0}-3 \omega_{1}$ | 80 | $I_{Y 0 X 1} J_{2}\left(A_{0}\right) J_{3}\left(A_{1}\right)$ | 0.084 |
| 12 | $4 \omega_{0}-2 \omega_{1}$ |  | $I_{Y 0 Y 1} J_{4}\left(A_{0}\right) J_{2}\left(A_{1}\right)$ | 0.027 |
| 13 | $3 \omega_{0}-\omega_{1}$ | 90 | $I_{X 0 X 1} J_{3}\left(A_{0}\right) J_{1}\left(A_{1}\right)$ | 0.103 |
| 14 | $3 \omega_{0}-4 \omega_{1}$ |  | $I_{X O Y 1} J_{3}\left(A_{0}\right) J_{4}\left(A_{1}\right)$ | 0.013 |
| 15 | $2 \omega_{0}$ | 100 | $I_{Y 0} J_{2}\left(A_{0}\right)$ | 0.423 |
| 16 | $\omega_{0}+\omega_{1}$ | 110 | $I_{X 0 X 1} J_{1}\left(A_{0}\right) J_{1}\left(A_{1}\right)$ | 0.269 |
| 17 | $2 \omega_{1}$ | 120 | $I_{Y 1} J_{2}\left(A_{1}\right)$ | 0.423 |
| 18 | $\omega_{0}-3 \omega_{1}$ | 130 | $I_{X 0 X 1} J_{1}\left(A_{0}\right) J_{3}\left(A_{1}\right)$ | 0.103 |
| 19 | $4 \omega_{0}-\omega_{1}$ | 140 | $I_{Y 0 X 1} J_{4}\left(A_{0}\right) J_{1}\left(A_{1}\right)$ | 0.033 |
| 20 | $2 \omega_{0}-4 \omega_{1}$ |  | $I_{Y O Y 1} J_{2}\left(A_{0}\right) J_{4}\left(A_{1}\right)$ | 0.027 |
| 21 | $3 \omega_{0}$ | 150 | $I_{X 0} J_{3}\left(A_{0}\right)$ | 0.199 |
| 22 | $2 \omega_{0}+\omega_{1}$ | 160 | $I_{Y 0 X 1} J_{2}\left(A_{0}\right) J_{1}\left(A_{1}\right)$ | 0.224 |
| 23 | $\omega_{0}+2 \omega_{1}$ | 170 | $I_{X O Y 1} J_{1}\left(A_{0}\right) J_{2}\left(A_{1}\right)$ | 0.224 |
| 24 | $3 \omega_{1}$ | 180 | $I_{X 1} J_{3}\left(A_{1}\right)$ | 0.199 |
| 26 | $\omega_{0}-4 \omega_{1}$ | 190 | $I_{X O Y 1} J_{1}\left(A_{0}\right) J_{2}\left(A_{1}\right)$ | 0.033 |
| 27 | $4 \omega_{0}$ | 200 | $I_{Y 0} J_{4}\left(A_{0}\right)$ | 0.065 |
| 28 | $3 \omega_{0}+\omega_{1}$ | 210 | $I_{X 0 X 1} J_{3}\left(A_{0}\right) J_{1}\left(A_{1}\right)$ | 0.103 |
| 29 | $2\left(\omega_{0}+\omega_{1}\right)$ | 220 | $I_{Y O Y 1} J_{2}\left(A_{0}\right) J_{2}\left(A_{1}\right)$ | 0.187 |
| 30 | $\omega_{0}+3 \omega_{1}$ | 230 | $I_{X 0 X 1} J_{1}\left(A_{0}\right) J_{3}\left(A_{1}\right)$ | 0.103 |
| 31 | $4 \omega_{1}$ | 240 | $I_{Y 1} J_{4}\left(A_{1}\right)$ | 0.065 |

${ }^{a}$ Assuming $\omega_{0}=50 \mathrm{kHz}$ and $\omega_{1}=60 \mathrm{kHz}$.
${ }^{b}$ Assuming $A_{0}=A_{1}=2.4048$. The $x$ and $y$ indices refer to the integer Bessel function indices in the $R_{k}$ column.

Figure 5.3.: Firsts frequency components of the Fourier analysis of the TimeDependent Intensity vector. Table reproduced from [117].
with

$$
\begin{align*}
\mathbf{C}^{T}(t) & =\left(\begin{array}{llllllll}
1 & \cos \left(\Omega_{0} t\right) & \sin \left(\Omega_{0} t\right) & \cos \left(\Omega_{1} t\right) & \sin \left(\Omega_{1} t\right) & \cos \left(\Omega_{2} t\right) & \sin \left(\Omega_{2} t\right) & \ldots
\end{array}\right),  \tag{5.27}\\
\mathbf{A}^{T} & =\left(\begin{array}{llllllll}
I_{d c} & 2 R_{0} \alpha_{0} & 2 R_{0} \beta_{0} & 2 R_{1} \alpha_{1} & 2 R_{1} \beta_{1} & 2 R_{2} \alpha_{2} & 2 R_{2} \beta_{2} & \ldots
\end{array}\right) . \tag{5.26}
\end{align*}
$$

It is worth to remember that the objective here is to calculate the elements of the $\mathbf{A}$ vector (which are related to $I_{X 0}, I_{Y 0}$, etc) from the measured timedependent intensity $I(t)$. The elements of the $\mathbf{C}(t)$ vector form a natural basis set for the intensity. The following integration can be used to determine the elements of the $\mathbf{A}$ vector:

$$
\begin{equation*}
\int_{0}^{T} \mathbf{C}(t) I(t) d t=\mathbf{A} \int_{0}^{T} \mathbf{C}(t) \mathbf{C}^{T}(t) d t=\mathbf{A K} \tag{5.28}
\end{equation*}
$$

where $\mathbf{K}=\int_{0}^{T} \mathbf{C}(t) \mathbf{C}^{T}(t) d t$ and it is a diagonal matrix if the chosen period $T$ is a complete number of cycles for both modulators 0 and 1. The vector $\mathbf{A}$ becomes:

$$
\begin{equation*}
A=\mathbf{K}^{-1} \int_{0}^{T} \mathbf{C}(t) I(t) d t \tag{5.29}
\end{equation*}
$$

In practice, since our digitized $I(t)$ consist of a long vector of point, we need to use a discrete version of Eq. (5.29) in order to determine $I_{X 0}, I_{Y 0}, I_{X 1}, I_{Y 1}, I_{X 0 X 1}$, $I_{X 0 Y 1}, I_{Y 0 X 1}$ and $I_{Y 0 Y 1}$. Given a digitized waveform I consisting in a vector of $T$ elements, we can generate a basis matrix $\mathbf{C}$ that in general contains at least eight different frequencies $\Omega_{k}$ (one for each one of the coefficients to determine). This matrix $\mathbf{C}$ has dimension ( $0: 2 \mathrm{~N}, 0: \mathrm{T}-1$ ) where N is the number of frequencies incorporated into the analysis. The elements of this basis matrix are constructed as

$$
\begin{align*}
& C_{0, i}=1.0 \\
& C_{2 k, i}=\cos \left(\Omega_{k} i \Delta t\right)  \tag{5.30}\\
& C_{2 k-1, i}=\sin \left(\Omega_{k} i \Delta t\right), \quad k=1, \ldots, N ; \quad i=0, \ldots, T-1,
\end{align*}
$$

where $\Delta t$ is the inverse of the sampling rate used for digitizing the waveform and $T \Delta t$ is the total time captured by the waveform digitizer.

The correction matrix $\mathbf{K}$ of dimension ( $0: 2 \mathrm{~N}, 0: 2 \mathrm{~N}$ ) is calculated from $\mathbf{C}$

$$
\begin{equation*}
\mathbf{K}=\mathbf{C C}^{T}, \tag{5.31}
\end{equation*}
$$

and in analogy with Eq. (5.29) the vector $\mathbf{A}$, of dimension ( $0: 2 \mathrm{~N}$ ), is calculated

$$
\begin{equation*}
\mathbf{A}=\mathbf{K}^{-1} \mathbf{C I} \tag{5.32}
\end{equation*}
$$

whose elements are $A_{0}=I_{d c} ; \quad A_{2 k}=\alpha_{k} ; \quad A_{2 k-1}=\beta_{k} k=1, \ldots N$.
According to Eq. (5.24) the magnitude of each frequency component is given by $\left(\alpha_{k}^{2}+\beta_{k}^{2}\right)^{1 / 2}$ and the sign of the component must be determined from the
measured phase $\phi_{m}$ and the relative modulator phases $\phi_{0}$ and $\phi_{1}$. The measured phase can be calculated as $\phi_{m}=\operatorname{atan} \beta_{k} / \alpha_{k}$.

We usually use 12 frequencies for this Fourier integration $(N=12)$ : $\omega_{0}$ to determine $I_{X 0}, 2 \omega_{0}$ to determine $I_{Y 0}, \omega_{1}$ to determine $I_{X 1}, 2 \omega_{1}$ to determine $I_{Y 1}$, $\omega_{0}+\omega_{1}$ and $\left|\omega_{0}-\omega_{1}\right|$ to independently determine $I_{X 0 X 1}, \omega_{0}+2 \omega_{1}$ and $\left|\omega_{0}-2 \omega_{1}\right|$ to independently determine $I_{X 0 Y 1}, 2 \omega_{0}+\omega_{1}$ and $\left|2 \omega_{0}-\omega_{1}\right|$ to independently determine $I_{Y 0 X 1}$, and $2 \omega_{0}+2 \omega_{1}$ and $\left|2 \omega_{0}-2 \omega_{1}\right|$ to independently determine $I_{Y 0 Y 1}$. The value of each of the eight coefficients is determined with the following equations:

$$
\begin{align*}
& I_{X 0}=\operatorname{sign}\left[\sin \left(\phi_{m}+\phi_{0}\right)\right] \frac{\left(\alpha^{2}+\beta^{2}\right)^{1 / 2}}{2 J_{1}\left(A_{0}\right)}{ }_{\Omega=\omega_{0}},  \tag{5.33a}\\
& I_{Y 0}=\operatorname{sign}\left[\cos \left(\phi_{m}+\phi_{0}\right)\right] \frac{\left(\alpha^{2}+\beta^{2}\right)^{1 / 2}}{2 J_{2}\left(A_{0}\right)}{ }_{\Omega=2 \omega_{0}},  \tag{5.33b}\\
& I_{X 1}=\operatorname{sign}\left[\cos \left(\phi_{m}+\phi_{1}\right)\right] \frac{\left(\alpha^{2}+\beta^{2}\right)^{1 / 2}}{2 J_{1}\left(A_{1}\right)} \quad \Omega=\omega_{1},  \tag{5.33c}\\
& I_{Y 1}=\operatorname{sign}\left[\cos \left(\phi_{m}+\phi_{1}\right)\right] \frac{\left(\alpha^{2}+\beta^{2}\right)^{1 / 2}}{2 J_{2}\left(A_{1}\right)} \quad \Omega=2 \omega_{1},  \tag{5.33d}\\
& I_{X 0 X 1}=\operatorname{sign}\left[\sin \left(\phi_{m}+\phi_{0}+\phi_{1}\right)\right] \frac{\left(\alpha^{2}+\beta^{2}\right)^{1 / 2}}{4 J_{1}\left(A_{0}\right) J_{1}\left(A_{1}\right)}{ }_{\Omega=\omega_{0}+\omega_{1}}+ \\
& +\operatorname{sign}\left[\sin \left(\phi_{m}+\phi_{0}-\phi_{1}\right)\right] \frac{\left(\alpha^{2}+\beta^{2}\right)^{1 / 2}}{4 J_{1}\left(A_{0}\right) J_{1}\left(A_{1}\right)_{\Omega=\omega_{0}-\omega_{1}}},  \tag{5.33e}\\
& I_{X 0 Y 1}=\operatorname{sign}\left[\sin \left(\phi_{m}+\phi_{0}+2 \phi_{1}\right)\right] \frac{\left(\alpha^{2}+\beta^{2}\right)^{1 / 2}}{4 J_{1}\left(A_{0}\right) J_{2}\left(A_{1}\right)}{ }_{\Omega=\omega_{0}+2 \omega_{1}}+ \\
& +\operatorname{sign}\left[\sin \left(\phi_{m}+\phi_{0}-2 \phi_{1}\right)\right] \frac{\left(\alpha^{2}+\beta^{2}\right)^{1 / 2}}{4 J_{1}\left(A_{0}\right) J_{2}\left(A_{1}\right)_{\Omega=\omega_{0}-2 \omega_{1}}},  \tag{5.33f}\\
& I_{Y 0 X 1}=\operatorname{sign}\left[\sin \left(\phi_{m}+2 \phi_{0}+\phi_{1}\right)\right] \frac{\left(\alpha^{2}+\beta^{2}\right)^{1 / 2}}{4 J_{2}\left(A_{0}\right) J_{1}\left(A_{1}\right)}{ }_{\Omega=2 \omega_{0}+\omega_{1}}+ \\
& +\operatorname{sign}\left[\sin \left(\phi_{m}+2 \phi_{0}-\phi_{1}\right)\right] \frac{\left(\alpha^{2}+\beta^{2}\right)^{1 / 2}}{4 J_{2}\left(A_{0}\right) J_{1}\left(A_{1}\right)}{\Omega=2 \omega_{0}-\omega_{1}},  \tag{5.33~g}\\
& I_{Y 0 Y 1}=\operatorname{sign}\left[\cos \left(\phi_{m}+2 \phi_{0}+2 \phi_{1}\right)\right] \frac{\left(\alpha^{2}+\beta^{2}\right)^{1 / 2}}{4 J_{2}\left(A_{0}\right) J_{2}\left(A_{1}\right)}{ }_{\Omega=2 \omega_{0}+2 \omega_{1}}+ \\
& +\operatorname{sign}\left[\sin \left(\phi_{m}+2 \phi_{0}-2 \phi_{1}\right)\right] \frac{\left(\alpha^{2}+\beta^{2}\right)^{1 / 2}}{4 J_{2}\left(A_{0}\right) J_{2}\left(A_{1}\right)}{ }_{\Omega=2 \omega_{0}-2 \omega_{1}}, \tag{5.33h}
\end{align*}
$$

Note that for the determination $I_{X 0 X 1}, I_{X 0 Y 1}, I_{Y 0 X 1}$ and $I_{Y 0 Y 1}$ we average the measurements of two different frequency components. Moreover we check that
the values obtained of each one of this two frequencies matches (within the error limits) the value of its companion. This ensures that the acquisition program can precisely resolve every frequency component.

According to the description of the triggering circuit given in section 5.3.2, we start the digitization whenever the reference signals of the modulators are in phase. As a result it would be reasonable for the reader to think that thank to the triggering circuit modulator phases $\phi_{0}$ and $\phi_{1}$ will close to zero. However this not the case, because PEMs controller puts out a constant phase that adds to the own optical phase of the modulator acting like an offset. This translates in no vanishing, but constant, values of $\phi_{0}$ and $\phi_{1} . \phi_{m}, \phi_{0}$ and $\phi_{1}$ are only used to measure the sign of the component of the I vector, and they need not be determined very accurately. The values of $\phi_{0}$ and $\phi_{1}$ are not same for every PEM, and need to be determined. To do so one can use sample with known optical behavior, i. e. samples for which we know the sign of the components of I vector, by recursively adjusting the values $\phi_{0}$ and $\phi_{1}$ till we get a correct sign for all the monitored frequency components.

### 5.6. Calibration

In other to measure the eight parameters $I_{X 0}, I_{Y 0}$, etc accurately the 2 -MGE needs to be calibrated. The calibration allows to determine all the characteristics of the PSG and the PSA, namely the angular orientation of each polarizer with respect to the longitudinal axis of its corresponding PEM $\left(\theta_{b 0}\right.$ and $\left.\theta_{b 1}\right)$, the control voltage needed to generate the PEM amplitudes such $A_{0}=A_{1}=2.4048$ (for this amplitude $J_{0}\left(A_{0}\right)=J_{0}\left(A_{1}\right)=0$ ), and the static strain-induced retardations $\delta_{0}$ and $\delta_{1}$ of each one of the PEMs.

The calibration is done in the straight-through configuration, without any sample. In this configuration the detected light intensity is given by:

$$
\begin{equation*}
I(t)=\mathbf{S}_{P S A}^{T} \mathbf{S}_{P S G} \tag{5.34}
\end{equation*}
$$

The result of this multiplication can be written using the basis vector notation as:

$$
\begin{equation*}
I(t)=\mathbf{B}_{\delta}^{T} \mathbf{I} \tag{5.35}
\end{equation*}
$$

where B

$$
\begin{gather*}
\mathbf{B}_{\delta}(t)^{T}=\left(\begin{array}{llllllllll}
1 & X_{0 \delta} & Y_{0 \delta} & X_{1 \delta} & Y_{1 \delta} & X_{0 \delta} X_{1 \delta} & X_{0 \delta} Y_{1 \delta} & Y_{0 \delta} X_{1 \delta} & Y_{0 \delta} Y_{1 \delta}
\end{array}\right),  \tag{5.36}\\
\mathbf{I}^{T}=\left(\begin{array}{llllllll}
1 & 0 & -2 C_{m 1} S_{m 0} \varepsilon_{b 1} & 0 & -2 C_{m 1} C_{m 0} \varepsilon_{b 0} & -1 & 0 & 0 \\
S_{m 1} S_{m 0}
\end{array}\right) . \tag{5.37}
\end{gather*}
$$

Note that the subscript $\delta$ in Eq. (5.35) denotes that the static retardation of the modulators is being considered (compare for example to Eq. (5.19)).

According to Eqs. (5.16) the static strain treated at first order can be included as follows:

$$
\left(\begin{array}{c}
1  \tag{5.38}\\
X_{0 \delta} \\
Y_{0 \delta} \\
X_{1 \delta} \\
Y_{1 \delta} \\
X_{0 \delta} X_{1 \delta} \\
X_{0 \delta} Y_{1 \delta} \\
Y_{0 \delta} X_{1 \delta} \\
Y_{0 \delta} Y_{1 \delta}
\end{array}\right) \simeq\left(\begin{array}{ccccccccc}
1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 1 & \delta_{0} & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & -\delta_{0} & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 1 & \delta_{1} & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & -\delta_{1} & 1 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 1 & \delta_{1} & \delta_{0} & 0 \\
0 & 0 & 0 & 0 & 0 & -\delta_{1} & 1 & 0 & \delta_{0} \\
0 & 0 & 0 & 0 & 0 & -\delta_{0} & 0 & 1 & \delta_{1} \\
0 & 0 & 0 & 0 & 0 & 0 & -\delta_{0} & -\delta_{0} & 1
\end{array}\right)\left(\begin{array}{c}
1 \\
X_{0} \\
Y_{0} \\
X_{1} \\
Y_{1} \\
X_{0} X_{1} \\
X_{0} Y_{1} \\
Y_{0} X_{1} \\
Y_{0} Y_{1}
\end{array}\right) .
$$

In shorthand notation is

$$
\begin{equation*}
\mathbf{B}_{\delta} \simeq \delta \mathrm{B} \tag{5.39}
\end{equation*}
$$

To include the effects of a non-zero $J_{0}\left(A_{0}\right)$ or $J_{0}\left(A_{1}\right)$ the basis elements $Y_{0}$ and $Y_{1}$ should be respectively replaced by $J_{0}\left(A_{0}\right)+Y_{0}$ and $J_{0}\left(A_{1}\right)+Y 1$ (see Eqs. ( 5.21 b ) and (5.21d)), so that a better approximation to (5.39) can be written as

$$
\mathbf{B}_{\delta} \cong \boldsymbol{\delta}\left(\begin{array}{c}
1  \tag{5.40}\\
X_{0} \\
J_{0}\left(A_{0}\right)+Y_{0} \\
X_{1} \\
J_{0}\left(A_{1}\right)+Y_{1} \\
X_{0} X_{1} \\
X_{0} J_{0}\left(A_{1}\right)+X_{0} Y_{1} \\
X_{1} J_{0}\left(A_{0}\right)+X_{1} Y_{0} \\
J_{0}\left(A_{0}\right) Y 1+Y_{0} J_{0}\left(A_{1}\right)+Y_{0} Y_{1}
\end{array}\right)
$$

that written as a matrix multiplication transforms into

$$
\begin{equation*}
\mathbf{B}_{\delta} \cong \delta \mathbf{J}_{0} \mathbf{B} \tag{5.41}
\end{equation*}
$$

where the $\mathbf{J}_{0}$ matrix is:

$$
\mathbf{J}_{0}=\left(\begin{array}{ccccccccc}
1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0  \tag{5.42}\\
0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
J_{0}\left(A_{0}\right) & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 \\
J_{0}\left(A_{1}\right) & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\
0 & J_{0}\left(A_{1}\right) & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\
0 & 0 & 0 & J_{0}\left(A_{0}\right) & 0 & 0 & 0 & 1 & 0 \\
0 & 0 & J_{0}\left(A_{1}\right) & 0 & J_{0}\left(A_{0}\right) & 0 & 0 & 0 & 1
\end{array}\right) .
$$

Chapter 5. Two-modulator generalized ellipsometer

Substituting Eq. (5.41) in Eq. (5.35) we get:

$$
\begin{equation*}
I(t)=\left[\boldsymbol{\delta} \mathbf{J}_{0} \mathbf{B}\right]^{T} \mathbf{I}=\mathbf{B}^{T} \mathbf{J}_{0}^{T} \boldsymbol{\delta}^{T} \mathbf{I} \tag{5.43}
\end{equation*}
$$

and from this it is deduced

$$
\begin{equation*}
\mathbf{I}_{\text {calibration }}=\mathbf{J}_{0}^{T} \boldsymbol{\delta}^{T} \mathbf{I} . \tag{5.44}
\end{equation*}
$$

Once the static strain and the possible non zero values for $J_{0}\left(A_{0}\right)$ and $J_{0}\left(A_{1}\right)$ are considered, the intensity coefficients initially given in Eq.(5.37) become more complicate:

$$
\begin{align*}
I_{X 0} & =0,  \tag{5.45a}\\
I_{Y 0} & =J_{0}\left(A_{1}\right) \cos \left(2 \theta_{m}\right)-2 \sin \left(2 \theta_{m}\right) \varepsilon_{b 1},  \tag{5.45b}\\
I_{X 1} & =0,  \tag{5.45c}\\
I_{Y 1} & =J_{0}\left(A_{0}\right) \cos \left(2 \theta_{m}\right)+2 \sin \left(2 \theta_{m}\right) \varepsilon_{b 0},  \tag{5.45d}\\
I_{X 0 X 1} & =-1,  \tag{5.45e}\\
I_{X 0 Y 1} & =-\delta_{1}-\delta_{0} \cos \left(2 \theta_{m}\right),  \tag{5.45f}\\
I_{Y 0 X 1} & =-\delta_{0}-\delta_{1} \cos \left(2 \theta_{m}\right),  \tag{5.45~g}\\
I_{Y 0 Y 1} & =\cos 2 \theta_{m}, \tag{5.45h}
\end{align*}
$$

where $\theta_{m}$ is the angle of the PSG with respect to the PSA $\left(\theta_{m}=\theta_{m 0}-\theta_{m 1}\right.$. With the exception of $I_{X 0 X 1}$ and $I_{Y 0 Y 1}$ if the system is well calibrated, the other coefficients of Eq.(5.45) are close to zero.

Equations (5.45) are used to identify the three basic calibration parameters of each PEM: the error in the polarizer angle with respect to the $45^{\circ} \varepsilon_{b i}$, the Bessel angle $A_{i}$ and the static strain $\delta_{i}(\lambda)$. If we first align parallel the PEMs $\left[\cos \left(2 \theta_{m}\right)=1\right.$ and $\left.\sin \left(2 \theta_{m}\right)=1\right]$ we get that $I_{Y 0}$ measures $J_{0}\left(A_{1}\right)$, while $I_{Y 1}$ measures $J_{0}\left(A_{0}\right)$. The objective of this first step of the calibration is to determine the control voltage as a function of the wavelength $V_{m i}(\lambda)$ required for setting $A_{i}=2.4048$ and thus to make $I_{Y 0}=I_{Y 1}=0$. Assuming that we know with anticipation the approximate voltage that makes $A \sim 2.4$, the automatic procedure to precisely find the desired control voltage is based on measuring at different wavelengths the values of $I_{Y 0}$ and $I_{Y 1}$ while applying different control voltages slightly deviated from our initial guess. For each wavelength the dispersion of $I_{Y 0}$ and $I_{Y 1}$ values with respect to the applied voltage is linearly fit, and fit results we get for every wavelength the voltage $V_{m 1}$ and $V_{m 0}$ that respectively makes $J_{0}\left(A_{0}\right)$ and $J_{0}\left(A_{1}\right)$ zero.

The second step of the calibration takes place with the PEMs aligned at $\pm 45^{\circ}$ with respect to one another $\left[\cos \left(2 \theta_{m}\right)=0\right]$. In this configuration the coefficients $I_{Y 0}$ measure $\varepsilon_{b 1}$ and $\varepsilon_{b 0}, I_{X 0 Y 1}$ and $I_{Y 0 X 1}$ measure $\delta_{1}$ and $\delta_{0}$, and we can use the coefficient $I_{Y 0 Y 1}$ to get a precise adjustment in the azimuthal alignment of
the two PEMs (this coefficient should vanish if the are orientated at $\pm 45^{\circ}$ ). The overall gain can be measured by coefficient $I_{X 0 X 1}$ for any value of $\theta_{m}$. Note that while $\delta_{i}$ and $V_{m i}$ are expected to change with $\lambda$, the overall gain and $\varepsilon_{b i}$ will approximately remain constant and we can average values taken at different wavelengths.

The relation between the Bessel angle $A$ and the modulator control voltage $V_{m}$ exhibits dispersion with photon energy or wavelength $\lambda$ and is of the form [129]:

$$
\begin{equation*}
V_{m i}(\lambda)=\frac{K_{i} A_{i} \lambda}{\sum_{k=0}^{\infty} \alpha_{k i} / \lambda^{2 k}} \tag{5.46}
\end{equation*}
$$

where $K_{i}$ is a constant. The static strain is a similar function of wavelength due to the dispersion of fused quartz and is given by [129]

$$
\begin{equation*}
\delta_{i}(\lambda)=\lambda \sum_{k=0}^{\infty} \beta_{k i} / \lambda^{2 k} \tag{5.47}
\end{equation*}
$$

The spectroscopic values of $V_{m i}$ and $\delta_{i}$ obtained in the calibration are respectively fitted using Eqs. (5.46) and (5.47). Fig 5.4 shows the control voltage calibrations $V_{m 0}$ and $V_{m 1}$ and the measured static strains $\delta_{0}$ and $\delta_{1}$ as a function of wavelength. The fitting have been done using Eqs (5.46) and (5.47) using four parameters for $V_{m i}$ and three parameters for $\delta_{i}$.

In addition to the overall, frequency-independent gain factor that is determined through the calibration. In practice there are additional particular gain factors for every frequency component that plays a role in the determination of the intensity vector. They are all caused by the limited bandwidth of the preamplifier for high frequencies and can be easily calculated if the performance characteristics (bandwidth and roll-off) preamplifier are known. In case they are not known they can be deduced by comparing different frequency components related to the same coefficient of the I vector (for example both the $\sim 20 \mathrm{kHz}$ and $\sim 220 \mathrm{kHz}$ components should give the same coefficient $I_{Y 0 Y 1}$. For our pre-amplifier the particular attenuation factors of each frequency component are calculated with the following roll-off curve:

$$
\begin{equation*}
A(\nu)=\frac{1}{\left(\nu / \nu_{0}\right)^{2}+1} \tag{5.48}
\end{equation*}
$$

where $\nu_{0}=356 \mathrm{kHz}$. The attenuation grow as we approach to the higher frequencies of interest [for example $A(20 k H z)=0.997$ and $A(220 k H z)=0.724$ ].

These calibrations completely characterize the PSG and PSA and need to be performed periodically to make measurements with high accuracy. The calibration process is automatized, so that a complete spectroscopic calibration routine


Figure 5.4.: Calibration data for the modulators used in the 2-MGE. Top panel, the modulator control voltage $V_{m 0}$ and $V_{m 1}$ required to give $A_{0}=A_{1}=2.4048$ where the solid lines represent the fits to the data using Eq. (5.46). Middle panel, static strain of each modulator as a function of wavelength, where the solid curves indicate the fits to the data by using Eq. (5.47). Bottom panel, the overall gain factor for measured at two different orientations of the PSG and the PSA.
that uses two different orientations of the PSG with respect to the PSA (a parallel or perpendicular orientation to find $V_{m i}$ and a $\pm 45^{\circ}$ orientation to find $\delta_{i}$ and $\left.\varepsilon_{b i}\right)$ is done in about 10 min . Fig. 5.4 shows the typical graph result of a calibration routine; in addition to the parameters shown in this figure, the values of $\varepsilon_{b i}$ are saved and residual errors in the alignment of the PEMs (through the parameter $\theta_{m}$ are also detected).

Calibration in a 2 -MGE is actually easier and probably more reliable than in a ellipsometers using a single photoelastic modulator because, as shown in Eqs.(5.45) the measured intensity coefficients, for a particular orientation of the PSG and the PSA, depend only on one calibration parameter. For example, for aligned PEMs $I_{Y 0}$ and $I_{Y} 1$ directly are $J_{0}\left(A_{1}\right)$ and $J_{0}\left(A_{0}\right)$, and thus we have a very sensitive method for measuring the deviation of $A_{i}$ from 2.4048 that is not available in single PEM instruments.

### 5.6.1. Using the calibration results into the measurement

To simplify the algebra in several calculations of section 5.5 it was considered that $\delta_{0}=\delta_{1}=\varepsilon_{b 0}=\varepsilon_{b 1}=J_{0}\left(A_{0}\right)=J_{0}\left(A_{1}\right)=0$. The calibration made in this section has allowed us to minimize $\varepsilon_{b 0}, \varepsilon_{b 1}, J_{0}\left(A_{0}\right)$ and $J_{0}\left(A_{1}\right)$ and it is safe to assure that are very small. However, although we have spectroscopically determined the static retardation $\delta_{0}$ and $\delta_{1}$ of the modulators we cannot reduce it.

Considering a general sample the detected intensity is given by Eq. (5.11), that without considering the static retardation of the modulators leads to the coefficients of the $\mathbf{I}$ vector given in Eq. (5.13), if we want to include the effect of a nonzero $\delta_{0}$ and $\delta_{1}$ we can use the strain matrix $\boldsymbol{\delta}$ :

$$
\begin{equation*}
I_{\delta-\text { corrected }}(t)=\boldsymbol{\delta}^{T} \mathbf{I} \tag{5.49}
\end{equation*}
$$

Chapter 5. Two-modulator generalized ellipsometer
so that Eqs. (5.13) now transform into:

$$
\begin{align*}
I_{d c}= & 1 \\
I_{X 0}= & m_{03}+\left(m_{02} C_{m 0}-m_{01} S_{m 0}\right) \delta_{0} \\
I_{Y 0}= & -C_{m 0} m_{02}+S_{m 0} m_{01}, \\
I_{X 1}= & -m_{30}+\left(m_{20} C_{m 1}-m_{10} S_{m 1}\right) \delta_{1}, \\
I_{Y 1}= & -C_{m 1} m_{20}+S_{m 1} m_{10}, \\
I_{X 0 X 1}= & -m_{33}+\left(m_{31} S_{m 0}-m_{32} C_{m 0}\right) \delta_{0}+\left(-m_{13} S_{m 1}+m_{23} C_{m 1}\right) \delta_{1}, \\
I_{X 0 Y 1}= & S_{m 1} m_{13}-C_{m 1} m_{23} \\
& +\left(m_{12} S_{m 1} C_{m 0}-m_{22} C_{m 1} C_{m 0}+m_{21} C_{m 1} S_{m 0}-m_{11} S_{m 1} S_{m 0}\right) \delta_{0}-\delta_{1} m_{33}, \\
I_{Y 0 X 1}= & -S_{m 0} m_{31}+C_{m 0} m_{32} \\
& +\left(m_{12} S_{m 1} C_{m 0}-m_{22} C_{m 1} C_{m 0}+m_{21} C_{m 1} S_{m 0}-m_{11} S_{m 1} S_{m 0}\right) \delta_{0}-\delta_{0} m_{33}, \\
I_{Y 0 Y 1}= & C_{m 0} C_{m 1} m_{22}-S_{m 0} C_{m 1} m_{21}-C_{m 0} S_{m 1} m_{12}+S_{m 0} S_{m 1} m_{11} \\
& \left.+\left(m_{32} C_{m 0}-m_{31} S_{m 0}\right) \delta_{1}+\left(m_{13} S_{m 1}-m_{23} C_{m 1}\right) \delta_{0}\right) . \tag{5.50}
\end{align*}
$$

## Chapter 6.

## Quartz assisted two-modulator generalized ellipsometer

The idea of complementing the 2-MGE setup by adding a quartz plate in each arm of the instrument started with the purpose of increasing the number of elements of the sample Mueller matrix that could be measured without changing the azimuthal orientations of the PSG and the PSA, which was required to do space resolved measurements.

### 6.1. Introduction

Due to the imperfect alignment of optical components that form the PSG and the PSA the light beam is not fully perpendicular to their surfaces. There will be a small, but usually non negligible, angle of incidence for light upcoming these optical elements (e.g. the polarizers and the PEMs). For example if we consider, as a rough estimation, that the PSG is a single optical element of $\mathrm{SiO}_{2}$ with a thickness of 50 mm (the thickness of the PEM's optical bar plus the polarizer is usually above this thicknesss), for a misalignment of $1^{\circ}$ the Snell's law indicates that the exiting beam of the PSG will be laterally displaced by about 0.6 mm from the entering beam. This sitution is referred as beam translation, and has been already described as a problem for imaging ellipsometers involving moving parts [132]. If the azimuthal rotation of the entire PSG is considered one might also take into account that the axis of rotation might not be perfectly parallel to the light beam nor with the surfaces normal. All this leads to deviations in the light beam that vary with the azimuthal position of the PSG that translate in small, but noticeable, differences in the spotted area of the sample. We will refer to this problem as beam translation. For the PSA the problem may be less dramatic if the elements that collect the light after the PSA cover are not too small, but still can cause that only fractions of the light probe would be analyzed.

In general these drawbacks do not represent a major problem if a complete measurement of the Mueller matrix is not required (then only one measurement configuration is used) or if the sample surface is homogeneous enough in reasonably big domains as often happens with thin films. However it is not compatible with samples that require measurements of the complete Mueller matrix with a
certain lateral resolution. Over the years, several different instruments have been proposed or build to measure complete Mueller matrices [73, 133-140]; however, Mueller polarimeters are not instruments of mainstream use and their presence in laboratories is still uncommon. In these instruments polarization modulation and analysis is commonly achieved used rotating compensators or liquid crystals with variable retardation. To perform space resolved measurements, instruments based on liquid crystals are adequate because they do not have moving parts and are compatible with CCD detectors [141]. However, the optical quality of liquid crystal based modulators is lower than than the one offered by PEMs [142], and, in general, these instruments do not reach the level accuracy and the sensitivity of PEM-based polarimeters.

In this chapter we describe an alternative technique to measure transmission or reflection Mueller matrices with a 2-MGE based on the use of thin crystal slabs with rotatory power (e.g. plates of quartz) as a substitute of the mechanical rotator to orientate the PSG and the PSA. This technique will be referred in this work as quartz-assisted two-modulator generalized ellipsometry, and will be described in detail in this chapter. Nevertheless, most of the contents of the previous chapter also apply to this technique so for a better comprehension this chapter should be studied independently.

The use of crystals with rotatory power for laser applications is quite common and, in fact, one can find available in the marked crystals ${ }^{1}$ cut such to provide a specific amount of rotation for characteristic laser wavelengths. However this optical rotation of the polarization has not been exploited in polarimeters or ellipsometers. The closer reference we have found to the subject is a paper by Azzam [143] which describes, but only from a theoretical point of view, the simulation of mechanical rotation by optical rotation and the possible application to a Fourier photopolarimeter (an instrument to measure the Stokes vector of a radiation). To our knowledge we present here the first experimental realization of a polarimeter that makes use of optical rotation as a substitutive of mechanical rotation.

Dealing with the same problem some previous experiments were done with another approach. This previous method consisted in adding one waveplate, tuned to the wavelength of measurement, in both arms of the instrument [144]. In that case the objective of the introduction of waveplates was not to measure more elements of the Mueller matrix (still 8 elements were measured), but instead they were used to strategically select the particular 8 Mueller matrix that we wanted to measure instead of being restricted to one of the schemes presented in Eqs. (5.14). Some results were obtained by this method and they will be reviewed in section 9.2, however here we will not give more details about this method as we consider that the one we present in this section deals with the same

[^20]experimental problem and is fairly superior. Quartz plates have the advantage that their induced polarization rotation is usually less sensitive to the alignment that retardation of the waveplates.

### 6.2. Description of the quartz-assisted 2-MGE

The quartz-assisted 2-MGE is similar to the 2-MGE described in the previous chapter. The basic difference consist on the addition of two quartz plates in the setup: one between the PSG and the sample (quartz 0) and the other between the sample and the PSA (quartz 1), so that light passes through a plate just before and after the sample. We use polished z-cut quartz $\left(\alpha-\mathrm{SiO}_{2}\right)$ plates, which means that their optical axis is perpendicular to their surface. For this type of cut and for light at normal incidence an uniaxial birefringent material such as quartz appears to be isotropic and the only optical effect which introduces is the optical rotation [145]. In our system quartz plates are mounted in two filter wheels (Thorlabs FW102B) conveniently placed after the PSG and before the PSA, as it is shown in Fig. 6.1. In practice the filter wheels are used to reach any of this 4 possible configurations which are essential for the method of measurement:

- Configuration I. Air / Sample / Air
- Configuration II. Quartz 0 / Sample / Air
- Configuration III. Air / Sample / Quartz 1
- Configuration IV. Quartz 0 / Sample / Quartz 1

Filter wheels allow a fast switch between the different configurations. In contrast to the $2-\mathrm{MGE}$ described in the previous chapter, here the mechanical rotators devoted to change the azimuthal positions of the PSG and the PSA are not needed for the measurement of the Mueller matrix. However, if they are present, they are still useful for the PEM alignment and to calibrate the static strain of the PEMs as it will be shown later.

An important attribute of the quartz plates is their thickness. They must be thin enough to not introduce any appreciable beam translation even if they are not perfectly aligned perpendicular to the beam. For a quartz plate of 1 mm with an angle of incidence of $1^{\circ}$ the beam translation of the beam caused by the quartz element is $12 \mu \mathrm{~m}$. Worthy to note that individual small quartz plates are easier to align with respect to the light beam that the entire PSG or PSA so angles of incidence $<1^{\circ}$ are not difficult to obtain.

The main difficulty arising from the addition of the quartz plates starts from the fact that the measured Mueller matrix contains contributions from ALL the optical elements between the PSG and the PSA. In this case, this includes the
sample (the element to be characterized) and the two quartz plates. Therefore for a successful measurement of the sample's Mueller matrix the contribution of the 2 quartz plates must be carefully determined.

One of the main applications of the quartz-assisted 2 -MGE is to perform Mueller matrix space resolved measurements of the sample. In our instrument the optical resolution of the space resolved measurements is mainly given by the diameter of the probe beam that illuminate the sample. Today's ellipsometers use either collimated or focused beams. In the case of collimated beams they diameters of $\sim 5 \mathrm{~mm}$ are usual, while for the case of focused beams the diameters can be below $100 \mu \mathrm{~m}$.

The focusing elements can be placed inside the space delimited by the PSG and PSA or can be external, i. e. elements placed in front of the PSG and behind the PSA. In the first case it is possible to use lenses with short focal lengths that facilitate a high local resolution but increases the polarization aberration. In this configuration the birefringence of the lenses affects the measurements, although in some cases it can be corrected [146]. When the focusing elements are external the anisotropies of these elements do not influence the ellipsometric result. This setups usually require comparatively long focal lengths of the optical elements, so that a beam with small divergence hits the sample and the lateral resolution is reduced. Our instrument uses external focusing elements (JovinYvon) that allows three beam spot sizes being $\sim 1 \mathrm{~mm}$ the bigger one and $\sim 90 \mu \mathrm{~m}$ the smaller one. For this system reducing the spot results in less light intensity reaching the detector, so the resultant signal-to-noise ratio is worse when compared to measurements with a bigger spot.

The sample is placed in a automated precision $\mathrm{x}-\mathrm{y}$ stage (see Fig. 6.1) that allows to map the sample by taking successive measurements of the Mueller matrix while the sample is moved to different positions. For spatially resolved measurements the instrument operates at a single wavelength, that is resolved by the monochromator. To increase the signal-to-noise ratio in these kind of measurements the monochromator could be substituted by an interference filter and the strong emission lines of a mercury lamp could be used as a light source.


Figure 6.1.: Photo of the 2-MGE built during this thesis. From left to right the picture first shows an optical fiber attached to the the focusing optics module, and it is followed by the first polarizer-PEM pair mounted in a mechanical rotation stage (PEMs are easily identifiable by the blue cables). The sample appears in the center of the photo, mounted in $x-y$ automatic translation stage. The elements just before and after the sample are filter wheels containing z-cut quartz rotators. After the second filter wheels comes the last PEM-polarizer pair, also attached to a mechanical rotation stage. Finally light is collected by another optical fiber that guides it to the monochromator.

Chapter 6. Quartz assisted two-modulator generalized ellipsometer

### 6.3. Effect of quartz plates in the measurement

A quartz plate can be described in terms of the Mueller matrix as:

$$
\mathbf{M}_{Q}=\left(\begin{array}{cccc}
1 & 0 & 0 & 0  \tag{6.1}\\
0 & \cos \mathrm{~T}_{R}+\mathrm{LB}^{2} \alpha & \mathrm{LBLB}^{\prime} \alpha+\mathrm{CB} \beta & -\mathrm{LBCB} \alpha+\mathrm{LB}^{\prime} \beta \\
0 & \mathrm{LBLB}^{\prime} \alpha-\mathrm{CB} \beta & \cos \mathrm{~T}_{R}+\mathrm{LB}^{\prime 2} \alpha & -\mathrm{LB}^{\prime} \mathrm{CB} \alpha-\mathrm{LB} \beta \\
0 & -\mathrm{LBCB} \alpha-\mathrm{LB}^{\prime} \beta & -\mathrm{LB}^{\prime} \mathrm{CB} \alpha+\mathrm{LB} \beta & \cos \mathrm{~T}_{R}+\mathrm{CB}^{2} \alpha
\end{array}\right)
$$

with $\alpha=\left(1-\cos \mathrm{T}_{R}\right) / \mathrm{T}_{R}^{2}, \beta=\sin \mathrm{T}_{R} / \mathrm{T}_{R}$ and $\mathrm{T}_{R}=\sqrt{\mathrm{LB}^{2}+\mathrm{LB}^{\prime 2}+\mathrm{CB}^{2}}$. For a z-cut plate, if the incident beam is parallel to the optical axis it behaves as an isotropic sample an no longer exhibits linear birefringence $\left(\mathrm{LB}=\mathrm{LB}^{\prime}=0\right)$. But, in general, it may be not possible to have a perfect alignment between the optic axis and the light beam so we assume that they are small ( $\mathrm{LB}, L B^{\prime} \ll$ $\mathrm{CB})$. Concerning these small linear birefringences their effect is assimilable to the case of ellipsometry with strained windows (see for example [146]). For a good orientation of the plates is easy to keep LB and $\mathrm{LB}^{\prime}$ below the $\sim 0.02$ rad range, and so we can consider them only to first order; then $T_{R}=C B$ and Eq.(6.1) becomes:

$$
\mathbf{M}_{Q i}=\left(\begin{array}{cccc}
1 & 0 & 0 & 0  \tag{6.2}\\
0 & \cos \mathrm{CB}_{i} & \sin \mathrm{CB}_{i} & A_{3 i} \\
0 & -\sin \mathrm{CB}_{i} & \cos \mathrm{CB}_{i} & A_{4 i} \\
0 & A_{1 i} & A_{2 i} & 1
\end{array}\right)
$$

where

$$
\begin{align*}
& A_{1 i}=-\mathrm{LB}_{i} \frac{\left(1-\cos \mathrm{CB}_{i}\right)}{\mathrm{CB}_{i}}-\mathrm{LB}_{i}^{\prime} \frac{\sin \mathrm{CB}_{i}}{\mathrm{CB}_{i}}  \tag{6.3a}\\
& A_{2 i}=-\mathrm{LB}_{i}^{\prime} \frac{\left(1-\cos \mathrm{CB}_{i}\right)}{\mathrm{CB}_{i}}+\mathrm{LB}_{i} \frac{\sin \mathrm{CB}_{i}}{\mathrm{CB}_{i}}  \tag{6.3b}\\
& A_{3 i}=-\mathrm{LB}_{i} \frac{\left(1-\cos \mathrm{CB}_{i}\right)}{\mathrm{CB}_{i}}+\mathrm{LB}_{i}^{\prime} \frac{\sin \mathrm{CB}_{i}}{\mathrm{CB}_{i}}  \tag{6.3c}\\
& A_{4 i}=-\mathrm{LB}_{i}^{\prime} \frac{\left(1-\cos \mathrm{CB}_{i}\right)}{\mathrm{CB}_{i}}-\mathrm{LB}_{i} \frac{\sin \mathrm{CB}_{i}}{\mathrm{CB}_{i}} \tag{6.3d}
\end{align*}
$$

with $i=0,1$ to refer to the first of the second quartz plates. $A_{1 i}, A_{2 i}, A_{3 i}, A_{4 i}$ are not independent, for example:

$$
\begin{align*}
& A_{1 i}=A_{4 i} \frac{1}{\sin \mathrm{CB}_{i}}+A_{2 i} \frac{\cos \mathrm{CB}_{i}}{\sin \mathrm{CB}_{i}}  \tag{6.4a}\\
& A_{3 i}=-A_{4 i} \frac{\cos \mathrm{CB}_{i}}{\sin \mathrm{CB}_{i}}-A_{2 i} \frac{1}{\sin \mathrm{CB}_{i}} \tag{6.4b}
\end{align*}
$$

The CB (which is twice the optical rotation, usually denoted by symbol $\rho$ ) of each plate can be adjusted to the following dispersion relation [20,147]:

$$
\begin{equation*}
\mathrm{CB}_{i}=\frac{m_{i} \lambda^{2}}{\left(\lambda^{2}-\lambda_{0, i}^{2}\right)^{2}} \tag{6.5}
\end{equation*}
$$

where $m_{i}$ and $\lambda_{0, i}$ are parameters that are determined through the calibration.
For each of the four configurations described in the previous section, the Mueller matrix $\mathbf{M}_{T}$ that describes the optical elements between the PSG and the PSA is:

- Configuration I. $\mathbf{M}_{T}=\mathbf{M}$
- Configuration II. $\mathbf{M}_{T}=\mathbf{M M}_{Q 0}$
- Configuration III. $\mathbf{M}_{T}=\mathbf{M}_{Q 1} \mathbf{M}$
- Configuration IV. $\mathbf{M}_{T}=\mathbf{M}_{Q 1} \mathbf{M M}_{Q 0}$
where $\mathbf{M}$ is given in Eq. (1.26) and $\mathbf{M}_{Q i}$ is given in Eq. (6.1) The results of these matrix multiplication are too large to be explicitly written here but are important to study the effect introduced by the quartz plates on the measurement. Therefore we will only write down the elements of the matrix $\mathbf{M}_{T}$ that are of interest.

Our measurement procedure uses as the fixed position of the PSG and PSA the angles: $\theta_{m 0}=0^{\circ} \theta_{m 1}=0^{\circ}$, that corresponds to the situation in which the modulation axis of the two modulators are parallel to the primary reference frame. All the calculations of this section are referred to this orientation. The choose of this particular orientation among all the cases given in section 5.4 is partially arbitrary: any of the cases in which the modulation axis of the PEMs are aligned parallel or perpendicular would be equally suitable. However, cases where the PEMs are aligned at $\pm 45^{\circ}$ with respect one to another would be less suitable because this positions are incompatible with a complete calibration.

According to Eq. (5.14a) in configuration $I$ the sample's Mueller matrix elements $m_{02}, m_{03}, m_{20}, m_{22}, m_{23}, m_{30}, m_{32}$ and $m_{33}$ can be measured. Configuration II permits to measure $m_{01}, m_{21}$ and $m_{31}$. With configuration III $m_{10}, m_{12}$

Chapter 6. Quartz assisted two-modulator generalized ellipsometer
and $m_{13}$ are determined. The remaining element, $m_{11}$, is accessible in configuration $I V$. In summary:

$$
\mathbf{M}=\left(\begin{array}{cccc}
1 & m_{01, I I} & m_{02, I} & m_{03, I}  \tag{6.6}\\
m_{10, I I I} & m_{11, I V} & m_{12, I I I} & m_{13, I I I} \\
m_{20, I} & m_{21, I I} & m_{22, I} & m_{23, I} \\
m_{30, I} & m_{31, I I} & m_{32, I} & m_{33, I}
\end{array}\right)
$$

The matrix elements correspondent to configuration $I$ are directly found from the coefficients of the intensity vector as shown in Eq. (5.14a). The rest of elements are found using the following equations:

$$
\begin{align*}
m_{01, I I}= & -\left[I_{Y 0, I I}+A_{2,0} m_{03, I}+m_{02, I} \cos \left(\mathrm{CB}_{0}\right)\right] \sin ^{-1}\left(\mathrm{CB}_{0}\right)  \tag{6.7a}\\
m_{21, I I}= & {\left[I_{Y 0 Y 1, I I}-A_{2,0} m_{23, I}-m_{22, I} \cos \left(\mathrm{CB}_{0}\right)\right] \sin ^{-1}\left(\mathrm{CB}_{0}\right) }  \tag{6.7b}\\
m_{31, I I}= & {\left[I_{Y 0 X 1, I I}-A_{2,0} m_{33, I}-m_{32, I} \cos \left(\mathrm{CB}_{0}\right)\right] \sin ^{-1}\left(\mathrm{CB}_{0}\right) }  \tag{6.7c}\\
m_{10, I I I}= & {\left[I_{Y 1, I I I}+A_{4,1} m_{30, I}+m_{20, I} \cos \left(\mathrm{CB}_{1}\right)\right] \sin ^{-1}\left(\mathrm{CB}_{1}\right) }  \tag{6.7d}\\
m_{12, I I I}= & {\left[-I_{Y 0 Y 1, I I I}+A_{4,1} m_{32, I}+m_{22, I} \cos \left(\mathrm{CB}_{1}\right)\right] \sin ^{-1}\left(\mathrm{CB}_{1}\right) }  \tag{6.7e}\\
m_{13, I I I}= & {\left[I_{X 0 Y 1, I I I}+A_{4,1} m_{33, I}+m_{23, I} \cos \left(\mathrm{CB}_{1}\right)\right] \sin ^{-1}\left(\mathrm{CB}_{1}\right) }  \tag{6.7f}\\
m_{11, I V}= & -\left[I_{Y 0 Y 1, I V}-A_{2,0}\left[m_{23, I} \cos \left(\mathrm{CB}_{1}\right)-m_{13, I I I} \sin \mathrm{CB}_{1}\right]\right.  \tag{6.7g}\\
& -\cos \left(\mathrm{CB}_{0}\right)\left[m_{22, I} \cos \left(\mathrm{CB}_{1}\right)-m_{12, I I I} \sin \left(\mathrm{CB}_{1}\right)+A_{4,1} m_{32, I}\right]  \tag{6.7h}\\
& \left.-\sin \left(\mathrm{CB}_{0}\right)\left[m_{21, I I} \cos \left(\mathrm{CB}_{1}\right)+A_{4,1} m_{31, I I}\right]\right] \sin ^{-1}\left(\mathrm{CB}_{0}\right) \sin ^{-1}\left(\mathrm{CB}_{1}\right) \tag{6.7i}
\end{align*}
$$

$\mathrm{CB}_{0}, \mathrm{CB}_{1}, A_{2,0}$ and $A_{4,1}$ are the wavelength-dependent parameters associated to the quartz plates that play a role in measurement of the complete Mueller matrix for a ( 0,0 ) [(PSG, PSA)] orientation of the PSG and PSA. Their values are determined during the calibration.

### 6.4. Calibration

The calibration in the quartz-assisted 2-MGE involves the determination of all the characteristics associated to a standard 2-MGE (most notably the Bessel voltages to make $J_{0}\left(A_{0}\right)=J_{0}\left(A_{1}\right)=0$ and the modulator's static strain $\delta_{0}$ and $\delta_{1}$, see section 5.6) and also the characteristic parameters of the quartz plates $\mathrm{CB}_{0}, \mathrm{CB}_{1}, A_{2,0}$ and $A_{4,1}$.

Table 6.1 shows the light intensity coefficients when the ellipsometer is aligned in the straight-through configuration without sample and with the modulation axis of the two modulators aligned parallel $\left(\theta_{m 0}=\theta_{m 1}=0^{\circ}\right)$ for the 4 configurations described in section 6.2. Configuration I permits to measure values of
$J_{0}\left(A_{1}\right)$ and $J_{0}\left(A_{0}\right)$ and it used to determine the control voltage required for setting $A_{i}=2.4048$ (for more details see section 5.6). Once this voltage has been found the configurations II and III are set to respectively determine $\cos \left(\mathrm{CB}_{0}\right)$ and $\cos \left(\mathrm{CB}_{1}\right)$ through the coefficient $I_{Y 0 Y 1}$. From this measurement the values of $\sin \left(\mathrm{CB}_{0}\right)$ and $\sin \left(\mathrm{CB}_{1}\right)$ can also be inferred.

The linear retardation parameters of the quartz plates that interfere the measurement (i.e. $A_{2,0}$ and $A_{4,1}$ ), appear mixed with the static retardation of the modulators $\delta_{0}$ and $\delta_{1}$ in coefficients $I_{X 0 Y 1}$ and $I_{Y 0 X 1}$, and are not directly distinguishable. The easier way to separate and quantify the linear retardation contributions of the modulators and the quartz plates is to align the PEMs at $\pm 45^{\circ}$ with respect to one another. This is the case described in section 5.6, where $I_{X 0 Y 1} I_{Y 0 X 1}$ become direct measurement of $\delta_{0}$ and $\delta_{1}$. Once they are known the coefficients $I_{X 0 Y 1} I_{Y 0 X 1}$ of configuration II allow to measure $A_{2,0}$ and $A_{4,0}$, while the same parameters in configuration III measure $A_{2,1}$ and $A_{4,1}$ (note that $\cos \left(\mathrm{CB}_{0}\right)$ and $\cos \left(\mathrm{CB}_{1}\right)$ are also known $)$.

Table 6.1.: Intensity coefficients for different calibration configurations. It is assumed that configuration $I$ is used first to find the Bessel voltages that satisfy $J_{0}\left(A_{1}\right)=J_{0}\left(A_{0}\right)=0$, and these voltages are used for the rest of the configuration (this is the reason why $J_{0}\left(A_{1}\right)$ and $J_{0}\left(A_{0}\right)$ do not appear in the other configurations).

| Coeff. | Config. I | Config. II | Config. III | Config. IV |
| :---: | :---: | :---: | :---: | :---: |
| $I_{X 0}$ | 0 | 0 | 0 | 0 |
| $I_{Y 0}$ | $J_{0}\left(A_{1}\right)$ | $2 \varepsilon_{b 1} \sin \left(\mathrm{CB}_{0}\right)$ | $2 \varepsilon_{b 1} \sin \left(\mathrm{CB}_{1}\right)$ | $2 \varepsilon_{b 1} \Xi_{1}$ |
| $I_{X 1}$ | 0 | 0 | 0 | 0 |
| $I_{Y 1}$ | $J_{0}\left(A_{0}\right)$ | $-2 \varepsilon_{b 0} \sin \left(\mathrm{CB}_{0}\right)$ | $-2 \varepsilon_{b 0} \sin \left(\mathrm{CB}_{1}\right)$ | $2 \varepsilon_{b 1} \Xi_{2}$ |
| $I_{X 0 X 1}$ | -1 | -1 | -1 | -1 |
| $I_{X 0 Y 1}$ | $-\delta_{1}-\delta_{0}$ | $-D_{0}-\delta_{1}-\delta_{0} \cos \left(\mathrm{CB}_{0}\right)$ | $-D_{1}-\delta_{1}-\delta_{0} \cos \left(\mathrm{CB}_{1}\right)$ | $-\nu-\delta_{1}-\delta_{0} H$ |
| $I_{Y 0 X 1}$ | $-\delta_{0}-\delta_{1}$ | $B_{0}-\delta_{0}-\delta_{1} \cos \left(\mathrm{CB}_{0}\right)$ | $B_{1}-\delta_{0}-\delta_{1} \cos \left(\mathrm{CB}_{1}\right)$ | $\mu-\delta_{0}-\delta_{1} H$ |
| $I_{Y 0 Y 1}$ | 1 | $\cos \left(\mathrm{CB}_{0}\right)$ | $\cos \left(\mathrm{CB}_{1}\right)$ | $\Xi_{3}$ |

In which the following identifications have been made:

$$
\begin{align*}
\Xi_{1} & =\cos \left(\mathrm{CB}_{0}\right) \sin \left(\mathrm{CB}_{1}\right)+\cos \left(\mathrm{CB}_{1}\right) \sin \left(\mathrm{CB}_{0}\right),  \tag{6.8a}\\
\Xi_{2} & =-\cos \left(\mathrm{CB}_{1}\right) \sin \left(\mathrm{CB}_{0}\right)-\cos \left(\mathrm{CB}_{0}\right) \sin \left(\mathrm{CB}_{1}\right),  \tag{6.8b}\\
\Xi_{3} & =\cos \left(\mathrm{CB}_{0}\right) \cos \left(\mathrm{CB}_{1}\right)-\sin \left(\mathrm{CB}_{0}\right) \sin \left(\mathrm{CB}_{1}\right),  \tag{6.8c}\\
\mu & =A_{2,0}+A_{1,1} \sin \left(\mathrm{CB}_{0}\right)+A_{2,1} \cos \left(\mathrm{CB}_{0}\right),  \tag{6.8d}\\
\nu & =A_{4,1}-A_{3,0} \sin \left(\mathrm{CB}_{1}\right)+A_{4,0} \cos \left(\mathrm{CB}_{1}\right) . \tag{6.8e}
\end{align*}
$$

Note that with this approach the configuration IV is not required for the calibration. An alternative calibration procedure would be desirable in case of an
instrument without azimuthal rotators attached to the PSG and PSA where changing the orientation of the PEMs to determine $\delta_{0}$ and $\delta_{1}$ might be not possible. In that case it is possible to construct the following system of equations:

$$
\begin{align*}
& -\delta_{1}-\delta_{0}=I_{X 0 Y 1, a} \\
& -A_{4,0}-\delta_{1}-\delta_{0} \cos \left(\mathrm{CB}_{0}\right)=I_{X 0 Y 1, b}, \\
& A_{2,1}-\delta_{0}-\delta_{1} \cos \left(\mathrm{CB}_{0}\right)=I_{Y 0 X 1, b} \\
& -A_{4,1}-\delta_{1}-\delta_{0} \cos \left(\mathrm{CB}_{1}\right)=I_{X 0 Y 1, c},  \tag{6.9}\\
& A_{2,1}-\delta_{0}-\delta_{1} \cos \left(\mathrm{CB}_{1}\right)=I_{Y 0 X 1, c} \\
& -\nu-\delta_{1}-\delta_{0} H=I_{X 0 Y 1, d} \\
& \mu-\delta_{0}-\delta_{1} H=I_{Y 0 X 1, d}
\end{align*}
$$

where, using Eqs. (6.4) :

$$
\begin{gather*}
\nu=A_{4,1} D_{1}+\left[A_{2,0} \frac{\cos \mathrm{CB}_{0}}{\sin \mathrm{CB}_{0}}+A_{4,0} \frac{1}{\sin \mathrm{CB}_{0}}\right] \sin \left(\mathrm{CB}_{1}\right)+A_{4,0} \cos \left(\mathrm{CB}_{1}\right),  \tag{6.10}\\
\mu=A_{2,0} B_{0}+\left[A_{2,1} \frac{1}{\sin \mathrm{CB}_{1}}+A_{4,1} \frac{\cos \mathrm{CB}_{1}}{\sin \mathrm{CB}_{1}}\right] \sin \left(\mathrm{CB}_{0}\right)+A_{2,1} \cos \left(\mathrm{CB}_{0}\right) .
\end{gather*}
$$

This is a linear system of equations with 6 unknown parameters: $\delta_{0}, \delta_{1}, A_{2,0}, A_{2,1}$, $A_{4,0}$ and $A_{4,1}$. Therefore the values of this parameters can be approximated by using a multiple linear regression analysis.

Figure 6.2 shows the measured cosines of the CB of the plates as a function of the wavelength as well as fit obtained with the dispersion of CB given in Eq. (6.5). The small differences in the measured values of $\cos C B$ for quartz 0 and quartz 1 are due a different thickness of the two plates ( $1.06 \pm 0.01 \mathrm{~mm}$ vs $1.02 \pm 0.01$ $\mathrm{mm})$. The bottom shows the measured linear birefringences $A_{2,0}, A_{2,1}, A_{4,0}$ and $A_{4,1}$ of the quartz plates as a function of the wavelength as well as the fit to the raw data by using Eq. (5.47). Note that the dispersion relation used for this fit is the same that was used to fit the static strain of the modulators.

To complete the calibration, coefficients $I_{Y 0}$ and $I_{Y 0}$ of configurations II or III can be used to measure the angles of the polarizer with respect to the PEMs ( $\varepsilon_{b 0}$ $\varepsilon_{b 1}$, defined in section 5.4) and the overall gain is measured by $I_{X 0 X 1}$.

### 6.5. Measurement schemes

The quartz-assisted 2-MGE offers two different modes of measurement: spectroscopic and space-resolved. The instrument can either work in any of these two modes, without significant instrumental changes between both. The characteristics of all the elements involved in the instrument were given in section 5.3.

### 6.5.1. Spectroscopic mode

The wavelength span of the instrument is $\sim 210-880$. The limitation in the UV is determined by the low output of Xe arc lamp in the UV and the absorp-


Figure 6.2.: Spectroscopic calibration of the 1 mm quartz plates. Top panel shows the $\cos \left(\mathrm{CB}_{i}\right)$, where the solid curves indicate the fits to the data using Eq. (6.5). Bottom panel shows the parameters $A_{2,0}, A_{2,1}, A_{4,0}$ and $A_{4,1}$ that are related to the small linear birefringence of the plates. The solid curves are the fits to the data using the model in Eq. (5.47).
tion properties of the optical materials involved (polarizers, optical fibers, etc.), whereas the infrared limitation is due to the cutoff of the PMT. The use of the quartz assisted technique does not narrows the wavelength span of the instrument since quartz is a high transparent material at these wavelengths; however, and due to the dispersive nature of the optical effects in quartz, a spectroscopic calibration of the quartz plates such the one presented in the previous section is needed.

According to Eqs. (6.7) the values of $\sin \mathrm{CB}_{0}, \sin \mathrm{CB}_{1}$, that appear in the denominator, are needed to calculate all the element of the Mueller matrix of the sample. Thus, it is evident that there will be an algebraically unfavorable situation whenever $\sin \mathrm{CB}_{i} \sim 0$. The value of CB increases when going to shorter wavelengths and is proportional to the thickness of the plate. For quartz, at the
wavelengths of measurement the case $\mathrm{CB}_{i} \simeq 0$ would only be a problem for extremely thin slabs, but the case $\mathrm{CB}_{i} \simeq 180^{\circ}$ occurs at $\sim 316 \mathrm{~nm}$ using 1 mm plates. In this situation a precise measurement of some elements of the Mueller matrix becomes impossible. On the contrary the situation CB $\sim 90^{\circ}$ (note that this corresponds to an optical rotation of $\sim 45^{\circ}$, that is a typical angle of rotation for instruments with mechanical rotators) is the more favorable for measurements.


Figure 6.3.: Evolution of $\sin \mathrm{CB}$ for two quartz plates with thicknesses 1 mm and 0.5 mm . Area marked in grey exemplifies wavelengths unsuitable for the measurement with the 1 mm plate but adequate for the 0.5 mm plate. Area in light red is inadequate for both plates.

Fig. 6.3 depicts the sin CB calculated using the fitting parameters found in the calibration for two quartz plates of different thickness. It can be seen that for each plate there is a zone suitable for spectroscopic measurements (for the 1 mm plate goes roughly goes from 340 nm to 850 nm ) because sin CB varies smoothly and does not take small values. For lower wavelength in the UV the CB quickly increases and the function sin CB starts oscillating; this zone is not suitable for spectroscopic measurements with elements of this thickness and requires thinner plates. The figure shows that with the 0.5 mm plate measurements down to 260 nm are possible but it offers a worse response (lower values of $\sin \mathrm{CB}$ ) for high wavelengths. To perform spectroscopic measurements in the far UV region a very thin quartz plate (e.g. 0.2 mm ) would be needed.

Fig. 6.4 shows the spectroscopic Mueller matrix measurement of a zero-order


Figure 6.4.: Spectroscopic Mueller matrix of a zero-order quarter wave plate designed for 532 nm . Different colours denote measurements taken at different configurations. For a quarter wave plate Mueller matrix element $m_{44}$ corresponds to the cosinus of the retardance.
plate designed to give a $\lambda / 4$ retardation at 532 nm which we use to test the measurement modes. The fast axis of the wavelength was placed so that it did not coincide with the $x$-y or the $45^{\circ}-135^{\circ}$ laboratory axis. The measurement was taken using the quartz assisted method with quartz plates of thicknesses $\sim 1 \mathrm{~mm}$. Note that the wavelength span of the measurement ( $400-700 \mathrm{~nm}$ ) falls within a favorable measurement zone for quartz plates of this thickness (see Fig. 6.3).

### 6.5.2. Spatially resolved mode

Spatially resolved measurements of the Mueller matrix are done at a single wavelength. Clearly, if possible, it is convenient to chose a wavelength or use the adequate plates so that $\left|\sin \mathrm{CB}_{0}\right|,\left|\sin \mathrm{CB}_{1}\right|>0.5$.

The fast modulation of the PEMs is incompatible with the relatively low speed of the available multi-detectors (e.g. CCD) [148]. To sort this difficulty in recent years some complex strategies to integrate modulation cycles of the PEMs for use
with CCDs have been published [142, 149-152], but they are methods of difficult application, and most of them suited for instruments using one single PEM. Because of this incompatibility, single detectors such as PMTs are the most suitable to work with PEMs. To do space resolved measurements the methodology to use with PMTs consist on mapping the sample point by point, being each measurement one "pixel" of the resulting images. The main requirement is that only light that has passed through a tiny portion of the sample reaches the detector, and this can be done either by reducing the spot size incident on the sample or by placing a pinhole in front of the PMT [83].


Figure 6.5.: Imaging Mueller matrix elements of the zero order 532 nm quarter wave plate that was spectroscopically measured at Fig. 6.4. The dimension of each frame is $4 \mathrm{~mm} \times 4 \mathrm{~mm}$, where the step size to generate each pixel is $40 \mu \mathrm{~m}$ $\times 40 \mu \mathrm{~m}$.

To map relatively big areas of the sample several thousands of Mueller ma-
trix measurements are required (values between 20000 and 40000 are common). Although the phase modulation technique allows very fast measurements, the obtainment of scanning measurement can be a time-consuming process because of the large amount of data that needs to be acquired sequentially. Moreover it is worth to remember that for every point of measurement the four configurations (I, II, III and IV) concerning the quartz are required to measure the Mueller matrix. To do so we effectively map the sample four times, one for each configuration I, II, III and IV and once the 4 complete sets of data have been acquired the final Mueller matrix mapping of the sample surface is computed. We have found that this method is tens of times more time efficient that accessing to each configuration individually for every point of the mapping, due to the limited speed of the filter wheels where the quartz plates are mounted.

The spatial resolution of the automatic X-Y translator in which the sample is mounted is higher that the optical resolution of the instrument. For any of the four configurations the automatic translator must be able to reproduce, at least within the optical resolution, the same $\mathrm{x}, \mathrm{y}$ mapping for any of the four configurations. To check this correspondence during the mapping in each configuration a transmission intensity record of each point is also acquired. It is obtained from the dc voltage that the PMT circuit gives to the PMT at each measurement point, the lower the voltage, the greater the transmittance. If the four intensity mappings, are point by point correspondent, we can be sure that both the X-Y translator movements and also the introduction of que quartz plates maintain the lateral precision of the measurement.

In summary the result of a space resolved measurement consist of a $15 \mathrm{x}-\mathrm{y}$ images that correspond to the 15 elements of the normalized sample Mueller matrix and 4 extra images that represent the map of the intensity of the transmitted for every configuration.

Fig. 6.5 shows the Mueller matrix of the test quarter wave plate. As expected, there are no dramatic changes in the values of the Mueller matrix elements all over the plate surface. but changes of the order 0.01 rad are clearly visible in the color scale. They can be probably attributed to defects in the optical material of the plates. The step size used to create this image ( $40 \mu \mathrm{~m} \times 40 \mu \mathrm{~m}$ ) may be beyond the optical resolution of the instrument, which means that although we can generate images with high pixel resolution, we may not be able to resolve objects with sizes $<80 \mu \mathrm{~m}$. The values found here fully coincide with those obtained in the spectroscopic measurement of Fig. 6.4.

Fig. 6.6 shows the calculated values of the orientation of the fast axis and the retardation of the plate as obtained from the Mueller matrix of Fig. 6.5. Both parameters fall within the expected values. As expected, the central part of the plate is the area that better approaches to an ideal quarter wave retardation, with some small inhomogeneities that cause retardation defects at the periphery.

## Fast axis



Retardation


Figure 6.6.: Mapped fast axis and retardation of analyzed quarter wave plate calculated from the Mueller matrix in Fig. 6.5. The arrow indicates the approximate colour that would corresponds to an ideal quarter wave retardation (1.571 rad).

## Part IV.

## Experimental measurements

## Chapter 7.

## Optical activity of $\alpha$-quartz

$\alpha$-quartz $\left(\mathrm{SiO}_{2}\right)$ is, without any doubt, the crystal that historically has been more investigated crystal from the point of view of its optical activity. Around 200 years ahead from the first observations of Arago [14] and Biot [15] in the early 1800's quartz is still used in nearly every text book to introduce the subject of optical activity. As a result of this long-term investigation some of the the parameters of optical activity are known with great accuracy [153,154], and even their variations with temperature and pressure have been investigated $[155,156]$. Nowadays quartz crystal is widely available commercially in a great diversity of forms and cuts. The optical rotation properties of quartz have, for example, applications in laser technology, or as a standard of calibration for commercial polarimeters. Also in this work we have take advantage of the rotation properties of quartz plates to devise the novel quartz-assisted 2-MGE technique introduced in Chapter 6 . However all this applications and nearly all the measurements of the optical activity of quartz reported in the literature during the last two centuries have been performed along the unique direction in the crystal (the c-axis because quartz is a uniaxial crystal) in which it does not shows linear birefringence, and CB is the only effect. Therefore most of the commercial quartz crystals are cut perpendicular to its c-axis.

Although both linear birefringence and optical activity have been separately studied using a variety of techniques, very few attempts have been made to quantify the optical activity in presence of linear birefringence. For transparent materials, an accurate measurement of CB cannot be made without a full account of the related linear birefringence; for absorbent materials the linear dichroism and CD must also be taken into account. In nature circular and linear birefringent effects often appear together, so a method to measure CB in presence of linear birefringence is highly desirable. One such measurement technique is the HighAccuracy Universal Polarimeter (HAUP) [54,56, 57, 157, 158] . This technique is a variation of the old technique of crossed-polarizers, and it involves making measurements at slightly deviated angles from the position of maximum extinction of the polarizers. Typically, a monochromatic light source is used and the values of the optical activity are determined from the evolution of the detected waveform as a function of the angle of rotation of the polarizers.

In this chapter, we present a method to measure the optical activity of crys-
tals based on the measurement of the Mueller matrix with the 2-MGE technique introduced in Chapter 5. Using this technique, it is possible to perform measurements in directions other than along the optic axis of the crystal, where the magnitude of the linear birefringence is significantly larger tan the magnitude of the optical activity. As an application of this method we have measured the components of the gyration tensor of a right-handed (RH) and left-handed (LH) quartz crystal. To our knowledge our investigation [159] is the first measurement of the optical activity of a crystal based on the spectroscopic measurement of the Mueller matrix for directions of the crystals where linear birefringence and CB appear together.

### 7.1. Experimental details of the measurement

The basis of our experimental approach is the measurement of transmission Mueller matrix different quartz plates, but here a precise control of the direction of light propagation through the crystal is of fundamental importance. The samples studied in this work consist of two z-cut (c-axis perpendicular to the optical faces of the sample) quartz crystals $\left(\mathrm{SiO}_{2}\right)$ with opposite handedness. This kind of quartz plates is widely available commercially at a low price since this is the most usual type of cut. Quartz is a uniaxial crystal that belongs to the trigonal system and to the non-centrosymmetric crystallographic point group ${ }^{1} 32$. Quartz is enantiomorphous, that is it may occur as either right- or left-handed. In order to be enantiomorphous a crystal must have no element of symmetry that changes the handedness. For quartz the atomic arranges provide a screw axis that produces a helical distribution of atoms, and the optical activity of the two types of handedness has opposite signs. We will present results separately for the left-handed ( LH ) plate and for the right-handed ( RH ) plate. The measured thickness the LH plate is $1.02 \pm 0.01 \mathrm{~mm}$ while the thickness of the RH plate is $1.06 \pm 0.01 \mathrm{~mm}$.

A general scheme of the sample positioning in the instrument is also shown in Fig. 7.1 and a more precise diagram of the geometry of the problem is shown in Fig. 7.2. Our quartz plates had the c-axis perpendicular to the optical faces of the sample (the most common cut). A light beam is incident upon the crystal at an angle of incidence $\phi$ with respect to its c-axis. $l, d$ and $\theta$ are respectively the light path through the crystal, the sample thickness and angle with respect to the c-axis of the wave normal for the wave propagating through the crystal. If $d$ and $\phi$ are known the light path $l$ can be found by taking the approximation that the wave normal (given by Snell's law) in the crystal coincide with the direction

[^21]

Figure 7.1.: Diagram of the quartz plates positioning inside the 2-MGE. The sample could be precisely tilted by an angle $\phi$.


Figure 7.2.: Detail of the off-normal incidence measurement of circular and linear birefringence.
of propagation

$$
\begin{equation*}
l=\frac{d}{\cos \theta}, \quad \theta=\arcsin \left(\frac{\sin \phi}{\bar{n}}\right) \tag{7.1}
\end{equation*}
$$

where $\bar{n}$ is the mean refractive index, that depends on $\theta$ and will be defined in the next section. The sample was tilted in order to perform measurements at different values for $\phi$ (which also means different values for $\theta$ ). The rotation was around the y-axis defined by the modulation axis of the PEMs when PSG and PSA are orientated at $0^{\circ}$.

In the case of quartz crystal, the CB is known to be nonzero in a wide frequency region which includes the whole studied spectrum ( 220 to 800 nm ) [147]. In this region quartz is not absorbent [160] and therefore all parameters related to the absorptive nature of the crystal (including the CD) are zero.

### 7.2. Method of measurement

To study the light propagation through the quartz plate at oblique incidence, i. e. for directions where linear birefringence is also present we can use the propagation description first given by [161], which uses the constitutive equations given by Born [162]. This model is widely used in the literature [64,145, 163,164] particularly in experimental work, and gives the possible values of the refractive index $n$, for a given direction of the wave normal. Although this approach is only approximate, it has been shown to be accurate as long as the birefringence of the investigated crystal is not huge [165], as this model describes the phenomenon of optical activity within an accuracy of the product of the linear birefringence in the direction of observation by the gyrotropy parameters [164]. In this context the two possible refractive indexes are given by the positive roots of the equation [145, 164]:

$$
\begin{equation*}
\left(n^{2}-n_{01}^{2}\right)\left(n^{2}-n_{02}^{2}\right)=G^{2} \tag{7.2}
\end{equation*}
$$

where $n_{01}$ and $n_{02}$ are the refractive indices of the eigenwaves in absence of optical activity. $G$ is the scalar gyration parameter and is a measure of the optical activity for the direction in question. $G$ as a function of direction is given by

$$
\begin{equation*}
G=g_{i j} l_{i} l_{j}, \tag{7.3}
\end{equation*}
$$

in which the convention of summing over repeated indices is used. $l_{i}$ and $l_{j}$ are the direction cosines of the normal wave and $g_{i j}$ are the components of the gyration tensor that describe the optical activity of the crystal. In the enantiomorphous uniaxial crystals the gyration tensor [145] is

$$
\mathbf{g}=\left[\begin{array}{ccc}
g_{11} & 0 & 0  \tag{7.4}\\
0 & g_{11} & 0 \\
0 & 0 & g_{33}
\end{array}\right]
$$

and $G$ takes the form [163]

$$
\begin{equation*}
G=g_{11} \sin ^{2} \theta+g_{33} \cos ^{2} \theta \tag{7.5}
\end{equation*}
$$

where $\theta$ is defined in (7.1).
The solutions of Eq. (7.2) are [163]:

$$
\begin{equation*}
n_{1,2}^{2}=\left[n_{01}^{2}+n_{02}^{2} \mp\left[\left(n_{02}^{2}-n_{01}^{2}\right)^{2}+4 G^{2}\right]^{1 / 2}\right] / 2, \tag{7.6}
\end{equation*}
$$

and for a uniaxial crystal $\left(\varepsilon_{11}=\varepsilon_{22} \neq \varepsilon_{33}\right)$ :

$$
\begin{equation*}
n_{01}^{2}=\varepsilon_{11}, \quad n_{02}^{2}=\frac{\varepsilon_{11} \varepsilon_{33}}{\left(\varepsilon_{11} \sin ^{2} \theta+\varepsilon_{33} \cos ^{2} \theta\right)} \tag{7.7}
\end{equation*}
$$

where $\varepsilon_{11}=\varepsilon_{o}=n_{o}^{2}$ and $\varepsilon_{33}=\varepsilon_{e}=n_{e}^{2}$. The refractive indices $n_{o}$ and $n_{e}$ of quartz are well-known and are given by the 5 parameters dispersion relation calculated in [166].

The total birefringence $\left(\mathrm{T}_{R}\right)$ of the system is a combination of linear (LB and LB') and circular (CB) birefringences:

$$
\begin{equation*}
\mathrm{T}_{R}^{2}=\mathrm{LB}^{2}+\mathrm{LB}^{\prime 2}+\mathrm{CB}^{2}, \tag{7.8}
\end{equation*}
$$

with

$$
\begin{equation*}
\mathrm{T}_{R}=\frac{2 \pi}{\lambda}\left(n_{2}-n_{1}\right) l . \tag{7.9}
\end{equation*}
$$

According to Eq.(7.6) and assuming that $G^{2} \ll n_{01}^{2} n_{02}^{2}, T_{R}^{2}$ can be approximated by:

$$
\begin{equation*}
\mathrm{T}_{R}^{2} \simeq \frac{4 \pi^{2}}{\lambda^{2}}\left[\left(n_{02}-n_{01}\right)^{2}+\frac{G^{2}}{n_{01} n_{02}}\right] l^{2} . \tag{7.10}
\end{equation*}
$$

Due to the geometry of our oblique incidence experiment (see Fig. 7.2) the sample tilting does not cause birefringence in direction $45^{\circ}$ to the $x$ axis ( $\mathrm{LB}^{\prime} \simeq 0$ ) and LB itself can be considered as a measure of the total linear birefringence. Therefore Eq. (7.8) can be simplified to:

$$
\begin{equation*}
\mathrm{T}_{R}^{2} \simeq \mathrm{LB}^{2}+\mathrm{CB}^{2} . \tag{7.11}
\end{equation*}
$$

Comparing Eq. (7.11) to Eqs. (7.10) and (7.8), we can make the following identifications,

$$
\begin{gather*}
\mathrm{LB}=\frac{2 \pi}{\lambda}\left(n_{02}-n_{01}\right) l  \tag{7.12}\\
\mathrm{CB}=\frac{2 \pi G l}{\lambda \bar{n}} \tag{7.13}
\end{gather*}
$$

where $\bar{n}$ stands for a mean refractive index $\bar{n}=\sqrt{n_{01} n_{02}}$.
In the special case of propagation along the optic axis $\left(\theta=0^{\circ}\right) \mathrm{CB}$ is proportional to the $g_{33}$ component of the gyration tensor:

$$
\begin{equation*}
g_{33}=\frac{\mathrm{CB}_{\theta=0^{\circ}} \lambda n_{o}}{2 \pi l} . \tag{7.14}
\end{equation*}
$$

For light propagation along the optical axis in an optically active uniaxial crystal or for light propagation in any direction in an optically active isotropic medium the eigenmodes of optical propagation through the material are left and right circularly polarized modes, both of which have their own refractive indices. In contrast, the eigenmodes in a uniaxial material without optical activity are linearly polarized modes that are mutually orthogonal. As a result, in a random direction in an optically active uniaxial crystal the eigenmodes are left and right elliptical polarization modes with a major axis that coincide with the directions
of the linearly polarized eigenmodes that would be present in the absence of optical activity.

The ellipticity $k$ that we introduced in Chapter 1 is the ratio of the minor to the major axis of the ellipse of polarization and it can be used to define the waves that travel through the crystal unchanged in form. Along the optic axis, $k=1$ which indicates that the eigenmodes are circularly polarized light. The quantity $k$ can be calculated as follows [145]:

$$
\begin{equation*}
k=\tan \left(\frac{\gamma}{2}\right) \tag{7.15}
\end{equation*}
$$

where $\gamma$ is an angle defined by the ratio between the circular and linear birefringences

$$
\begin{equation*}
\tan \gamma=\frac{\mathrm{CB}}{\mathrm{LB}} \tag{7.16}
\end{equation*}
$$

With the relations of Eqs. (7.12) and (7.13) this ratio can be written as:

$$
\begin{equation*}
\frac{\mathrm{CB}}{\mathrm{LB}}=\frac{G}{\bar{n}\left(n_{02}-n_{01}\right)}, \tag{7.17}
\end{equation*}
$$

and substituting Eq. (7.5) here we can derive an expression to calculate $g_{11}$ from experimental measurements of the ratio $\mathrm{CB} / \mathrm{LB}$ :

$$
\begin{equation*}
g_{11}=\frac{1}{\sin ^{2} \theta}\left[\bar{n}\left(n_{02}-n_{01}\right) \frac{\mathrm{CB}}{\mathrm{LB}}-g_{33} \cos ^{2} \theta\right] . \tag{7.18}
\end{equation*}
$$

### 7.2.1. A model for the dispersion of the gyration tensor components

The first phenomenological models to describe the optical activity appeared early in the $20^{t h}$ century $[153,167]$, where the first tenuous connections were made between optical rotation and the interaction between atoms. One of the more successful approaches was carried out by Chandrasekhar using classical theory to study the optical rotation of quartz [20,147]. In this model there are two coupled oscillators that represent the smallest unit of the optically active crystal. The oscillators are assumed to be identical and undamped (that is, the crystal is non-absorbing). This coupled oscillator-model can also be treated quantum mechanically and describes the optical rotation $\rho$ of quartz in a wide frequency region:

$$
\begin{equation*}
\rho=\frac{k \lambda^{2}}{\left(\lambda^{2}-\lambda_{0}^{2}\right)^{2}} . \tag{7.19}
\end{equation*}
$$

The optical rotation $\rho$ is related to CB by $\rho=1 / 2 \mathrm{CB}$, and $k$ and $\lambda_{0}$ are the two unique parameters of the model.

To measure the components of the gyration tensor $G$ we propose to use this dispersion model to parameterize $g_{11}$ and $g_{33}$ :

$$
\begin{equation*}
g_{i i}=\frac{A_{i} \lambda^{3}}{\left(\lambda^{2}-B_{i}^{2}\right)^{2}}, \tag{7.20}
\end{equation*}
$$

where $A_{i}$ and $B_{i}$ are the parameters to be determined. Note that this model can be straightforward derived from (7.19) if we consider that $G=\lambda \bar{n} \rho / \pi l$ (see Eq. (7.13)) and we use the approximation that $\bar{n}$ is constant. We have found that Eq. (7.20) is a good fit for our experimental $g_{11}$ and $g_{33}$ data. It is interesting to note that while for quartz $\mathrm{CB}\left(\propto\left(n_{-}-n_{+}\right)\right)$varies by more than one order of magnitude in the interval range from 200 nm to 800 nm , while the variation of the mean refractive index $\bar{n}\left(\propto\left(n_{-}+n_{+}\right)\right)$in this same interval is only around the $7 \%$. Thus it is not strange that if Eq. (7.19) constitutes a good fit for CB, Eq. (7.20) can be used to fit the gyration tensor with success.

### 7.3. Data analysis of the Mueller matrix of quartz



Figure 7.3.: (Color online). Spectroscopic Mueller matrix for a RH plate of quartz at small incidence angles $(\phi)$.

## Chapter 7. Optical activity of $\alpha$-quartz



Figure 7.4.: (Color online). Spectroscopic Mueller matrix for a LH plate of quartz at small incidence angles ( $\phi$ ).

The spectroscopic Mueller matrices measured at different small incident angles for RH and LH plates are respectively presented in Fig. 7.3 and Fig. 7.4. In both cases, and for all the orientations of the plates, Mueller matrix elements $m_{01}, m_{02}$, $m_{03}, m_{10}, m_{20}$ and $m_{30}$ are approximately zero in all the studied spectral range (note that these elements have been multiplied by a factor 10). This indicates that at this wavelength range and for these small angles of incidence there is not significant diattenuation and the Mueller matrices of the quartz plates basically take the form of a general retarder given in Eq. (3.24).

However, a crystal with retardation properties that vary rapidly with wavelength may induce depolarization if the system collects a band of wavelengths rather than a single wavelength. Since the retardation is proportional to the inverse of the wavelength, a band of wavelengths will result in a different retardation for each wavelength, resulting in partially polarized light [168]. To minimize this depolarization, the monochromator slit width can be reduced and/or a smaller core optical fiber used at the input of the monochromator [169]. Fig. 7.5 depicts a comparison between the fraction of polarized light $\beta$ (see Eq. (1.33)at normal incidence and at $60^{\circ}$ incidence. As it can be seen in the figure the depolarization becomes particularly significant for big angles of incidence, where the
observed retardation has an important variation with wavelength, and it needs to be considered.


Figure 7.5.: (Color online). Fraction $\beta$ of polarized light for normal and oblique incidence. For the same conditions the RH plate is a bit more depolarizing because it is a slightly thicker than the LH plate.

In order to consider the depolarization of the measured Mueller matrices we have factorized them according to the Lu-Chipman decomposition given as in Eq. (4.23). This factorization allows the separation of the depolarization properties from the retardation and dichroic properties. For a uniaxial crystal the Mueller matrices $\mathbf{M}_{R}$ and $\mathbf{M}_{D}$ commute [170], so the Lu-Chipman decomposition can be used to factor the experimental Mueller matrix. Once the retardation matrix $\mathbf{M}_{R}$ has been calculated we use Eqs. (4.32d), (4.32e) and (4.32f) to extract the values of linear and circular birefringence.

When dealing with samples that introduce significant retardations, such as crystals, the determination of the sample Mueller matrix may not be sufficient to determine all its birefringent characteristics. In Eq. (3.24) it can be seen that many of the Mueller matrix elements contain trigonometric functions. Furthermore, the arccos, which is multi-valued, is required to determine the total birefringence (see Eq. (4.33)). For systems with small retardations this does not constitute a problem since the total retardation must fall in the $[0, \pi]$ interval. However, for systems with big retardances (e.g. some crystals) this problem prevents us from determining the birefringences unless we know their order. For example at 252 nm the Mueller matrix of the left handed plate of quartz with an
angle of incidence of $9^{\circ}$ is the following:

$$
\mathbf{M}=\left[\begin{array}{cccc}
1 & 0.002 & 0.003 & 0.006  \tag{7.21}\\
0.005 & 1.002 & 0.009 & 0.010 \\
0.001 & 0.011 & 1.000 & 0.002 \\
0.000 & 0.010 & -0.006 & 0.988
\end{array}\right]
$$

This matrix is nearly the same as the identity Mueller matrix, which, one would obtain in transmission for an isotropic medium. Of course, the sample is not isotropic, although its Mueller matrix appears to be that of an isotropic medium at this wavelength. The importance of the order must always be considered when analyzing these results, since a single Mueller matrix is not always sufficient to provide the characterization of a polarization element.

For large angles of incidence a significant linear diattenuation appears in the measured Mueller matrices as they no longer have all the vanishing elements shown in Eq. (3.24). In this case the linear diattenuation is not due to an anisotropic absorbance in quartz, but rather is caused by the different reflectivities of s- and p- polarized light [171]. The polar decomposition allows us to obtain the linear diattenuation of the sample from the matrix factor $\mathbf{M}_{D}$ [82]. The top panel of Fig. 7.6 displays the linear diattenuation for a quartz plate at an angle of incidence of $60^{\circ}$.

The bottom panel of Fig. 7.6 shows the total retardance $T_{R}$ as determined using Eq. (4.33) for the RH quartz plate with a incidence angle of $60^{\circ}$. At this large incident angle the linear birefringence of quartz is big compared to $\pi$ and, the total retardance goes through several oscillations between 0 and $\pi$ as a function of wavelength. To determine the real $\mathrm{T}_{R}$ we should also know their order to unfold the "folded" spectrum of Fig. 7.6. Fortunately we do not need to know the order of the birefringence to determine the ellipticity and the gyration tensor of quartz. Eqs. (7.16) and (7.18) show that it is only necessary to determine the ratio of CB to LB rather than determining LB and CB separately. That is, we do not have to determine $\mathrm{T}_{R}$. In terms of Mueller matrix elements, the ratio $\mathrm{CB} / \mathrm{LB}$ is given by:

$$
\begin{equation*}
\frac{\mathrm{CB}}{\mathrm{LB}}=\frac{\mathrm{m}_{R_{12}}-\mathrm{m}_{R_{21}}}{\mathrm{~m}_{R_{32}}-\mathrm{m}_{R_{23}}} . \tag{7.22}
\end{equation*}
$$

Therefore, the ratio $\mathrm{CB} / \mathrm{LB}$ can be straightforward determined once the matrix $\mathbf{M}_{R}$ is known. Although Eq. (7.22) is a significant simplification, each of the Mueller matrix elements must be measured accurately. Obviously, a large error occurs when the denominator $\left(\mathrm{m}_{R_{32}}-\mathrm{m}_{R_{23}}\right)$ approaches to zero, which occurs at several wavelengths for these samples. Fortunately, as it will be discussed in the next section, these points can be easily identified and can be removed from the calculation. Another experimental issue is that the retardation oscillation frequency increases at small wavelengths (See bottom panel of Fig. 7.6), and


Figure 7.6.: Linear diattenuation and "folded" total retardation $\mathrm{T}_{R}$ induced by a 1 mm thick, z -cut, RH quartz plate at an incidence angle of $60^{\circ}$.
the consecutive maxima and minima may be separated only by few nanometers. Thus, a good spectral resolution as determined by the monochromator must be sufficient to resolve these oscillations. Thinner samples would also reduce the total retardation and therefore avoid this problem.

### 7.4. Results

We have examined the retarding properties of the quartz plates at angles slightly deviated from the position of normal incidence by measuring their Mueller matrices. These measurements show the decrease on the ellipticity $k$ as we deviate from the optic axis. Figs. 7.3 and 7.4, respectively show the normalized Mueller matrix for a RH and a LH plate of quartz at small angles of incidence. Small deviations from the condition of normal incidence translate in obvious differences in the matrices, which indicate that even for situations close to the propagation along the optical axis the LB becomes comparable to CB. Qualitatively these matrices resemble the matrix for a general retarder given in Eq. (3.24), where
the different handedness is denoted by the opposite signs that Mueller matrix elements $m_{12}, m_{21}, m_{13}, m_{31}$ take in Figs. 7.3 and 7.4. Elements $m_{23}$ and $m_{32}$ do not change in sign but this is due to the geometry of our oblique incidence measurements. The only elements that are not sensitive to the handedness of quartz, no matter the orientation of the plates, are the diagonal elements.


Figure 7.7.: (Color online). Spectroscopic evolution of the ellipticity $k$ for small angles of incidence.

Fig. 7.7 shows the ellipticity $k$ for the RH and LH plates of quartz calculated from the values of LB and CB according to Eq. (7.15). The ellipticity describes the polarization state of the waves that propagate unchanged through the crystal. As expected, at normal incidence, the ellipticity is 1 for the RH and -1 for the LH , and it quickly diminishes as the angle of incidence becomes more oblique. The points missing in Fig. 7.7 are points placed at wavelengths at which the total retardance $T_{R}$ is close to $2 \pi$ where accurate measurement of the ellipticity is not possible.

The measurement of the 2 independent components of the gyration tensor of quartz requires the orientation of the samples in two different configurations. The first measurement is made at normal incidence $\left(\theta=\phi=0^{\circ}\right)$ where $g_{33}$ is calculated using Eq. (7.14). This is the usual measurement configuration where the sample does not show LB, so the determination is straight-forward. The determination of $g_{11}$ is obtained using Eq. (7.18) and requires that the measurements be taken at a large incidence angle to avoid small values for the $\sin \theta$ that would translate in large errors in $g_{11}$. We have chosen $\phi=60^{\circ}\left(\theta \sim 33^{\circ}\right)$ which is close to the maximum incidence angle for which we can guarantee that both the extraordinary and ordinary rays are collected by the detector.


Figure 7.8.: Measured components of the gyration tensor ( $\circ$ ) and fitted dispersion relation (straight line) for the RH plate.

The experimental values and the fitted curves for the $g_{11}$ and $g_{33}$ components of the RH and the LH quartz plates are shown in Figs. 7.8 and 7.9. The fitting to the model of Eq. (7.20) has been performed with a weighted Levenberg-Marquadt nonlinear minimization procedure using the reduced $\chi^{2}$ as a figure of merit. Not all the experimental points have been used for the fitting (see bottom panels of Figs. 7.8 and 7.9), since we have only used those points that correspond to wavelengths for which the ratio $\mathrm{CB} / \mathrm{LB}$ can be measured with highest accuracy. In particular we have only considered points for which $\left|\mathrm{m}_{R_{32}}-\mathrm{m}_{R_{23}}\right| / 2 \geq 0.98$, i. e. points for which the absolute value of the denominator of (7.22) is near its maximum value.

Table 7.1 summarizes the fitting parameters for the four fits presented in Figs.

Chapter 7. Optical activity of $\alpha$-quartz


Figure 7.9.: Measured components of the gyration tensor ( $\circ$ ) and fitted dispersion relation (straight line) for the LH plate.
7.8 and 7.9. In all cases the dispersion relation given in Eq. (7.20)constitutes a good fit for experimental $g_{33}$ and $g_{11}$ data. Although the accuracy of $g_{11}$ is considerably lower than $g_{33}$, it is clear that the shape of the dispersion curves for both components are fairly similar.

A comparison between our data and some of the previously published data is presented in Table 7.2 at two wavelengths. There are few experimental measurements of $g_{11}$ of quartz available in the literature and most of them are restricted to a single wavelength. With the exception of ref. [55], our results agree with literature values for $g_{33}$ within the stated error limits, and the agreement for $g_{11}$ is within 2 standard deviations with all the literature values.

We also calculated the experimental ratio of the two gyration tensor compo-

Table 7.1.: Fitted parameters for the model in (7.20) with $\lambda$ expressed in nm .

| Component | LH | RH |
| :---: | :--- | :--- |
| $g_{11}$ | $A_{1}=0.0277 \pm 0.0017$ | $A_{1}=-0.0298 \pm 0.0015$ |
|  | $B_{1}=105.6 \pm 5.7$ | $B_{1}=91.1 \pm 4.2$ |
|  | $\chi^{2}=0.82$ | $\chi^{2}=0.59$ |
| $g_{33}$ | $A_{3}=-0.0609 \pm 0.0002$ | $A_{3}=0.0604 \pm 0.0005$ |
|  | $B_{3}=97.54 \pm 0.05$ | $B_{3}=97.33 \pm 0.05$ |
|  | $\chi^{2}=1.33$ | $\chi^{2}=0.94$ |



Figure 7.10.: (Color online). Ratio of the two independent gyration tensor components $g_{11} / g_{33}$.
nents $g_{11} / g_{33}$. In the nonabsorptive wavelength regime this ratio should be -0.5 due to a symmetry argument that applies to the tetrahedron building units of crystalline quartz [172]. Fig. 7.10 shows our spectroscopic experimental results for the $g_{11} / g_{33}$ ratio. They are consistent with the symmetry argument as most points are within $5 \%$ of the theoretical ideal value, and the calculated mean value is $-0.486 \pm 0.028$. Using this value in Eq. (7.5) we can deduce that there is no optical rotation $(G=0)$, that is, there exists an isotropic point for the optical activity that is cylindrically symmetric around the optic axis, for propagation directed $\sim 55^{\circ}$ from the optic axis.

We have presented a technique for measuring the optical activity in uniaxial crystals in situations where both linear birefringence and circular birefringent ef-

Table 7.2.: Comparison of data for the components of the gyration tensor of quartz at room temperature determined from this study and from selected values in the literature.

| Wavelength (nm) | Ref. | $\left\|g_{33}\right\| \times 10^{-5}$ | $\left\|g_{11}\right\| \times 10^{-5}$ |
| :---: | :---: | :---: | :---: |
| 632.8 | $[57]$ | $10.1 \pm 0.2$ | $5.9 \pm 0.4$ |
|  | $[158]$ | 10.11 | 6.11 |
|  | $[55]$ | $13.6 \pm 0.5$ | $5.7 \pm 0.5$ |
|  | $[173]$ | 10.528 | 5.39 |
|  | $[157]$ | - | $\sim 5.2$ |
|  | This work* | $10.06 \pm 0.07$ | $4.8 \pm 0.5$ |
| 510 | $[161]$ | $12.96 \pm 0.2$ | $5.82 \pm 0.4$ |
|  | $[157]$ | - | $\sim 6.5$ |
|  | This work* | $12.81 \pm 0.08$ | $6.1 \pm 0.6$ |
| * Values obtained from the average of RH and LH results |  |  |  |

* Values obtained from the average of RH and LH results
fects are present. Hence this technique allows one to measure the CB of opticallyactive crystals in directions different from the optical axis. This technique has been applied to the measurement of the two independent components of the gyration tensor of RH and LH crystalline quartz. The basis of the technique is the measurement of the ellipticity for the waves that transmit unchanged in their state of polarization through the crystal. For directions out of the optical axis the ellipticity of these waves is very small and precise measurements require a highly sensitive experimental approach, such as provided by a transmission 2-MGE experiment. To our knowledge this this is the first description of a systematic methodology to obtain spectroscopic measurements of the optical activity in crystals for direction out of the optic axis.

The presented results show the ability of the 2-MGE to deal with the phenomenon of optical activity in presence of linear birefringence. Although the 2MGE is a suitable instrument for this kind of measurements, we have presented a general approach to the measurement procedure for any instrument capable of measuring a sample Mueller matrix. We start from the basis that the Mueller matrix of the crystal can be measured and we demonstrate how effects inherent to the optical activity can be extracted from the matrix even when much higher linear birefringence is present. Nowadays there exist several different instruments that are capable of determining the Mueller matrix of a sample, so this method could be applied in several laboratories.

Another advantage of this technique is that different components of the gyration tensor of the crystal can be determined without having to use samples cut
according to several different crystallographic planes. This is because we do not exclusively analyze light transmitted in directions perpendicular to these planes, besides we also use a more general oblique incidence configuration where more than one component of the gyration tensor is contributing to the observed optical activity. As an example we have measured the $g_{11}$ and $g_{33}$ components of quartz using only a $z$-cut crystal with light incident at $60^{\circ}$ from the optic axis. In this configuration both components $g_{11}$ and $g_{33}$ contribute to the optical activity, but as $g_{33}$ can be well determined from normal incidence measurements the only remaining incognita is $g_{11}$.

We envision that a similar approach to the one presented here can be used for several other crystals with point groups $3,4,6,32,422$ and 622 (International notation), which are optically active and have linear birefringence. Moreover, this technique may also be useful for crystals with point groups $\overline{4}$ and $\overline{4} 2 m$ which do not show optical activity for light propagating along the optic axis, but do show optical rotation in directions where there is also birefringence.

## Chapter 8.

## Chiral induction by hydrodynamic effects

In recent years the induction of chirality by hydrodynamic forces has been focus of interest and debate in chemistry and has arosed new experimental efforts in an attempt of understanding a phenomenon that still is quite unknown and that, until very recently, remained mostly unexplored.

The action of mechanical forces originated by flows on chemical phenomena was generally considered to be insignificant and, thus the experimental observation that flows can play a certain role in the emergence of supramolecular chirality was surprising and, even at some point, remained controversial. Two pioneering reports, published in 1976 [174] and 1993 [175], initiated the investigation on two different species of molecular aggregates. However, they were widely refuted because their claimed results were apparently not easily reproducible and the mainstream opinion associated them to measurement errors. The first report which demonstrated the inductive effect of flows on supramolecular chirality, was published by the group of Ribó in 2001 [176]. It showed that the direction of stirring in a swirling vortex can be irreversibly transferred to the chiral sign of the CD-spectra for a certain type of porphyrin J-aggregates.

Many investigations about this subject have been carried out in some supramolecular species of J-aggregates. These type of aggregates tend to be thin and longshaped particles, that, even in solution, if there exists some level of orientation show relatively strong linear polarization-dependent optical effects. Solutions of these aggregates constitutes a good example of an optically anisotropic liquid, in which the measurement of the optical activity by using the Mueller matrix method described in this thesis is a fairly more accurate approach that conventional chiroptical methods.

During the development of this thesis a large number of experiments on this subject were performed, but in this chapter we will only present the most remarkable ones. It was important to study this phenomenon extensively because of the number of factors that a priori could be involved in the experiments (species of aggregates or type particles, the level of aggregation and the size of aggregates, the geometry of the cuvette, the stirring speed, etc), but also, and none less important, to assure the reproducibility of everyone of the described measurements.

### 8.1. Spectroscopic measurements

### 8.1.1. Porphyrin aggregates

Porphyrin aggregates considered herein have an UV/Vis absorption band that is clearly separated form those of their monomers, so that the dichroic contributions can be easily studied in spite of the presence of monomer species in the solution. We have measured J-aggregates of the porphyrins $\mathrm{TPPS}_{1}, \mathrm{TPPS}_{2 o}, \mathrm{H}_{4} \mathrm{TPPS}_{3}$ and $\mathrm{H}_{4} \mathrm{TPPS}_{4}$. All of them are achiral amphiphilic porphyrins and their J-aggregates have have thin long acicular shapes (nanoribbons)

In Ref. [176] it was shown for first time that the chiral sign (understood as the sign of CD spectra) of the $\mathrm{H}_{2} \mathrm{TPPS}_{3}$ porphyrin could be selected by the direction of vortical stirring during formation of the J-aggregates. This was an irreversible induction of CD and it did not manifested in the other cited porphyrins, but constituted the first clear evidence that supramolecular chirality in porphyrin J-aggregates could be induced by hydrodynamic forces.


Figure 8.1.: Scheme of the experimental disposition used in the measurements involving a stirred cuvette. The cuvette was equipped with a $5 \mathrm{~mm} \times 2 \mathrm{~mm}$ magnetic bar that, that in conjunction with a magnetic stirrer, permitted stirring in CW and CCW directions at a selectable speed. The light beam was directed to the central part of the generated vortex.

In the framework of this thesis we observed a surprising new type of induction when a cuvette containing a water solution of J-aggregates was in situ stirred while taking polarimetric measurements. In these J-aggregates the stirring effect caused and instantaneous and reversible emergence of CD, i.e. the observation of a certain sign or just the opposite one depended on the direction of stirring during the measurement. This effect has been reported in a number of recent publications that contained interpretations not really supported by experiments [177-179]. Our approach to the problem has been directed toward obtaining as much information as possible from polarimetric experiments on stirred solutions.

Fig. 8.1 shows the basic experimental setup used for the measurements presented in this section: the solution was contained in a 10 mm pathlength square section quartz UV/Vis cuvette. The cuvette was in situ stirred at a constant speed using small magnetic bars in clockwise (CW) or counter-clockwise (CCW) during the spectroscopic measurement of the Mueller matrix.

We first observed this reversible mechanical induction of supramolecular optical activity in J-aggregates of $\mathrm{H}_{4} \mathrm{TPPS}_{4}$ and it is the porphyrin that we have studied more in detail. The size and shape of the $\mathrm{H}_{4} \mathrm{TPPS}_{4} \mathrm{~J}$-aggregates depend on the procedure of their preparation ${ }^{1}$. The $\mathrm{H}_{4} \mathrm{TPPS}_{4} \mathrm{~J}$-aggregates, as inferred from the AFM analysis, are straight nanoribbons about 200 nm long, 50 nm wide, and with a thickness of 3.2 nm (bilayer structure), but those in aged solutions can show lengths in the $\mu \mathrm{m}$ range and thicknesses of 4.8 nm (see Fig. 8.2).


Figure 8.2.: AFM topography with section analysis of aged $\mathrm{H}_{4} \mathrm{TPPS}_{4} \mathrm{~J}$ aggregates.

Spectroscopic Mueller matrix measurements of a solution prepared from $25 \mu \mathbf{l}$ of an aged solution (one year old) of $\mathrm{H}_{4} \mathrm{TPPS}_{4} 1.4 \cdot 10^{-3} \mathrm{M}$ were taken at each nm in the interval from 350 nm to 800 nm , so that, at the end, 451 normalized Mueller matrices were measured for each sample configuration (stagnant, CW and ACW). Fig. 8.3 depicts spectroscopic generalized ellipsometry data in the interval range between 400 and 600 nm for the aggregated porphyrin. Considerable difference between the stagnant solution and the CW and CCW stirred solutions is in general observed except for the elements of the diagonal. Also for the elements of the anti-diagonal ( $m_{03}, m_{12}, m_{21}, m_{30}$ ) important differences between the CW and CCW directions are seen.

[^22]

Figure 8.3.: Detail of the experimental normalized Mueller matrices for CW stirring, CCW stirring and stagnant (no stirring).

Fig. 8.4 shows the CD and CB wavelength dependencies obtained as a result of the analysis following the analytic inversion method introduced in section 4.1 of the experimental matrices shown in Fig. 8.3. The J-aggregates of $\mathrm{H}_{4} \mathrm{TPPS}_{4}$ in stagnant solutions show a permanent low intense CD signal because this molecule is chiral, but by vortex stirring a much higher intense CD signal emerges. Thus, the natural chirality of the porphyrin is overruled by the mechanical stirring effect and the chiral sign is determined by the stirring direction. Also, the wavelength dependence of the measured CD spectra is bisignated and is Kramers-Kronig consistent with the measured the CB spectra. To illustrate this consistency we display in an inset of the Fig. 8.4 the qualitative theoric CB spectrum corresponding to a CD bisignated band of opposite signs calculated by a two-oscillator Lorentz model that is Kramers-Kronig consistent.

The reversible flow-induced CD was also detected for the micelle-like aggregates of the porphyrin $\mathrm{TPPS}_{1}$, the J-aggregates of the $\mathrm{TPPS}_{2 o}$ porphyrin, but the CD spectra of the $\mathrm{H}_{4} \mathrm{TPPS}_{3} \mathrm{~J}$-aggregates was not sensitive to the stirring [180]. In the case of the $\mathrm{H}_{4} \mathrm{TPPS}_{3}$, there exists the irreversible mechano-chiral effect [176], but once obtained, the CD spectra remains unchanged independently of the stirring direction. Although the J-aggregates of the porphyrins showing the irreversible and the reversible effect belong to different compounds, $\left(\mathrm{H}_{4} \mathrm{TPPS}_{3}\right.$ and $\mathrm{H}_{4} \mathrm{TPP} S_{4}$ respectively), there is no chemical reason to expect different signs of the CD for the chiral exciton arising by folding or bending of the mesophases of these two different compounds.


Figure 8.4.: Spectroscopic determination of CD and CB as a function of the stirring direction. The inset shows that a Kramers-Kronig transformation of a CD bisignated band yields the same type of profiles measured for CB.

We will discuss in detail the explanation underlying in these experimental measurements at the ending part of this chapter, because in following sections we will present new experimental data that will offer us additional information about the phenomenon. However at this point, and in order to motivate the contents of the following sections, it is worth to anticipate that our hypothesis is that the emergence of chirality through the effect of the hydrodynamic forces originated by a swirling vortex must be related to the feasibility of the flow to orient, fold and twist the particles in the fluid. Consequently, the shape and the elasticity of the aggregates and also the characteristics of the flows in the cuvette may have influence on the occurrence of this phenomenon.

## Cuvette geometry. Cylindrical flasks

The evidence that the flow regime inside the cuvette was responsible for the observed induction of supramolecular chirality made us suspect that the experi-
ment might yield different results placing the solution in containers of different geometry. To study this dependence the stirring experiments on solutions of the $\mathrm{H}_{4} \mathrm{TPP} S_{4}$ porphyrin were repeated in the same contions as in the previous chapter but this time using cylindrical flasks as containers.

In the case of an square section cuvette the sign and intensity of the CD showed a low dependence on the rotation speed, for example they were quite similar for 1500 rpm and 5000 rpm (see left column of Fig. 8.5). In contrast, we found that the CD values in the cylindrical flask were strongly depending on the rotation speed. At low rotation speeds the CD signals did not have a well defined sign and showed a noisy pattern (top right panel of Fig. 8.5). However, at high rotation speed a clear bisignated CD signal emerged, but it was of opposite sign (bottom right panel of Fig. 8.5) to those obtained in the square section cuvette


Figure 8.5.: Comparison of the CD spectra obtained at different stirring speeds for square section and cylindrical section cuvettes.

The differences in the determined CD spectra can be explained because in the case of the cylindrical flask, due to the presence of an ascending chiral flow not appearing in a square section cuvette, stirring would be expected to lead to two mechano-chiral effects of opposite sign: a descending vortex in the middle region and an ascending vortex around the walls. Therefore, the CD signal observed in our transmission experiments (light propagation orthogonal to the long flask axis) must correspond to the addition of two CDs of opposite sign. Obviously, as the rotation speed of the magnetic bar increases more liquid is expelled towards the walls, the gradients of shear rate at the flask wall increases
and its effect overcomes that of the central vortex. Fig. 8.6 shows and schematic representation of the main contributing chiral flows for a square section cuvette and a a cylindrical flask. The observed different flow regimes as a function of the geometry of container explains the previous results on the irreversible stirring effect observed during the formation of the J-aggregates of $\mathrm{H}_{4} \mathrm{TPPS}_{3}$. In those experiments $[176,181]$ solutions were stirred at high rotation speeds in small cylindrical tubes ( 0.8 cm diameter), where the correlation between the CD sign and the stirring direction must be attributed to the ascending flow.


Figure 8.6.: Schematic representation of the flow pattern for two different cuvette geometries.

According to our interpretation, in the square section cuvette the CD signal would only emerge in the middle of the cuvette and not at its walls. Therefore, the chiral bias observed would be caused by the gradients of shear rate in the descending vortex in the center of the cuvette, because the irregular flow in the walls would yield a racemic mixture or a small chiral bias of folded/bent mesophases. In the case of the square-section cuvette the descending vortical flow acts as an attractor of the particles in solution. This was also confirmed by observation of the polarization-independent transmittance changes upon stirring. In the square-section cuvette, stirring led to significant changes in the absorbance values compared to stagnant conditions (see Fig. 8.7); in the central part of the cuvette there was an increase of absorbance in the J-aggregate bands ( 490 nm and 705 nm ), but no absorbance differences were detected for the monomer absorption ( 434 nm ). In contrast, measurements took on cylindrical flask did not show such absorbance changes.

Space resolved measurements presented later in this chapter are a visual demonstration that in square-section cuvettes the CD emergengence only takes place in the central part of the cuvette.


Figure 8.7.: Absorbance of stirred and stagnant solutions of long-shaped $\mathrm{H}_{2} \mathrm{TPPS}_{4} \mathrm{~J}$-aggregates in a square section UV/Vis cuvette ( 1 cm ). There are changes for the J-aggregate absorption ( 490 nm and 705 nm bands) but not for the diprotonated monomer ( 434 nm and 650 nm bands).

### 8.1.2. Rossette nanotubes

The induction of optical activity upon stirring was also observed in a different system from the porphyrin J-aggregates presented above. We carried out experiments with stirred solutions of a new class of adaptive nanotubular architectures know as rossete nanotubes (RNT) resulting from the self-assembly and self-organization of biologically inspired materials [182-184]. The results presented here correspond to solutions of organic compounds featuring 2 synthetic heterobicyclic base ( $\mathrm{G} \wedge \mathrm{C}$ ) that contain the hydrogen bonding arrays of the DNA bases guanine and cytosine on opposite faces of the molecule. This twin system self-assembles spontaneously, to form a six-membered supermacrocycle (rosette) maintained by 18 H -bonds, which in turn self-organizes into stable and architecturally complex 1-D helical stacks defining an unoccluded central channel, the rosette nanotube. The diameters of the nanotubes are about 5 nm , while the can have lengths of several hundreds of nm. The nanotube RNT samples were provided by H. Fenniri (Alberta, Canada).

Fig. 8.8 shows the measured spectroscopic Mueller matrix normalized to the $m_{00}$ element for CW and CCW stirring of a methanol RNT solution contained in a square section cuvette. Samples were scanned from 235-340 nm in intervals of 1 nm . The CD profiles shown in Fig. 8.9a were calculated from the analytic inversion of the Mueller matrix in Fig. 8.8 and evidence the induction of chiral sign as function of the direction of stirring. Note from this figure that the measurement with 2-MGE in stagnant configuration on RNTs solutions did not yield any discernible value of CD. However we the sensitivity of our 2-MGE at wavelengths around 240 nm is not sufficient to discern CD from noise in samples with circular dichroic signals below 10 mdeg. The total in-plane linear dichroism


Figure 8.8.: Normalized spectroscopic Mueller matrix of a solution of RNTs obtained for CW and CCW stirriging and in stagnant.

LDm is taken as $\mathrm{LD}_{m}=\left(\mathrm{LD}^{2}+\mathrm{LD}^{\prime 2}\right)^{1 / 2}$ and is displayed in Fig. 8.9b , and, analogously the in-plane linear birefringence LBm is $\mathrm{LB}_{m}=\left(\mathrm{LB}^{2}+\mathrm{LB}^{\prime 2}\right)^{1 / 2}$ and is shown in Fig. 8.9c. The greater values of LDm and LBm obtained for CW and CCW stirring show that, as result of the vortical flows, the long shaped RNTs increase their degree of orientation with respect to the stagnant configuration.


Figure 8.9.: Comparison of the retrieved optical effects for CW stirring (solid line), CCW stirring (dashed line) and stagnant (dotted line).

An interesting fact is that RNTs that are not made of twin compounds (i.e. they feature only a single $G \wedge C$ ) do not show any change in $C D$ upon stirring, some-
thing similar to what happened to the $H_{4} T P P S_{3}$ described in the previous section. At the moment we have not extracted any final conclusion of these different behaviours, but we think that they can be significant in the near future for a further understanding of the phenomenon.

### 8.2. Space-resolved measurements in square section stirred cuvettes

Space-resolved measurements were performed following to the methodology described in chapter 6 . The objective was to obtain a map of the optical polarization properties of the entire cuvette in order to be able to detect differences on the measured effects at the different parts of the cuvette. We were interested in performing such mappings because spectroscopic measurements taken at different zones of the cuvette (for example comparing the central part to a lateral) had already indicated us differences in the measured effects [180]. In this case spaceresolved measurement were only possible in square section cuvettes, because for the cylindrical flasks measurements are only possible at the center of the cylinder because of the reflections produced by the non-planar flask surface in the laterals.

Experiments shown in this section where performed with a solution of $\mathrm{H}_{4} \mathrm{TPPS}_{4}$ J-aggregates in stagnant and stirred conditions. Figs. 8.10, 8.11 and 8.12 show, respectively, the Mueller matrices for no stirring, CW stirring and CCW stirring. All three measurements were recorded with the same sample a using a 1.5 mm diameter for the beam spot. Each mapping was done in about 20 minutes with a pixel resolution of $0.8 \mathrm{~mm} \times 0.8 \mathrm{~mm}$.

The space resolved measurements in square section cuvettes confirmed that the induction of optical activity takes place in the central part of the cuvette (see Fig. 8.13), in coincidence with the central chiral descending vortex. Fig. 2 shows that at 485 nm the switch of optical activity is clearly visible in the CD maps, because this wavelength corresponds to the high-energy peak of a bisignated CD absorption band. For CB the change of sign can also be noted, but is less appreciable because this wavelength does not correspond to any peak of CB.

The values of LD and $\mathrm{LD}^{\prime}$ (or alternatively LB and $\mathrm{LB}^{\prime}$ ) presented in Fig. 8.13 can be used to construct a vectorial representation of the measured projections of linear dichroism (see Fig. 8.14). The length of the vectors, $l$, and their orientation with respect to the horizontal laboratory axis, $\theta$ have been calculated according to

$$
\begin{align*}
l & \propto \sqrt{\mathrm{LD}^{2}+\mathrm{LD}^{\prime 2}},  \tag{8.1}\\
\theta & =\frac{1}{2} \arctan \frac{\mathrm{LD}^{\prime}}{\mathrm{LD}} . \tag{8.2}
\end{align*}
$$

The vectors drawn can be correlated with the average orientation of the nanophases in the solution.
8.2. Space-resolved measurements in square section stirred cuvettes


Figure 8.10.: Case of a stagnant solution.

Figure 8.15 shows a superposition of the CD false color map and calculated axes of linear dichroism. For both, CW and CCW stirring, there appears to be a small bend of the vortex axis at the top of cuvette. This fact allows areas with weaker flows (top-left corner for CW stirring and top-right corner for CCW stirring), where the particles can maintain more stable orientation, yielding greater values of LD and LD' that translate in longer vectors.

Chapter 8. Chiral induction by hydrodynamic effects


Figure 8.11.: Case of a CW stirred solution.

### 8.3. Discussion

The phenomenon reviewed in this chapter represents the selection of the chirality of a supramolecular structure, i.e. a process of chiral symmetry breaking. Only recently we have been able to agglutinate all the experimental data necessary to describe accurately this phenomenon. Research on the physico-chemical interactions underlying this chiral induction is now in progress.

The first published interpretations $[178,179]$ about vortex-induced optical activity attributed this phenomenon to a cholesteric-like alignment of the particles


Figure 8.12.: Case of a CCW stirred solution.
in the stirring vortex analog to the twisted-crystal structure described in section 3.2.4. According to their interpretations the observed CD would be due to the macroscopic arrangement of the aggregates, but not due to structural changes in the interacting chromophores of the aggregates. This is an elegant explanation but it was published without any further demonstration. Even leaving aside the fact that it is questionable that a vortical stirring would result, in a square-section cuvette, to a perfect helical arrangement of the nanofibers in the light path as the one they described, we do not think that such arrangement is responsible for the measured signals. There exist a cumulative of reasons for this opinion:

Chapter 8. Chiral induction by hydrodynamic effects


Figure 8.13.: Optical effects calculated from the Mueller matrices.
i) A supposed vortex-induced macroscopic helical arrangement would have an approximate pitch of twice the dimensions of the cuvette ( 20 mm for the 10 mm pathlength cuvette used), which is several orders of magnitude greater that the wavelength of light. Therefore, from the point of view of the light beam it would be completely inhomogeneous medium. In contrast, the measured spectroscopic Mueller matrices do not have the more exotic profiles found in systems composed of a sequence of optical elements. In fact we have found that their interpretation according to the homogeneous model introduced in chapter 3 yields the expected simple and Kramers-Kronig consistent values of linear dichroism and linear birefringence of a molecular absorption band.
ii) The noise level of LD, LB, LD', LB' measurements suffers an important increase when switching from a stagnant measurement condition to a stirred situation. This is easily understandable as, immediately, when the stirring starts violent flows appear in the cuvette. The stirring mechanism supposes a new source of "noise" for the measurement of Mueller matrix, that acts by quickly changing the position and orientation of particles in solution. However, the noise


Figure 8.14.: Representation of the calculated axes of linear dichroism for the space-resolved measurements of the cuvette in stagnant and while stirring. The length of vectors is function of the magnitude of linear dichroism.
pattern of CD, although also shows some logic increase, seems to be much less affected by the mechanical stirring than those of linear effects. In case that the measured CD signals while stirring were result of an special helical arrangement of the fibers, one would expect noise levels associated to measured CD greater than the observed ones. Fig. 8.16 shows that this behaviour is also appreciable when comparing the standard deviations associated to the measurement of certain Mueller matrix elements. Experimental of Mueller matrix elements $m_{03}$, that for this system is mainly contributed by CD, and $m_{01}$, here dominated by LD, evolve to very different standard deviations when the stirring starts. While for $m_{01}$ the measured standard deviation becomes order of magnitudes greater, the changes in the standard deviation of $m_{03}$ are much more subtle.
iii) There is no reason to think that an helical arrangement is not possible in a cylindrical section cuvette. We think that in a cylindric flask such macroscopic helical arrangement could be more perfect than in the standard squared-section cuvettes. Therefore, if such macroscopic arrangement was responsible for the CD signal we should observe the same type of chiral induction in cylindrical flasks.

Another group [177] has recently pointed that the deposition of chiral aggregates on the cuvette walls during prolonged stirring could have significant contributions to the measured signals. While we agree that a deposition of a certain

Chapter 8. Chiral induction by hydrodynamic effects


Figure 8.15.: Superposition of the CD color map (already presented in Fig. 2) with a vectorial representation of in-plane projection of the linear dichroism.


Figure 8.16.: Comparison of the standard deviations associated to the measurement of the Mueller matrix elements $m_{01}$ and $m_{03}$ in stagnant and CCW stirring conditions. Element $m_{01}$ is mainly contributed here by LD and element $m_{03}$ is mainly contributed by CD.
amount of chiral aggregates in the cuvette walls can occur, non of our experiments indicate that deposited aggregates have significant contribution on the observed effect.

Rather than rigid structures, the J-aggregates of the studied porphyrins are soft-matter nanophases that have been demonstrated to be foldable by effect of hydrodynamic forces [181]. The elastic characteristics of the $\mathrm{H}_{4} \mathrm{TPPS}_{4}$ and $\mathrm{H}_{4} \mathrm{TPPS}_{3} \mathrm{~J}$-aggregates were studied by scanning probe microscopy techniques
yielding typical results for soft polymers and without significant differences both porphyrins [180]. The effect of mechanical forces on the conformational shape of macromolecules is well known. Gradients of shear rate and viscous drag forces acting on the moving particles exert a net torque on the particles that may lead to folding and twisting. This effect will be greater when the elastic modulus of the particles is low, i.e. the reversible stationary effect described in this thesis would be due to a pure mechanical effect upon elastic mesophases and the irreversible effect would be due to a "freezing".

The fact that can be surprising about the experiments reported in this chapter is that the deformation of a classical object (although nanosized) to a chiral shape is being detected at the level of the electronic transition ${ }^{2}$. The apparent size scale paradox between macroscopically induced hydrodynamic forces and the changes detected in the electronic transitions disappears when it is taken into account that the phenomenon of light absorption involves the interaction of the photon with an exciton of the solid (the nano- or microparticles in suspension). The exciton structure may depend on the distortion originated by the hydrodynamic forces, i.e. the modified excitonic absorption may be intrinsically chiral. According to our perspective, the understanding of the nanosized aggregates of the solution would be not so different to those of metallic macroscopic helices (a free electron in a helix) where optical activity had already been detected by the year 1920, $[185,186]$. In our opinion the mechanical effect that manifests in the induction of optical activity by stirring would be assimilable to the effect of stress in certain symmetry classes of achiral crystals that makes them exhibit optical activity, a phenomenon that is referred as piezo-rotation or piezo-gyration [64, 187]. As piezo-gyration can only occur for certain groups of symmetry, our assumption that mechanical stress leads to changes of the gyration tensor could explain our experimental results that show that the stirring effect is observed in solutions of certain porphyrin J-aggregates or RNTs but not in others of the same family, if inducible and non-inducible systems belonged to different symmetry point groups. Future work should be oriented to identify the symmetry changes that occur in the nanophases when they are bend and/or fold under gradients of shear rates of flows.

[^23]
## Chapter 9.

## Measurements on heterogenous solid-state samples

This chapter groups experiments carried out in solid heterogenous samples, for which the optical properties have been scanned thorough the surface.

### 9.1. Chiral domain in crystallizations

One of the most promising applications of Mueller matrix polarimetry is to investigate chiral domains in transparent solid crystalline samples. When the structure or composition of the sample varies from point to point, imaging or mapping techniques are the only way to discern the different optical effects that may be present in the sample.

Many of experiments involving the study the chiral symmetry breaking occurring at solidification processes have been performed in sodium chlorate $\left(\mathrm{NaClO}_{3}\right)$. One of the main reasons for this election is that sodium chlorate belongs to the cubic tetrahedral class, and, therefore, as it is isotropic, the posterior identification of crystallites of different handedness is very easy. One of the famous experiments with $\mathrm{NaClO}_{3}$ was done by Kondepudi [188], who noticed that the stirring of $\mathrm{Na}^{+}$and $\mathrm{ClO}_{3}^{-}$ions solution caused all the subsequent crystals to adopt the same chirality. More recently Viedma [189] presented, in a related experiment, that an initially equal mix of crystals in a supersaturated solution can achieve complete homochirality thanks to the action of grinding and stirring the crystals. With the experimental approach used in this thesis, we can go beyond cubic crystals and investigate other anisotropic solids that are of interest for the study of chiral symmetry breaking processes.

In the experiments of this section we studied achiral substances in the liquid phase, that, after solidification, yielded chiral enantiomorphs. The main difference with the experiments cited before is that, in our case, we were interested in systems made of one single chemical component ${ }^{1}$ so that from a thermodynamic point of view the problem can be merely presented as liquid-solid phase transition. As from an effective point of view, left-handed and right-handed enantiomers are

[^24]energetically identical and none of them should be preferred over the other the identification and study of possible chiral domains after crystallization may be in close correlation with symmetry breaking scenarios.

### 9.1.1. Benzil polycrystalline films

Benzil is a molecular crystal; its molecules do not exhibit any optical activity when they are in solution [190], but they crystallize in a chiral enantiomorphous group of symmetry $D_{3}^{4}$ or $D_{3}^{6}$ (the same group of symmetry than $\alpha$-quartz). The crystal of benzil is built upon a hexagonal lattice and the unit cell accommodates three molecules disposed spirally around the trigonal axis [191]. Therefore the benzil crystal is uniaxial and shows optical activity, which at visible regions of the spectra manifest not only in CB, but also in CD because benzil molecules have absorption bands.


Figure 9.1.: CD and CB of a benzil crystal for light propagating propagating parallel to the optical axis. These measurements were obtained from spectroscopic measurements of the Mueller matrix with the 2-MGE in a small benzil crystal grown by evaporation.

Fig. 9.1 shows the CD and CB spectra of a benzil crystal for light propagating parallel to the optic axis. A clear CD absorption band centered around 400 nm can be clearly seen, while at higher wavelengths the crystal shows no CD, but its optical activity can still be traced with CB. As it is common for anisotropic crystals (see the case of quartz in chapter 7), optical activity in benzil can be easily measured for light propagating along the optical axis, but in other directions is more difficult because linear birefringence and linear dichroism effects become much greater.

Benzil (1,2-diphenyl-1,2-ethanedione, Aldrich) was recrystallized twice from toluene, as means of purification, yielding six-sided prisms of about 5 mm in length. Later they were manually ground in an agate mortar to get a fine powder. $2-3 \mathrm{mg}$ of the powder were sandwiched between a blank microscope slide and a glass cover slips. The slide was raised to a temperature above the melting point of benzil (378K) in a hot stage, held for 15 minutes, and left to cool at room


Figure 9.2.: The picture on the left shows one of the studied thin polycrystalline film of benzil under standard illumination. The picture on the right was captured after putting the same sample between crossed polarizers.
temperature. A polycrystalline film is eventually developed from the metastable melt. Fig. 9.2 shows the type of samples we obtained after this preparation process.

Most of the surface of the sample was scanned by the 2-MGE using the assistance of quartz plates to maintain a high lateral resolution during the measurement. Imaging representations of the transmission Mueller matrix of the sample in regions around $10 \times 10 \mathrm{~mm}$ were obtained for several different samples (see one of them in Fig. 9.3). The pixel size used for generating these mapping plots is $60 \mu \mathrm{~m}$, which is slightly below the optical resolution of the measurement (85-90 $\mu \mathrm{m})$. In our case, this resolution is determined by the size of the light spot and the width and height of the monochromator slits placed before the detector.

Fig. 9.4 shows the $\mathrm{CD}, \mathrm{CB}$ and fraction $\beta$ of polarized light ( 1 for completely polarized and 0 for completely depolarized) calculated from the Mueller matrix in Fig. 9.3. Well delimited zones with different values of CD and CB are observed. In Fig. 9.5 the small area delimited by a black square in Fig. 9.4 is presented with more detail. In this figure the vectorial representation of the projections of linear birefringence, calculated with Eqs. (8.1) and (8.2) but, in this case, using LB and LB' instead of LD and LD', has been superposed to the color map of CD. This figure confirms that the boundaries between areas of positive and negative CD (mostly delimited by strait lines) do not correspond with changes on the orientation of the projected directions of the optic axis. Thus, zones in distinct false colour correspond to enantiomorphs otherwise in comparable optical orientations.

In Fig. 9.6 we show the positive and negative CD domains found experimental for several different benzil samples. In general the formation of well-defined and spatially segregated chiral domains can be appreciated. The boundaries between areas of opposite sign of CD in benzil are in almost all cases straight lines. These results were interesting for us because some theoretical works about dynamics of the Frank chiral amplification model that leads to homochirality already predict the formation of separate "colonies" of the two kinds of enantiomers bounded

Chapter 9. Measurements on heterogenous solid-state samples


Figure 9.3.: Mapping of the normalized Mueller matrix of a thin polycristaline film of benzil measured at 400 nm . It covers an area of $10 \times 10 \mathrm{~mm}$ with an step size of $60 \mu \mathrm{~m}$.
by racemic surfaces $[192,193]$. Such colonies could only survive if the two areas were connected by a strait. These works describe that any initial curvature in these boundaries would make the boundary move towards the side from which it is concave, and, as a consequence, one species will eventually shrink, and the other will survive, giving situations where only one of the enantiomers would be present (for example case c of Fig. 9.6 seems to be near to this situation).

### 9.1.2. Other polycrystalline films

Analogous experiments to those of benzil have been repeated with other organic compounds, in particular with benzophenone and binaphthyl. Thesse samples were prepared in a similar way to benzil thin films, but taking into account the heating/cooling process must be adjusted to the melting temperature of each substance. Nevertheless optical measurements in this compounds have carried


Figure 9.4.: Values of $\mathrm{CD}, \mathrm{CB}$ and $\beta$ obtained from the Mueller matrix in Fig. 9.3 using the inversion process of section 4.2. The black square indicates the area that is studied with more detail in Fig. 9.5.


Figure 9.5.: Superposition of the CD color map (squared area in Fig. 9.4) with a vectorial representation of in-plane projection of the linear birefringence, also calculated from the Mueller matrix.
many more problems than in the case of benzil. There are probably a variety of reasons for this different behavior; one of them is that this samples tend to show a more grained structure, where the crystallites appear to be very small


Figure 9.6.: Chiral domains obtained experimentally for several different benzil thin film samples. Colour red indicates positive CD, while negative values are represented in blue. Areas without a well-defined red or blue colour correspond to air blisters that were present in the samples. Case f) corresponds to the sample that has been presented in more detail in this section.
and, eventually, superposed one to another. All these structural facts probably lead to scattering phenomena or interface effects that complicate the optical measurement. For example Fig. 9.7 shows that the depolarization introduced by these samples is quite stronger than in the case of benzil, as it is revealed when we compare $\beta$ values, as calculated from Eq. 1.33 of Fig. 9.7 and Fig. 9.4. Another important issue is that these compounds do not have an accessible absorption band of CD in the visible as benzil (see Fig. 9.1). The identification of chiral domains in these compounds in the visible part of the spectra (which are the only wavelengths were the samples are transparent enough) should be done with CB , which usually is is more difficult to recognize than CD because CB is more masked by linear birefringence than CD.

Despite these difficulties, our intention is to pursue more experimental work in this topic in order to identify other solids or other preparation methods that permit extending the study we initiated with benzil to other organic compounds.


Figure 9.7.: Detected intensity and fraction of polarized light ( $\beta$ ) for crystallizations of binaphthyl and benzophenona. Binaphthyl sample was scanned at 390 nm aver an area of $12 \times 12 \mathrm{~mm}$, while benzophenona sample was scanned at 370 nm over an area of $6 \times 6 \mathrm{~mm}$. Round areas with fractions of polarized light close to 1 correspond to air blisters.

### 9.2. Meteorites

Although this section is situated at the end of this thesis, the measurements presented herein were chronologically obtained before most of the data shown in previous chapters. Also they are the only measurements presented in this thesis that were not obtained with the instrument described in Part III, that refers to the 2-MGE available in the facilities of the Faculty of Physics of the University of Barcelona. Instead, measurements of this section were also performed with a transmission 2-MGE, but this time with one of the instruments built and kept by Gerald. E. Jellison at the Oak Ridge National Laboratory (Tennessee, EEUU).

In contrast to previous chapters, the conclusions we will reach in this section are somewhat speculative, mostly because the difficulty associated to meteorite
samples is beyond our current technical capabilities for the analysis of complex samples with polarimetry. Therefore, this section, rather than offering conclusive results it opens new perspectives for original and unprecedented work in polarimetry. The contents of this section are a summary of the results published in [194].

### 9.2.1. Motivation

The introduction here of our work on meteorites here may sound a little awkward, but it supposes a continuation of our the work about the study of processes of chiral induction that can act upon an specific chemical process showing spontaneous mirror symmetry breaking we have carried in this thesis. Furthermore the investigation reported here could be be included in the more general topic of the understanding of the chemical evolution towards biological homochirality (see section 2.3). In fact, strictly speaking, our more evident objective here was almost the same that in the previous section: the identification of chiral bias and chiral domains in a solid samples. The major difference is that instead of using artificially-prepared samples in the laboratory, we have to deal with natural solid samples with much more complex characteristics.

We started the work on meteorites, or more precisely in carbonaceous chondrites ${ }^{2}$ with the hypothesis that the action of shear force gradients during the accretion of materials and solidification processes during the formation of planetesimal bodies and comets could lead to a chiral bias (enantiomeric excess) of chiral fractures, screw distortions or step-kink chiral centers on the surfaces of inorganic matrices, which could later transfer a chiral bias to organic reactions.

Carbonaceous chondrites are the remnants of first chemical stages where chiral sign selection and chirality amplification could have occurred. This is suggested in $\mathrm{CM} 2^{3}$ meteorites by the presence of significant chiral organic compounds $[195,196]$ and principally because a real chiral bias has been detected in aminoacids [37, 197, 198] and organic kerogenic materials [199]. Further, in CM chondrites inorganic matrices as those proposed in some abiotic reaction scenarios are ubiquitously present in CM chondrites. Many studies of CM2 reported the presence of products of aqueous alteration, e.g. phyllosilicates, namely Fe-rich serpentines and poorly characterized phases (PCPs) in chondrules, aggregates, and within fine-grained rims.

[^25]
### 9.2.2. Experiment

Meteorite samples, or, more in general petrological samples, represent one of most difficult types of samples one can investigate in polarimetry. They usually are heterogeneous even at a microscopic scale, which means that the optical characteristics of two near points, even if they are separated only by a few microns, may be different. Also there are difficulties associated to the manipulation and cut of the rocks, in order to obtain specimens thin enough to have the level of transparency enough to allow some light reach the detector.

Measurements were carried out with a transmission 2-MGE that shares the fundamentals indicated chapter 5 . The method involving quartz rotators described in Chapter 6 was not applied to these measurements because, chronologically it was developed later. In this work we were mainly interested in the measurement of CB which, assuming that the measured Mueller matrix corresponds to the Mueller matrix of a retarder (Eq. (3.24)) can be obtained from the difference between the difference between $m_{12}$ and $m_{21}$ matrix elements, as shown in Eq. (3.46a). Therefore our main objective here was the determination of $m_{12}$ and $m_{21}$. As the 2-MGE is not able to measure simultaneously $m_{12}$ and $m_{21}$ for any relative orientation of the PSG and PSA (see Eqs. (5.14)) we addressed this issue by placing two zero-order quarter wave plates (QWP) tuned to the measuring wavelength ( 546 nm ) just before and after the sample that permit to select which 8 elements were measured simultaneously. The details about the QWP method we used are available in Ref. [144], we do not devote more attention to them in this thesis, because in case we were able to repeat the measurement right now we would follow the quartz rotators technique described in Chapter 6

The measurements were performed on three mineralogic pristine cuts ( $\sim 30 \mu \mathrm{~m}$ thickness) of the CM2 meteorites Murchison (ASU828-A-1), Murray (ASU635-A-3) and Cold-Bokkeveld (ASU578-A-2) on loan from the Center for Meteorite Studies of the Arizona State University.

### 9.2.3. Results

Characteristic images of the CB mapping of portions of the meteorite thin sections surface calculated according to $\mathrm{CB} \sim\left(m_{12}-m_{21}\right) / 2$ are shown in the series b of Figs. 9.8, 9.9 and 9.10. Most of the transparent area do not show CB, however, at specific sites of the sample positive and negative CB values are detected. The statistical errors associated to the experimental measurement in the points with zero CB values should, in principle, give a Gaussian distribution centered at zero. Points with CB should give a bimodal distribution at positive and negative values and a chiral bias should be indicated by a distribution different to $50 \%$ for both signs. The series c of Figs. 9.8, 9.9 and 9.10 show the distribution of CB values for the MU, MR and COB thin sections. We have excluded
from this statistic points which exhibited a percent transmittance below $0.65 \%$, which corresponds to opaque points in which the measurement has a poor signal to noise ratio (darker points of series a of the Figs. 9.8, 9.9 and 9.10). Also very transparent points (percent transmittance above $20 \%$ ) which are, most of them, attributable to areas out the meteorite sections were not considered in the recount.


Figure 9.8.: Measurements of a section of a Murchison thin cut ( $\sim 30 \mu m$ thickness); a) optical density measured at 546 nm ; b) CB mapping of a) (the dimension of each frame is $12 \mathrm{~mm} \times 6 \mathrm{~mm}$, where each pixel measures $30 \mu \mathrm{~m} \times 30 \mu \mathrm{~m}$ ); c) Distribution of the CB values of the image pixels in b), once eliminated the non-transparent and the more transparent pixels; $d$ and e SEM images of specific areas (marked in b) and EDX composition analysis .

The results of the Murray sample sample (Fig. 9.9) indicate the absence of a significant sign bias in the CB. In contrast the Murchison sample (Figs. 9.8 and 9.11 ) shows in addition to the Gaussian distribution value around zero a bimodal distribution with a clear bias towards negative CB values. In fact the Murray sample probably also shows an additional bimodal distribution with a similar


Figure 9.9.: Measurements of a section of a Murray thin cut ( $\sim 30 \mu m$ thickness); a) optical density measured at 546 nm ; b) CB mapping of a) (the dimension of each frame is $4 \mathrm{~mm} \times 4 \mathrm{~mm}$, where each pixel measures $20 \mu m \times 20 \mu \mathrm{~m}$ ); c) Distribution of the CB values of the image pixels in b), once eliminated the non-transparent and the more transparent pixels.


Figure 9.10.: Measurements of a section of a Cold-Bokkeveld thin cut ( $\sim 30 \mu m$ thickness); a) optical density measured at 546 nm : b) CB mapping of a) (the dimension of each frame is $2 \mathrm{~mm} \times 2 \mathrm{~mm}$, where each pixel measures $20 \mu \mathrm{~m} \times$ $20 \mu \mathrm{~m})$; c) distribution of the CB values of the image pixels in b, once eliminated the non-transparent and the more transparent pixels
bias than that of the Murchison sample, but of low significance which could be also attributed to experimental errors. The presence of a biased CB distribution is clearly observed in the Cold-Bokkeveld sample (Fig. 9.10), however, in this last case the number of measured points with good transparency values is low for a chiral bias estimation, but it shows clearly the bimodal distribution of CB values.

In order to identify the materials corresponding to the areas of the Murchison meteorite that showed optical activity, the thin sections were carefully analyzed by SEM-EDX. The series d and e of Fig 9.8 constitute a representative example of the composition of the areas exhibiting strong optical activity. The results indicate that CB signals are not originated by intrinsically chiral crystals such as heptahydrated sodium sulfate or quartz crystals. Specifically, CB signals are

## Chapter 9. Measurements on heterogenous solid-state samples



Figure 9.11.: Distribution of the CB values of other sections of the Murchison thin cut than that of Figs. 9.8, once eliminated the non-transparent and the more transparent pixels, showing in all cases the bias towards negative CB values.
originated in phyllosilicates sites and poorly characterized phases ${ }^{4}$. Although CB active sites coincide with the location of serpentines (phyllosilicate structures of the meteorite where aminoacids have been detected), it is not reasonable to assume that the relatively high CB contributions could arise from the low enantiomeric excesses of organic chiral compounds, which commonly show low CB values at the visible wavelength region. The distribution of chirality of microsites of different sign with a statistical chiral bias is a chemical reasonable scenario, which agrees with the strong heterogeneous structure of carboneaceous chondrites and with the different distribution of enantiomeric excesses of chiral aminoacids in different samples of CM2 Pizzarello [37, 199].

We think that is likely that strong shear gradients could originate a surface chiral bias by the formation of screw chiral distortions and step-kinks by the effect of flow shear gradients. For example, it is known that hydrothermal chrysotile, which shows similarities with the serpentines identified some meteorites [200], forms chiral nanotubes by the curling to spirals of the crystalline flakes of the phylosilicate bilayered structure. Moreover, the effect of mechanical forces on the structure of these phyllosilicates it probably significant; for instance, it has been previously reported that grinding on phylosilicates and specifically on serpentines originate transitions from crystalline to amorphous phases by the creation of structural defects [201]. The obtained results, albeit preliminary, are not in

[^26]contradiction with our hypothesis and should be considered as a stimulus for further work on the topic.

In this section we have presented a first experimental approach to the study of the chiral properties of solid state meteorite samples. As commented above these measurements involve an important technical complexity together with a higher difficulty than encountered by any previous work done in this field. The experimental procedure presented constitutes a good starting point and describes the problems that need to be considered and improved in further research. It should be noticed that, in addition to the feasibility to obtain results concerning to the detection of chirality, the used methodology may be a new tool for the petrographical characterization of meteorites.

## Chapter 10.

## Conclusions

The conclusions of this thesis have been grouped in three categories that correspond to the three parts in which the main contents of this work have been divided: theory (Part II), experimental sets (Part III) and measurements (Part IV).

### 10.1. Theory

The constitutive relations of an optically active (gyrotropic) and anisotropic medium are considerably more complex than those of a non-gyrotropic medium and, consequently, the resolution of the Maxwell equations with gyrotropic constitutive relations supposes a great added difficulty with respect to the nongyrotropic case. In practice, the solutions of Maxwell equations for a gyrotropic medium can only be handled numerically and, therefore, it is difficult to construct a model based on the dielectric and gyration tensors of the medium that describes, in terms of the Jones or Mueller matrices, the interaction of light with anisotropic chiral media.

1. The infinitesimal matrix representation of an optical system introduced by Jones permits the obtention of an analytic Jones or Mueller-Jones matrix. This matrix describes light transmission through any homogenous anisotropic chiral medium as a function of the thickness-dependent optical effects of the medium. This method has the limitation that it does not take into account the effect of the interfaces between different media. However, we have shown that for light at normally incident onto a bulk anisotropic sample, the change in the polarization occurring only at the interfaces is usually small enough to be safely omitted.
2. In contrast to what some authors have suggested, from the work of R.C. Jones it is not possible to infer the existence of any new type of birefringence or dichroism (which were eventually known as Jones birefringence and Jones dichroism in the literature) independent from the common birefringences or dichroisms. Jones used for his optical calculus two different in-plane projections of linear birefringence ( $\mathrm{LB}, \mathrm{LB}^{\prime}$ ) and linear dichroism ( $\mathrm{LD}, \mathrm{LD}^{\prime}$ ), but they do not identify different physical phenomena.
3. We have introduced the analytic equations that permit the inversion of the most general Jones or Mueller-Jones matrix obtained with the Jones infinitesimal formalism. These equations can be applied to the interpretation of experimental Mueller matrices if they do not include depolarization effects.
4. Mueller matrix product decompositions can be used to factorize experimental matrices into more simple terms. This can serve for the interpretation of the experimental Mueller matrices, and they are specially interesting for depolarizing media. However, product-decompositions are order-dependent, i.e., they also depend on the order in which the matrix factors are multiplied, which, in practice, means that not always the optical properties of original Mueller matrix can be derived from the optical properties of the individual factors. In the case of the polar decomposition we have derived a set of conditions that, if fulfilled, assure the commutativity of the diattenuating and retarding factors of the decomposition. They can be used as a criteria for the applicability of the polar/Lu-Chipman decompositions.
5. A new product decomposition, which we have named "pseudopolar decomposition", has been proposed. It is a product factorization that yields a diattenuating factor, a retarding factor and infinite succession of correction terms than account for the non-commutative optical properties of the medium. In case of media with commutative optical effects the pseudopolar decomposition simplifies to the polar decomposition.

### 10.2. Experimental sets

The time-dependent intensity generated by the 2 -MGE is complicated and can be written as a Fourier expansion with an infinite number of terms of different frequency. However, it can be described by a dc term and eight independent functions allowing to determine eight different elements of the Mueller matrix of the sample in a given configuration. The choice of the measured Mueller matrix elements is entirely determined by the azimuthal angles of the PSG and the PSA. A complete Mueller matrix determination requires four measurement configurations.
6. A spectroscopic 2-MGE, a type of polarimeter based on the use of two free runing PEMs operating at its own distinct frequencies, has been entirely built. It has been designed to be used either in transmission or in reflection, and in both cases the angle of incidence can be adjusted. Acquisition and calibration routines for the different measurement modes, together with a complete graphical user interface, have been programmed to operate the instrument.
7. We have demonstrated the usefulness of the 2-MGE to investigate optical activity in a wide spectral range from UV, to near-IR. The use of PEMs as elements with variable retardance allows a high purity and efficiency in the modulation that, together with the synchronous signal acquisition, is the basis for the high sensitivity and high resolution that the measurements of optical activity requires.
8. We have developed a novel technique that can expand and improve the capabilities of $2-\mathrm{MGE}$ or other types of ellipsometers/polarimeters. This new experimental proposal is based on the use of an element with optical rotation as a substitute of the precision mechanical rotation stages that are typically used to change the azimuthal orientation of the PSG and the PSA. This technique was conceived to solve the beam translation problem ocurring when the azimuthal angles of the PSG and PSA are changed.
9. The optical rotation method has been put in practice in the 2-MGE by using two thin $z$-cut $\alpha$-quartz plates as elements with rotatory power. With the assistance of the quartz plates, the $2-\mathrm{MGE}$ is able to measure all the elements of the normalized Mueller matrix while keeping the PSG and PSA static. The four different measurement configurations needed to measure a Mueller matrix with a 2 -MGE are reached by selectively introducting/removing the quartz elements present at both arms of the instruments. Besides spectroscopic measurements, the quartz elements have permitted us to obtain spatially resolved measurements of the Mueller matrix that were impossible with the mechanical rotators because of the beam translation problem. Calibration routines for the use of quartz-plates have also been built.

### 10.3. Experiments

The determination of optical activity supposes in many occasions an experimental challenge. Optical activity signals arising at molecular and supramolecular levels, of interest in chemistry, tend to be so small that they can be below the resolution of many polarimeters or ellipsometers. In anisotropic crystals, optical activity may be even more difficult to detect, because their linear birefringence is orders of magnitude greater, and hides the effect of optical activity, despite they can exhibit much intense optical activity than molecular non-crystalline substances.
10. We have conducted the first experimental determination of the gyration tensor of a crystal ( $\alpha$-quartz) based on spectroscopic Mueller matrix measurements. We faced the long-standing problem associated to the determination of optical activity of quartz in birefringent directions of the crystal,
in order to be able to measure the $g_{11}$ component of the gyration tensor. The experimental approach we have used is based on the obtention of the transmission Mueller matrix measurements of $z$-cut quartz plates at oblique angles of incidence. The same methodology could be used to measure components of the gyration tensor of other crystals.
11. Spectroscopic Mueller matrix measurements have been applied to the study the induction and switching of chirality in solutions of certain soft-matter nanoribbons under the effect of a stirring vortex. We have shown that the handedness of the induced optical activity depends on the direction of stirring in a completely reversible process, which supposes a modification of the supramolecular chirality of the nanophases.
12. Spatially resolved measurements of the Mueller matrix in square section cuvettes containing these solutions have revealed that the induction of optical activity takes place only in the central part of the cuvette, and have permitted to establish a spatial correlation between chiral flows and the emergence of supramolecular chirality. Our experiments suggest that the measured supramolecular chirality arises due to a bending/folding of the nanoribbons under chiral hydrodynamic force. Such mechanical effect would be produced by the flow shear gradients and the observed phenomenon could be described as piezorotation. We have also found out the importance of the design of stirred containers to optimize such chiral inductions.
13. The obtention of measurements of optical activity with spatial resolution and in anisotropic samples represents an important step forward towards the characterization of chiral materials. To our knowledge this analysis has not been carried out with any other technique. We have used it to examine chiral domains in thin polycrystalline solidifications of benzil from a melt. Domains corresponding to different crystals enantiomorphs have been unambiguously identified through measurement of CD and CB. It has been verified that the boundaries between different domains are straight lines.
14. We have presented a first experimental approach to the study of the chiral properties of solid state meteorite samples. These measurements involve an important technical complexity because the samples are highly heterogenous, and tend to be opaque. Differences in the optical response of samples corresponding to different meteorites have been observed, and a chiral bias has been glimpsed in a Murchison meteorite sample. The experimental procedure presented constitutes a good starting point and describes the problems that need to be considered and improved in further research.

### 10.4. Future perspectives

In this final section we discuss, without a particular order, several directions in which the work presented in this thesis could be extended. Some of them are topics that are already under investigation at the moment of writing this section.

One of the main areas that we think that still requires an intense study is the measurement of optical activity in anisotropic samples in reflection. It would require a quite different vision from transmission studies, because while transmission measurements are based on long path-lengths measurements and they can be described by a set of "thickness-dependent" parameters, in reflection, optical activity is purely an interface effect. This is a quite virgin area of research, where only a small number of measurements in isotropic optical active samples have been reported [202, 203]. However if appropriate models are constructed (probably using the Berreman formalism), the analysis of anisotropic chiral samples in reflection should be feasible. It would be really worthy to start a two-sided, theoretical and experimental, work to examine in which kind of samples (if any) analyzing the reflected beam (instead of the transmitted beam, as is has been done in this thesis) would be better for the characterization of optical activity.

In our opinion, many of the more interesting future applications of Mueller matrix polarimetry would require having instruments with imaging capabilities. Having a Mueller polarimeter acting as a high resolution optical microscope would nearly be a dream made true. In fact we are not so far from this objective because there are many recent works $[67,204]$ (and even the instrumental part of this thesis) that propose experimental setups going in this direction. The main issue here is that, while getting non-precise, nearly qualitative, measurements of the Mueller matrix of a sample is relatively easy from an experimental point of view, applications that require a high accuracy, precision and resolution, such the measurement of optical activity, need a much higher experimental effort that is not always compatible with the requirements imposed by microscopy. For example, the photoelastic modulators we employ are an excellent source of phase modulation, but at the moment, due to speed mismatches, are not compatible with CCD detectors usually employed in microscopy.

A really appealing apparatus to develop, and the "natural" continuation of the instrumental work presented in this thesis would be a polarimeter/ellipsometer running with 4 PEMs at different resonant frequencies (2 of them in then PSG and the other 2 in the PSA). This instrument would be able to measure all the elements of the Mueller matrix in a single measurement and in a completely static setup. Another interesting objective to pursue would be extending the optical activity measurements on the Mueller matrix determination to other wavelength regions. In particular, it would be very attractive to go to the IR because it would permit the measurement of optical activity associated with molecular vibrations in anisotropic samples. Some experimental work aimed at determining
the Mueller matrix in the IR domain is planned in our research group.
The construction and tuning of the 2-MGE took to the author of this thesis more time than any other topics discussed in this work. Considerable efforts were also put in the development of theoretical tools for the analysis of experimental data. At the moment these instrumental and theoretical tools have been exploited for a restricted number of samples. We are aware that there is an almost endless list of chiral samples susceptible to be studied by Mueller polarimetry. A very clear example are chiral crystals, where there are many gyration tensors waiting to be measured. Besides the measurements on quartz presented in this thesis, we have also worked (and still there is some work in process) in the measurement of the gyration tensor of benzil and $\mathrm{AgGaS}_{2}$. Anyway, it must be remarked that having a set of theoretical and experimental tools does not mean that the analysis of a new chiral sample can be done on a routine basis; but rather each sample supposes a new challenge.

Going to more particular applications, there are several fields that we think that could benefit from optical activity measurements pioneered in this thesis Liquid crystals is one of the main research areas where we believe that our tools could have a more straightforward application. The study of the chiroptical properties of liquid crystals represents a vast area of investigation and it also attracts the interest from the industry. Many more applications could located in the biosciences, where the there are lots of biopolymers, for example DNA, with chiral anisotropic structures. Another novel and very active area of research that demands of advanced optical activity measurements is nanotechnology. For example, in the characterization of chiral photonic crystals or for a further understanding of the interesting optical properties of 2D and 3D chiral periodic structures in the sub-wavelength scale known as metamaterials.

Part V.
Appendices

## Appendix A.

## Resum en català

## A.1. Introducció

L'objectiu d'aquesta tesi és estudiar la quiralitat de medis anisòtrops mitjançant polarimetria de matriu de Mueller. Per arribar a aquest objectiu haurem de tractar aspectes diversos i en bona mesura multidisciplinaris, però tots ells importants per a la consecució d'aquesta tesi. Per exemple, veurem formalismes per la descripció de la propagació de la llum polaritzada per aquest tipus de medis, però també ens fixarem en qüestions plenament instrumentals com ara la construcció d'un polarímetre o el • lipsòmetre de matriu de Mueller, i, finalment, buscarem una interpretació dels nostres resultats en aspectes de química supramolecular.

L'activitat òptica és la manifestació de com interaccionen els medis quirals amb la llum polaritzada. Va ser observada per primer cop pel astrònom François Arago (en català Francesc Aragó) quan va descriure una rotació en la direcció de polarització quan llum polaritzada linealment travessava un cristall de quars al llarg del seu eix òptic. En l'actualitat, gairebé 200 anys després d'aquest descobriment, els científics coneixen moltes substàncies i materials que posseeixen activitat òptica i la seva mesura ha esdevingut d'una importància essencial en molts àrees de recerca: química, biologia, ciència de materials, farmàcia, etc.

Els primers treballs experimentals sobre activitat òptica estaven basats en la mesura de la rotació de la direcció de polarització quan llum polaritzada linealment travessava alguns medis quirals. Aviat es va descobrir que aquesta rotació òptica, referida a vegades també com poder rotatori, es podia explicar per una diferencia en els índex de refracció del medi per llum polaritzada circularment cap a la esquerra o cap a dreta, una diferència que es coneix habitualment com birefringència circular ( $\mathrm{CB}^{1}$ ). No va ser fins dècades més tard, després dels treballs d'Aimé Cotton, que va començar la mesura de les diferències d'absorció per llum polaritzada circularment cap a l'esquerra i per llum polaritzada circularment cap a la dreta, que es coneix amb el nom de dicroisme circular.

Malgrat que en l'actualitat el fenomen de l'activitat òptica és àmpliament conegut, la seva mesura encara presenta dificultats pels científics, especialment

[^27]en medis anisòtrops. Una raó és que els efectes de CB i CD són en general petits, i, a més a més, en medis anisòtrop hi ha birefringència lineal i/o dicroisme lineal que són ordres de magnitud més gran i emmascaren l'efecte de CB i CD. Un exemple de gran importància és la mesura de $C B$ en un cristall per direccions diferents de l'eix òptic, ja que la petita pertorbació que causa la CB sobre la polarització de la llum, en comparació amb el gran pertobació causada per la birefrigència lineal, és difícilment detectada des d'un punt de vista experimentals. Els aparells comercials disponibles en l'actualitat no es poden usar per mesurar CD i/o CB mostres anisòtropes, perquè estan dissenyats sota la premissa que el medi estudiat és òpticament isòtrop, és a dir, que té les mateixes propietats òptiques en totes les direccions.

Mitjançat una tècnica com la polarimetria es pot mesurar experimentalment CD i CB. Per medis isòtrops els instrument destinats a la mesura de CB son simples i estan constituïts únicament per una font de llum i dos polaritzadors, entre els quals es situa la mostra. Pel cas CD l'aparell haurà de disposar també d'algun element retardador o compensador en el camí òptic. Quan es tracta de medis anisòtrops veurem que la mesura de CD i CB esdevé més complexa i, idealment, s'haurà de basar en la determinació de la matriu de Mueller mostra.

## A.2. Propagació de la llum en medis òpticament actius $\mathbf{i}$ anisòtrops

La descripció de la propagació de la llum per medis òpticament actius i anisòtrops no està exempta de dificultats. La teoria electromagnètica es va fonamentar degudament a partir de la formulació de les equacions de Maxwell, però tot i això, Maxwell no va considerar els medis quirals. No s'ha trobat una solució analítica per a la resolució de les equacions de Maxwell emprant equacions constitutives més generals d'un medi anisòtrop i quiral i el problema s'ha de tractar numèricament.

Per facilitar la resolució de les equacions en medis anisòtrops s'han desenvolupat alguns mètodes matricials que permeten sistematitzar els càlculs i, a l'hora, asseguren el compliment de les condicions de contorn lligades a les interfícies de canvi de medi. El mètode més conegut és el desenvolupat per Berreman [69], que ha estat àmpliament difós dins l'àmbit de l'el • lipsometria, ja que és fàcilment adaptable a sistemes multicapa. Malgrat aquests avantatges tampoc amb aquest mètode es poden trobar equacions analítiques i generals que descriguin con evoluciona la polarització de la llum quan travessa un medi quiral i anisòtrop.

En aquesta tesi farem servir un mètode alternatiu de tipus semi-fenomenològic que no requereix considerar les equacions constitutives del medi. Aquest mètode va ser introduït per R. C. Jones en l'àmbit del seu formalisme per descriure llum polaritzada [75]. Es basa en una representació infinitesimal del medi a partir
dels vuit variable necessàries per descriure el comportament de qualsevol sistema òptic complex que no introdueixi despolarització ${ }^{2}$ [76]. Aquestes vuit variables es defineixen com:

- retard isòtrop de fase,

$$
\begin{equation*}
\eta=\frac{2 \pi}{\lambda}\left(n_{x}+n_{y}\right) l \tag{A.1a}
\end{equation*}
$$

- retard isòtrop d'amplitud,

$$
\begin{equation*}
\kappa=\frac{2 \pi}{\lambda}\left(k_{x}+k_{y}\right) l, \tag{A.1b}
\end{equation*}
$$

- projecció horitzontal del dicroisme lineal,

$$
\begin{equation*}
\mathrm{LD}=\frac{2 \pi}{\lambda}\left(k_{x}-k_{y}\right) l, \tag{A.1c}
\end{equation*}
$$

- projecció horitzontal de la birefringència lineal,

$$
\begin{equation*}
\mathrm{LB}=\frac{2 \pi}{\lambda}\left(n_{x}-n_{y}\right) l, \tag{A.1d}
\end{equation*}
$$

- projecció a $45^{\circ}$ del dicroisme lineal,

$$
\begin{equation*}
\mathrm{LD}^{\prime}=\frac{2 \pi}{\lambda}\left(k_{45}-k_{135}\right) l \tag{A.1e}
\end{equation*}
$$

- projecció a $45^{\circ}$ de la birefringència lineal,

$$
\begin{equation*}
\mathrm{LB}^{\prime}=\frac{2 \pi}{\lambda}\left(n_{45}-n_{135}\right) l, \tag{A.1f}
\end{equation*}
$$

- dicroisme circular,

$$
\begin{equation*}
\mathrm{CD}=\frac{2 \pi}{\lambda}\left(k_{-}-k_{+}\right) l, \tag{A.1g}
\end{equation*}
$$

- birefringència circular,

$$
\begin{equation*}
\mathrm{CB}=\frac{2 \pi}{\lambda}\left(n_{-}-n_{+}\right) l \tag{A.1h}
\end{equation*}
$$

[^28]on $n$ indica l'índex de refracció, $k$ és el coeficient d'extinció, $l$ és el camí òptic dins del medi i $\lambda$ és la longitud d'ona en el buit. Els subíndex especifiquen la polarització de la llum segons $x, y$, a $45^{\circ}$ de l'eix $x$, a $135^{\circ}$ de l'eix $x$, circularment cap a la dreta + , o cap a l'esquerra - És interessant remarcar que LB i $\mathrm{LB}^{\prime}$ (de la mateixa manera que LD i $\mathrm{LD}^{\prime}$ ) no descriuen efectes físics diferents, sinó diferents projeccions de la birefringència (o del dicroisme) [96].

Aquest mètode, almenys en la forma en que es presenta en aquesta tesi, només està descrit per llum polaritzada en incidència normal sobre la mostra, i no té en compte les reflexions que puguin ocórrer a les interfícies de la mostra. Tot i aquestes limitacions, es tracta d'un sistema molt útil per estudiar la transmissió de llum a través d'un medi homogeni, anisòtrop i amb activitat òptica, ja que empra àlgebra senzilla i ofereix resultats analítics que són fàcilment relacionables amb l'experiment.

La matriu de Jones J que descriu la transmissió de llum a través de qualsevol tipus de medi homogeni i que no introdueixi despolarització es pot obtenir a partir de l'exponencial [78]:

$$
\begin{equation*}
\mathbf{J}(\omega, z)=\exp [z \mathbf{N}(\omega)] \tag{A.2}
\end{equation*}
$$

on $\omega$ és la freqüència, $z$ és la distància dins del medi $(z=0$ indicaria el punt de la interfície). La matriu $\mathbf{N}$ és el generador infinitesimal de $\mathbf{J}$, i satisfà:

$$
\begin{equation*}
\mathbf{N}(\omega)=\lim _{z \rightarrow 0} \frac{\mathbf{J}(\omega, z)-\mathbf{I}}{z} \tag{A.3}
\end{equation*}
$$

$\mathbf{N}$ és una matriu infinitesimal que es pot construir a partir d'una versió infinitesimal de les variables definides a les Eqs. (A.1). La matriu de Jones de l'Eq. (A.2) es pot calcular com [75,170]:

$$
\mathbf{J}=e^{-i \chi / 2}\left(\begin{array}{cc}
\cos \frac{\mathrm{T}}{2}-\frac{i \mathrm{~L}}{\mathrm{~T}} \sin \frac{\mathrm{~T}}{2} & \frac{\left(\mathrm{C}-i \mathrm{~L}^{\prime}\right)}{\mathrm{T}} \sin \frac{\mathrm{~T}}{2}  \tag{A.4}\\
-\frac{\left(\mathrm{C}+i \mathrm{~L}^{\prime}\right)}{\mathrm{T}} \sin \frac{\mathrm{~T}}{2} & \cos \frac{\mathrm{~T}}{2}+\frac{i \mathrm{~L}}{\mathrm{~T}} \sin \frac{\mathrm{~T}}{2}
\end{array}\right),
$$

on hem definit un retard complex per cada parella de birefringència i dicroisme:

$$
\begin{gather*}
\chi=\eta-i \kappa  \tag{A.5}\\
\mathrm{~L}=\mathrm{LB}-i \mathrm{LD}^{\prime}  \tag{A.6}\\
\mathrm{L}^{\prime}=\mathrm{LB}^{\prime}-i \mathrm{LD}^{\prime},  \tag{A.7}\\
\mathrm{C}=\mathrm{CB}-i \mathrm{CD}, \tag{A.8}
\end{gather*}
$$

and $\mathrm{T}=\sqrt{\mathrm{L}^{2}+\mathrm{L}^{\prime 2}+\mathrm{C}^{2}}$.

La matriu de Mueller $\mathbf{M}$ corresponent a la matriu de Jones ${ }^{3}$ de l'Eq. (A.4) es pot calcular a partir de la relació [3]:

$$
\begin{equation*}
\mathbf{M}=A\left(\mathbf{J} \otimes \mathbf{J}^{*}\right) A^{-1}, \tag{A.9}
\end{equation*}
$$

on $\otimes$ indica un producte de Kronecker i $A$ és

$$
A=\frac{1}{\sqrt{2}}\left(\begin{array}{cccc}
1 & 0 & 0 & 1  \tag{A.10}\\
1 & 0 & 0 & -1 \\
0 & 1 & 1 & 0 \\
0 & i & -i & 0
\end{array}\right)
$$

La matriu de Mueller resultant es pot calcular doncs analíticament [41, 170] i l'hem presentada a l'Eq. (3.21) d'aquesta tesi.

## A.3. Inversió i descomposició de les matrius de Mueller

En aquesta secció ens interessarem en l'obtenció dels paràmetres òptics rellevant definits a l'apartat anterior a partir d'una matriu de Mueller experimental. Distingirem dos mecanismes, un consistent en la inversió analítica de la matriu de Jones o de Mueller-Jones descrita a l'apartat anterior, i l'altre basat en descomposicions algebraiques de la matriu de Mueller.

## A.3.1. Inversió analítica d'un matriu de Mueller-Jones corresponent a un medi homogeni

Per conveniència expressarem els elements d'una matriu de Jones en forma polar:

$$
\mathbf{J}=e^{i \theta_{00}}\left(\begin{array}{cc}
r_{00} & r_{01} e^{i\left(\theta_{01}-\theta_{00}\right)}  \tag{A.11}\\
r_{10} e^{i\left(\theta_{10}-\theta_{00}\right)} & r_{11} e^{i\left(\theta_{11}-\theta_{00}\right)}
\end{array}\right)
$$

Per tal de determinar LB, LB', CB, LD, LD' and CD a partir d'una matriu de Jones expressada segons l'Eq. (A.11) es poden emprar les següents equacions [99]:

$$
\begin{gather*}
\mathrm{LB}=\operatorname{Re}\left[i \Omega\left(r_{00}-r_{11} e^{i\left(\theta_{11}-\theta_{00}\right)}\right)\right]  \tag{A.12a}\\
\mathrm{LB}^{\prime}=\operatorname{Re}\left[i \Omega\left(r_{01} e^{i\left(\theta_{01}-\theta_{00}\right)}+r_{10} e^{i\left(\theta_{10}-\theta_{00}\right)}\right)\right],  \tag{A.12b}\\
\mathrm{CB}=\operatorname{Re}\left[\Omega\left(r_{01} e^{i\left(\theta_{01}-\theta_{00}\right)}-r_{10} e^{i\left(\theta_{10}-\theta_{00}\right)}\right)\right],  \tag{A.12c}\\
\mathrm{LD}=-\operatorname{Im}\left[i \Omega\left(r_{00}-r_{11} e^{i\left(\theta_{11}-\theta_{00}\right)}\right)\right], \tag{A.12d}
\end{gather*}
$$

[^29]\[

$$
\begin{align*}
\mathrm{LD}^{\prime} & =-\operatorname{Im}\left[i \Omega\left(r_{01} e^{i\left(\theta_{01}-\theta_{00}\right)}+r_{10} e^{i\left(\theta_{10}-\theta_{00}\right)}\right)\right]  \tag{A.12e}\\
\mathrm{CD} & =-\operatorname{Im}\left[\Omega\left(r_{01} e^{i\left(\theta_{01}-\theta_{00}\right)}-r_{10} e^{i\left(\theta_{10}-\theta_{00}\right)}\right)\right] \tag{A.12f}
\end{align*}
$$
\]

amb

$$
\begin{equation*}
K=\left[r_{00} r_{11} e^{i\left(\theta_{11}-\theta_{00}\right)}-r_{01} r_{10} e^{i\left(\theta_{01}-\theta_{00}\right)} e^{i\left(\theta_{10}-\theta_{00}\right)}\right]^{-1 / 2} \tag{A.13}
\end{equation*}
$$

i on $\Omega=\mathrm{T} K /[2 \sin (\mathrm{~T} / 2)]$ i $\mathrm{T}=2 \cos ^{-1}\left[K\left(r_{00}+r_{11} e^{i\left(\theta_{11}-\theta_{00}\right)}\right) / 2\right]$. Els símbols Re i Im indiquen, respectivament, la part real i la part imaginària.

Les Eqs. (A.12) permeten invertir d'una manera directa i ràpida una matriu de Jones. Les mateixes equacions es poden fer servir per invertir una matriu de Mueller-Jones si es té en compte que els factors $r_{00}, r_{01}, r_{10}, r_{11}$, $e^{i\left(\theta_{01}-\theta_{00}\right)}, e^{i\left(\theta_{11}-\theta_{00}\right)}$ and $e^{i\left(\theta_{11}-\theta_{00}\right)}$ són facilment calculables a partir d'una matriu de Mueller-Jones, tal com es mostra a [7]. Tot i això, aquestes equacions, per elles soles, no són útils des d'un punt de vista experimental, ja que, en general, el resultat d'un experiment de polarimetria és una matriu de Mueller, però no estrictament una matriu de Mueller-Jones.

Per invertir una matriu de Mueller experimental primer haurem de calcular la matriu de Mueller-Jones més propera a la matriu de Mueller experimental. Això es pot fer emprant l'anomenada matriu de coherència que fou introduïda per Cloude [100]. Els valors propis de la matriu de coherència permeten calcular la matriu de Mueller-Jones estimada corresponent a una matriu de Mueller experimental, tot seguint el procediment descrit a [99]. La validesa d'aquest procés estimatiu dependrà de si la matriu de Mueller experimental original era, ja d'entrada, propera a una matriu de Mueller-Jones. En general els resultats només seran fiables en el cas que el medi estudiat no introdueixi despolarització.

## A.3.2. Descomposició de les matrius de Mueller

En aquest apartat descriurem un metodologia d'anàlisi de les matrius de Mueller completament diferent. El nostre objectiu aquí és descomposar la matriu de Mueller experimental en components més simples i que tinguin una interpretació física més senzilla. Aquests procediments de descomposició no es basen en un model teòric que descrigui la interacció d'una ona electromagnètica amb una mostra, sinó que estan basats en les diferents propietats algebraiques que presenten les matrius de Mueller corresponents a diferents tipus de medi.

Qualsevol matriu de Mueller $\mathbf{M}$ es pot descomposar en el producte d'un diatenuador $\mathbf{M}_{D}$, un retardador $\mathbf{M}_{R}$ i un despolaritzador $\mathbf{M}_{\Delta}$

$$
\begin{equation*}
\mathbf{M}=\mathbf{M}_{\Delta} \mathbf{M}_{R} \mathbf{M}_{D} \tag{A.14}
\end{equation*}
$$

Aquesta factorització es coneix amb el nom de de descomposició de Lu-Chipman [82] i es pot aplicar a totes les matrius de Mueller experimentals. Si s'assumeix
que no hi ha despolarització $\left(\mathbf{M}_{\Delta}=\mathbf{I}\right)$ la descomposició passa a tenir només dos termes i es coneix generalment com descomposició polar [102,103].

Els tres factors de la descomposició de Lu-Chipman de l'Eq. (A.14) en general no commuten entre ells, és a dir, el seu producte depèn de l'ordre en què els termes es multipliquen. Això fa que la descomposició no sigui única, i que es puguin definir diferents variacions de la descomposició en funció de l'ordre en què els factos són multiplicats [105]. Aquest fet fa que els factors resultant de la descomposició puguin perdre la seva interpretatibilitat física, en el sentit que les dicroisme de la matriu $\mathbf{M}_{D}$ i la birefringència de la matriu $\mathbf{M}_{R}$ poden no coincidir, respectivament, amb el dicroisme i la birefringència de la matriu de Mueller M original.

## Descomposició pseudopolar

La descomposició pseudopolar està molt relacionada amb la descomposició de Lu-Chipman, però ofereix un tractament per a la no commutabilitat dels efectes òptics involucrats en el càlcul. Aquesta descomposició es pot deduir a partir la forma exponencial d'una matriu de Jones o de Mueller-Jones [vegeu Eq. (A.2)] si es fa servir la formula de Zassenhaus [113] que indica que l'exponencial de la suma de dos operadors que no commuten es pot obtenir com un producte infinit dels exponencials dels operadors i dels seus commutadors. Per fer més simple l'ús de la descomposició, ens quedarem només ens els dos primers termes de la succesió. En termes de la matriu de Mueller la descomposició pseudopolar truncada la podem escriure com

$$
\begin{equation*}
\mathbf{M} \cong \mathbf{M}_{\Delta} \mathbf{M}_{R} \mathbf{M}_{D} \mathbf{M}_{1 C} \mathbf{M}_{2 C} \tag{A.15}
\end{equation*}
$$

on $\mathbf{M}_{R}$ és un retardador que té la mateixa birefringència que $\mathbf{M}$ i $\mathbf{M}_{D}$ és una diatenuador amb el mateix dicroisme que $\mathbf{M} . \mathbf{M}_{1 C}$ i $\mathbf{M}_{2 C}$ son els dos primers termes d'una successió infinita de matrius, on cada terme d'ordre superior s'apropa més a la matriu unitat. Anomenem a les matrius $\mathbf{M}_{1 C}$ i $\mathbf{M}_{2 C}$ com matrius de correcció.

El càlcul de tots els factors de la descomposició pseudopolar a partir de d'una matriu de Mueller experimental es realitza a través d'un procés iteratiu que es detalla a [170]. És important remarcar un altre cop que malgrat que $\mathbf{M}_{R}$ i $\mathbf{M}_{D}$ que apareixen tant l'Eq. (A.15) com també a l'Eq. (A.14) i que en ambdós casos $\mathbf{M}_{D}$ es refereix a un diatenuador i $\mathbf{M}_{R}$ a un retardador, només en el cas de la descomposició pseudopolar les propietats d'aquests elements corresponen a les de la matriu original.

En un sistema que només presenti efectes òptics que commuten la descomposició pseudopolar es transforma automàticament en la descomposió polar (si no considerem despolarització) o de Lu-Chipman (si la despolarització és considerada), ja que, en aquestes condicions, els infinits termes de la successió con-
vergeixen cap a la matriu identitat $\mathbf{M}_{1 C}=\mathbf{M}_{2 C}=\ldots=\mathbf{I}$. Posat que coneixem l'estructura del les matrius de correcció [170] és possible derivar les condicions que asseguren la commutabilitat dels efectes òptics. Són aquestes:

$$
\begin{align*}
& \mathrm{CBLD}^{\prime}-\mathrm{LB}^{\prime} \mathrm{CD}=0, \\
& \mathrm{CDLB}-\mathrm{LDCB}=0,  \tag{A.16}\\
& \mathrm{LBLD}^{\prime}-\mathrm{LB}^{\prime} \mathrm{LD}=0
\end{align*}
$$

Aquestes condicions mostren que en un medi anisotrop i quiral ( $\mathrm{CD}, \mathrm{CB} \neq 0$ ) sempre hi haurà algun grau de no commutativitat. Un medi medi aquiral ( $\mathrm{CD}=$ $\mathrm{CB}=0$ ) tindrà propietats òptiques commutatives si els eixos de birefringencìa i dichroisme lineal coincideixen, ja que aleshores també es satisfà $\mathrm{LBLD}^{\prime}-\mathrm{LB}^{\prime} \mathrm{LD}=$ 0 .

## A.4. Dispositius experimentals

Els aparells que permeten la mesura d'elements de la matriu de Mueller d'una mostra es coneixen com polarímetres o el • lipsòmetres. Tots ells tenen es basen en una configuració elemental comuna: primer la llum travessa un generador d'estats de polarització (PSG), després interacciona amb la mostra i seguidament travessa un analitzador d'estats de polarització (PSA). La diferències més importants entre diferent tipus de el • lipsòmetres o polarímetres venen donades pels elements òptics que composen el PSG i el PSA. Les característiques del PSG i el PSA també determinen quins elements de la matriu de Mueller (eventualment tots) són mesurats en cada tipus d'aparell.

En aquesta tesi hem emprat un el •lipsòmetre generalitzat de doble modulador (2-MGE) [117,118] per obtenir mesures completes de la matriu de Mueller normalitzada. Aquest instrument, de construcció pròpia, utilitza dos parells de polaritzador-modulador fotoelàstic, un en el PSG i l'altre en el PSA. Els moduladors fotoelàstics (PEMs) son dispositius ressonants d'alta freqüència (en el nostre instrument $\sim 50 \mathrm{kHz}$ el del PSG i $\sim 60 \mathrm{kHz}$ el del PSA) que introdueixen un desfasament periòdic entre les components del camp elèctric. Com que en aquest treball estem interessat en realitzar mesures d'activitat òptica que requereixen una alta sensibilitat és molt convenient fer PEMs com a moduladors de la polarització de la llum enlloc d'altres tipus de moduladors, ja que els PEMs posseeixen un alt grau de puresa, eficiència i estabilitat en la modulació.

Les taules 5.1 i 5.2 enumeren tots els components òptics i electrònics que composen el 2-MGE, mentre que les Figs. 5.1 i 5.2 són, respectivament, representacions esquemàtiques del muntatge òptic i de les connexions electròniques de l'aparell. La llum que arriba al detector del 2-MGE és una complicada funció
temporal que es pot expressar com

$$
\begin{align*}
I(t)= & I_{d c}+I_{X 0} X_{0}+I_{Y 0} Y_{0}+I_{X 1} X_{1}+I_{X 0 X 1} X_{0} X_{1} \\
& +I_{X 0 Y 1} X_{0} Y_{1}+I_{Y 0 X 1} Y_{0} X_{1}+I_{Y 0 Y 1} Y_{0} Y_{1} . \tag{A.17}
\end{align*}
$$

Els termes $I_{d c}, I_{X 0}, I_{Y 0}, \ldots$ són constats que multipliquen les funcions base que depenen del temps:

$$
\begin{align*}
& X_{0}=\sin \left[A_{0} \sin \left(\omega_{0} t+\phi_{0}\right)\right]  \tag{A.18a}\\
& Y_{0}=\cos \left[A_{0} \sin \left(\omega_{0} t+\phi_{0}\right)\right]  \tag{A.18b}\\
& X_{1}=\sin \left[A_{1} \sin \left(\omega_{1} t+\phi_{1}\right)\right]  \tag{A.18c}\\
& Y_{1}=\cos \left[A_{1} \sin \left(\omega_{1} t+\phi_{1}\right)\right] \tag{A.18d}
\end{align*}
$$

on $A_{0}$ i $A_{1}$ són les amplituds de modulació, i les freqüències dels moduladors són $2 \pi \omega_{0}$ i $2 \pi \omega_{1}$.

La informació sobre la mostra està continguda els vuit paràmetres $I_{X 0}, I_{Y 0}$, $I_{X 1}$, etc que normalment s'expressen ja normalitzats a la intensitat continua de la llum $I_{d c}$. En funció de l'orientació azimutal del PSG i el PSA aquests vuit paràmetres es poden relacionar directament a vuit elements de la matriu de Mueller de la mostra. Per exemple, en el nostre instrument si el PSG i el PSA s'orienten a $\left(0^{\circ}, 0^{\circ}\right)$ dels eixos primaris de referència de l'aparell, els elements de la matriu de Mueller mesurables són:

$$
\mathbf{M}=\left(\begin{array}{cccc}
1 & \bullet & -I_{Y 0} & I_{X 0}  \tag{A.19}\\
\bullet & \bullet & \bullet & \bullet \\
-I_{Y 1} & \bullet & I_{Y 0 Y 1} & -I_{X 0 Y 1} \\
-I_{X 1} & \bullet & I_{Y 0 X 1} & -I_{X 0 X 1}
\end{array}\right)
$$

on els símbols (•)indiquen elements no mesurables en la configuració donada. Tanmateix, si es fan servir 4 configuracions de mesura diferents [per exemple orientacions $\left(0^{\circ}, 0^{\circ}\right),\left(0^{\circ}, 45^{\circ}\right),\left(45^{\circ}, 0^{\circ}\right)$ i $\left(45^{\circ}, 45^{\circ}\right)$ ] tots els elements de la matriu de Mueller normalitzada són mesurables.

## A.4.1. Rotació òptica aplicada a la mesura

Un dels usos principal del 2-MGE durant aquesta tesi ha estat la mesura espectroscòpica de la matriu de Mueller, però a mesura que augmentava la complexitat de les mostres que caracteritzaven ens vam adonar que el 2-MGE tenia problemes a l'hora mesurar mostres heterogènies al llar de la seva superfícies, és a dir, on les propietats òptiques variaven de punt a punt. De fet el problema sorgeix quan canvia l'angle azimutal PSG i/o el PSA: com que l'eix de rotació no coincideix exactament amb la direcció del feix de llum, es produeix una desplaçament del feix de llum sobre la mostra. A la pràctica això significa que en diferents configuracions de mesura s'estan analitzant punts lleugerament diferents de la mostra.

El desplaçament del feix depèn, evidentment de la qualitat de l'alineament, però també del gruix del PSG i el PSA. En el nostre cas en tenir el PSG i el PSA gruixos de desenes de mm es fa pal • lès que si s'aplica la rotació del PSG i el PSA el instrument disposa d'una resolució lateral pobre, cosa suposa un problema en mostres heterogènies.

Per solucionar aquest problema hem ideat un sistema on no es necessari moure el PSG i el PSA. En comptes d'això introduïm làmines de quars cristal • lí amb l'eix òptic perpendicular a la superfície com elements generadors d'una rotació òptica. La intenció és substituir la rotació mecànica del PSG i el PSA per la rotació òptica generada per un element amb CB. Utilitzem dos conjunts làmines de quars de diferents gruixos muntats cadascun d'ells en una roda de filtres, una situada entre el PSG i la mostra (quars 0), i l'altra entre la mostra i el PSA (quars 1). Les rodes de filtres permeten posar o treure les làmines de quars en el camí òptic d'una manera automàtica i ràpida.

Tal com mostra l'Eq. A.12c, el gruix de les làmines de quars és proporcional a l'angle de rotació òptica ( CB es el doble de la rotació òptica). Però a més a més CB és un efecte dispersiu, és a dir, per un determinat gruix del cristall de quars la rotació òptica varia amb la longitud d'ona [159]. Des del punt de vista del 2-MGE la rotació òptica "ideal" per a qualsevol longitud d'ona seria $45^{\circ}$, que per a una làmina d'1 mm de gruix es produeix aproximadament a 430 nm . Tot i això també és possible realitzar mesures amb angles de rotació òptica diferents de $45^{\circ}$ si l'efecte de les làmines de quars s'ha calibrat amb antelació. Per angles de rotació òptica arbitraris, en comptes de mesurar elements de la matriu de Mueller aillats, mesurarem combinacions de elements. Amb tot, com que el $2-\mathrm{MGE}$, ja és capaç de mesurar 8 elements de la matriu de Mueller sense necessitat de fer servir làmines de quarts, podrem determinar tots elements de la matriu de Mueller si es fan servir les següents configuracions:

- Configuració I. PSG / Mostra / PSA,
- Configuració II. PSG / Quars 0 / Mostra / PSA,
- Configuració III. PSG / Mostra / Quars 1 / PSA,
- Configuració IV. PSG / Quars 0 / Mostra / Quars 1 / PSA.

Els elements de la matriu de Mueller que mesurem a cada configuració venen donats per:

$$
\mathbf{M}=\left(\begin{array}{cccc}
1 & m_{01, I I} & m_{02, I} & m_{03, I}  \tag{A.20}\\
m_{10, I I I} & m_{11, I V} & m_{12, I I I} & m_{13, I I I} \\
m_{20, I} & m_{21, I I} & m_{22, I} & m_{23, I} \\
m_{30, I} & m_{31, I I} & m_{32, I} & m_{33, I}
\end{array}\right) .
$$

Noteu que amb la configuració I mesurem 8 elements de la matriu de Mueller, amb cadascuna de les configuracions II i III mesurem 3 nous elements i només
fem servir la configuració IV per mesurar l'element restant. La característica més important d'aquest mecanisme de mesura és que en tot moment el PSG i el PSA romanen en la mateixa posició i les làmines de quars no produeixen cap desplaçament del feix apreciable. Les rodes de filtres permeten accedir d'una manera ràpida i automàtica a qualsevol de les configuracions esmentades.

A part de l'efecte de rotació òptica les làmines de quars poden produir també algunes pertorbacions a la mesura si el seu eix òptic no es perfectament paral $\cdot$ lel al feix lluminós. En aquest cas, es farà manifesta alguna traça de birefringència lineal. Per poder calibrar i corregir aquest petit efecte, es poden fer servir les mateixes estratègies fetes servir a el • lipsometria generalitzada per calibrar les finestres d'un el • lipsòmetre [146].

## A.5. Mesures experimentals

## A.5.1. Quars

Quars és un cristall enantiomorfic molt comú a la Terra. Històricament la seva activitat òptica ha estat molt estudiada, ja que les primeres mesures d'activitat òptica de començaments del començaments del segle XIX es van fer en quars [14]. Tot i això la majoria de mesures que s'han fet sobre l'activitat òptica del quars tenen en comú que s'han realitzat per llum propagant-se a través de la única direcció del cristall que no mostra birefringència lineal (l'eix òptic) [153]. Tanmateix és ben conegut en cristal • lografia que els quars també mostra activitat òptica (en aquest cas només CB i no CD, ja que és un cristall transparent) en altres direccions del cristall. En aquestes circumstancies, però, la mesura esdevé molt més complexa.

La dependència del CB segons la direcció de propagació de la llum a través d'un cristall de quars es pot expressar rigorosament en termes del tensor de girotropria. En el cas del quars aquest tensor té dos elements diferents: $g_{33}$ per direccions paral • lel a l'eix òptic (senzill de mesurar) i $g_{11}$ per direccions perpendicular a l'eix òptic (difícil de mesurar):

$$
\mathbf{g}=\left[\begin{array}{ccc}
g_{11} & 0 & 0  \tag{A.21}\\
0 & g_{11} & 0 \\
0 & 0 & g_{33}
\end{array}\right]
$$

En aquest treball ens hem centrat en la mesura de l'element $g_{11}$ del quars. El nostre procediment de mesura està basat en la mesura de la matriu de Mueller en transmissió per incidència obliqua de la llum sobre la superfície del cristall. La Fig. 7.2 mostra la geometria bàsica del nostre experiment. Amb aquesta geometria hem demostrat que l'element $g_{11}$ es pot determinar experimentalment si els següents paràmetres són coneguts amb antelació: l'element $g_{33}$, l'angle
d'incidència i els índex de refracció ordinari i extraordinari del quars. Els detalls sobre com calcular aquest element del tensor es poden consultar a [159] o al capítol 7 d'aquesta tesi.

Les Figs. 7.8 i 7.9 mostren, respectivament, les mesures experimentals espectroscòpiques dels components del tensors girotropia d'un cristall de quars "dretà" $(\mathrm{RH})$ i un cristall "esquerrà" (LH). Tal com s'espera ambdós enantiomorfs tenen signes oposat en els elements del tensor de girotropia.

## A.5.2. Induccions quirals per efectes hidrodinàmics

L'estudi de les induccions quirals com a conseqüència d'efectes hidrodinàmics engloba tot un seguit d'experiments realitzats en el transcurs d'aquesta tesis que tenen en comú l'estudi l'activitat òptica de solucions que s'agiten en sentit horari (CW) o anti-horari (CCW) durant la mesura de la matriu de Mueller. El interès en aquest experiments rau en el fet que per certes solucions que contenen estructures supramoleculars de formes allargades i mides nanomètriques s'indueix una activitat òptica supramolecular que ve determinada pel sentit de l'agitació. Es tracta d'un fenomen totalment reversible i que dura només temps en el qual l'agitació està activa, cosa que la fa diferent induccions lligades a agitacions descrites prèviament $[176,188]$. Aquest efecte ha estat focus de certes controvèrsies i en diverses publicacions recents [177-179] s'ha especulat amb diverses interpretacions de les observacions experimentals. La nostra aproximació al problema ha estat obtenir tanta informació com fos possible sobre el fenòmen a partir de les nostres mesures experimentals amb el 2-MGE, cosa que finalment ha donat lloc a diferents publicacions [97, 180, 205].

Els experiments es van desenvolupar en cubetes de 10 mm de camí òptic i agitades emprant petites barretes magnètiques. En la major part d'experiments que hem realitzat les cubetes contenien J-agregats de porfirines. En certes porfirines no es mostrava cap efecte en agitar (per exemple en la porfirina $\mathrm{H}_{4} \mathrm{TPPS}_{3}$ ) mentre que en d'altres, com la $\mathrm{H}_{4} \mathrm{TPPS}_{4}$, apareixia una forta inducció d'activitat òptica. Els J-agregats d'aquestes porfirines son estructures allargades de matèria tova que arriben a fer uns quants centenar de nm de longitud i pocs nm de gruix. S'ha vist com els gradients de fluxos poden esculpir (tòrcer o doblegar) la forma d'aquestes estructures [181], i, possiblement, modificar els patrons d'ordre de les interaccions electròniques que sostenen l'agregat.

Mitjançant mesures espectroscòpiques de la matriu de Mueller en aquestes solucions [97] vam comprovar com la inducció d'activitat òptica es podia resseguir tant en els valors de CD com els CB calculat a partir de la matriu de Mueller (vegeu Fig. 8.4). Això va permetre confirmar que un procés d'inducció i canvi d'activitat era el responsable de l'efecte observat. També hem realitzat mesures de la matriu de Mueller de les cubetes agitades amb alta resolució espaial, tot fent servir el mètode descrit a l'apartat A.4.1. Les mesures de CD fetes amb
resolució espaial, tal com la mostrada a la Fig. 8.13 demostren que la inducció té lloc a la part central de la cubeta, en coincidència amb el vòrtex descendent generat per l'agitador. Addicionalment l'escanejat de la matriu de Mueller al llarg de tota la cubeta, també permet veure l'evolució de les anisotropies lineals a les diferents parts, i, en certa manera, això por conèixer in situ el patró de fluxos dins de la cubeta (vegeu un exemples a les Figs. 8.14 i 8.15, on es fa servir les projeccions de dicroisme lineal per una representació vectorial dels eixos de dicroisme).

Experiments anàlegs als descrits en aquest apartat s'han repetit en flascons o cubetes cilíndriques en substitució de les de secció quadrada i hem observat canvis en el signe dels senyals induïts. Atribuïm aquestes diferències al fet que per una cubeta cilíndrica, degut a la seva major simetria, també hi ha un flux quiral ascendent que actua en sentit contrari que el flux quiral descendent. Aquest fet es descriu detalladament a la secció 8.1.1 d'aquesta tesi.

## A.5.3. Mostres sòlides heterogènies

Una de les aplicacions més útils de les mesures de la matriu de Mueller amb resolució espaial és l'estudi de mostres sòlides heterogènies que tenen propietats òptiques canviants al llarg de la seva superfície.

## Dominis quirals en cristal • litzacions primes de compostos orgànics

La possibilitat d'identificar dominis quirals en cristal • litzacions preparades al laboratori a partir fusions de compostos orgànics dona peu a estudiar els processos de trencament i evolució de la simetria quiral que tenen lloc durant un canvi de fase com és la solidificació.

A partir de fusions de diversos compostos orgànics, com ara benzil, benzofenona, o binaftil vam preparar pel • lícules primes policristal • lines de compostos que formen cristalls quirals i enantiomorfics. Això significa poden cristal • litzar en dues formes diferents, una imatge especular de l'altra, i cadascuna de elles amb signes oposats d'activitat òptica. Com que el cost energètic per formar els dos enantiomorfs és idèntic s'hauria d'esperar els dos enantiòmers es formarien amb la mateixa probabilitat. Tanmateix se sap que poden ocorren processos, com ara certes reaccions autocatalítiques, que trenquen aquesta simetria quiral.

Els millors resultats d'aquestes mesures es van obtenir per cristal • litzacions de benzil. Aquest cristall té una banda prou intensa de CD al voltant de 400 nm (vegeu Fig. 9.1) que facilita la identificació dels dominis quirals si les mesures es fan a aquesta longitud d'ona. La Fig. 9.3 mostra un exemple de les matrius de Mueller obtingudes i a la Fig. 9.4 es poden apreciar clarament els dominis quirals identificats en aquesta mostra. Els dominis quirals identificats en la majoria de mostres (Fig. 9.6) estan separats per línies rectes. Segons certs models teòrics
sobre processos de trencament de simetria $[192,193]$ aquest tipus de fronteres és la única manera on dominis quirals de diferent signe poden sobreviure en sistemes que evolucionen cap a la homoquiralitat.

## Meteorits

La possibilitat de realitzar mesures polarimètriques acurades i amb una gran resolució espaial en mostres de condrites carbonàcies (un tipus de meteorit amb abundant presencia de compostos de carboni i on s'han detectat aminoàcids) representarien un pas important en la recerca dels orígens de l' homoquiralitat terrestre. Com a primera incursió en aquest àmbit d'estudi vam realitzar algunes mesures preliminars [194] en talls prims i polits dels meteorits Murchison, Murray i Cold Bekkeveld. La nostra hipòtesi a l'hora de planejar l'estudi era que els processos d'estrès (gradients de cisalla principalment) en que aquests cossos van estar sotmesos durant la seva solidificació podrien haver donar lloc a un excés enantiomeric fractures o distorsions en la matriu inorgànica del meteorit que posteriorment es podria haver transmès a les reaccions orgàniques.

L'objectiu de les nostres mesures era dons similar a la mencionada a l'apartat anterior: la identificació de dominis quirals en mostres sòlides. En aquest cas però la dificultat era molt més gran ja que les mostres estudiades eren altament heterogènies i caldria una resolució submicromètrica per poder-les caracteritzar adequadament. Quan vam analitzar estadísticament els resultats obtinguts per la mostra de Murchison (Fig. 9.8) van observar una desviació en CB no esperada que no apareixia per les altres mostres. Això semblaria refermar la nostra hipòtesi, ja que precisament al meteorit Murchison és on s'han descrit amb més claredat la desviació quiral del aminoàcids continguts en ell [37, 197, 198].

Les mesures en condrites carbonàcies representen tot un desafiament experimental i requeririen estudis molt més amplis i amb més recursos. Per ara les nostres mesures han posat de relleu una forma d'estudiar aquestes mostres que fins ara no s'havia ni plantejat i els resultats obtinguts haurien de servir com a estímul per futurs treballs en aquest àmbit.

## A.5.4. Conclusions

La resolució de les equacions de Maxwell que descriuen la interacció d'un medi anisòtrop i giròtrop amb llum no es pot tractar analíticament en la majoria de casos i es difícil modelitzar les matrius de Jones o de Mueller a partir del tensor del tensor dielèctric i de girotropia del medi. Tanmateix, es pot fer servir la representació infinitesimal del medi introduïda per Jones i basada en efectes òptics bàsics per descriure el comportament de qualsevol component òptic. Aquest mètode genera equacions analítiques molt útils per interpretar resultats experimentals. A més, en contra del suggerit per alguns autors, del treball original de

Jones no es pot inferir l'existència de cap nou tipus de birefringència i dicroisme. LB i LB' (al igual que LD i LD') són només projeccions del mateix fenomen físic.

Hem determinat equacions analítiques que permeten invertir qualsevol matriu de Jones o de Mueller-Jones. Aquesta inversió és pot aplicar a qualsevol matriu de Mueller experimental si no inclou una important despolarització i permet la determinació els efectes físics d'interès, com ara CD i CB. Una altra alternativa per a la interpretació de les matrius de Mueller experimentals és l'ús d'una descomposició tipus producte com la descomposició polar o de Lu-Chipman que permet factoritzar una matriu de Mueller termes més simples. El problema que presenten les descomposicions és que, en general, els factors de la descomposició depenen de l'ordre en què són multiplicats. En aquest treball hem derivat les condicions que ha de satisfer un medi, per assegurar la conmutatibilitat dels factors de la descomposició. Quan aquestes condicions es compleixen la descomposició pot ésser aplicada. El treball en aquesta àrea ens ha portat a descriure una nova descomposició i batejar-la com descomposició pseudopolar.

Dins d'un àmbit ja plenament instrumental hem construït un el • lipsòmetre espectroscòpic conegut com 2-MGE. La característica més distintiva del 2-MGE és l'ús de dos PEMs funcionant independentment, cosa que li permet ésser capaç de mesurar la matriu de Mueller d'una mostra, ja sigui en transmissió o en reflexió. Hem demostrat que aquest tipus d'aparell pot ser utilitzat per mesures d'activitat òptica que requereixen una gran sensibilitat, i que pot ser usat tant en líquids com en sòlids, en un ampli rang de longituds d'ona, des de el UV fins al IR proper.

Hem desenvolupat un nou mètode de mesura que pot expandir i/o millorar les capacitats del 2-MGE o d'altres tipus d'el • lipsòmetres o polarímetres. Aquest nou mètode experimental es basa en la utilitació d'elements amb rotació òptica com substituts de les unitats de rotació mecànica que es fan servir per orientar el PSG i el PSA i així evitar problemes relacionats amb la desviació del feix. Aquest mètode ens permet realitzar mesures de la matriu de Mueller amb alta resolució espacial que no serien possibles emprant les unitats de rotació mecànica.

Mitjançant la mesura de la matriu de Mueller hem realitzat nombrosos experiments dirigits a determinar l'activitat òptica de diversos sistemes. Podem destacar que per primer cop s'ha mesurat completament el tensor de girotropia d'un cristall com el quars a partir de mesures espectroscòpiques de la matriu de Mueller. Un altre grup de mesures importants han estat les realitzades en solucions agitades que contenien nanopartícules allargades de matèria tova. En aquest cas es van realitzar tant mesures espectroscòpiques com de tipus "mapping" que han permès estudiar amb detall i/o aclarir aspectes desconeguts d'un fenomen fins ara controvertit d'inducció de quiralitat supramolecular per efectes hidrodinàmics.

Un dels àmbits de mesura que creiem que pot tenir mes perspectives de futur es la caracterització mitjaçant la matriu de Mueller de sòlides i heterogènies, és
amb propietats òptiques que varien al llarg de la superfície. Exemple d'això han estat la determinació de dominis quirals en pel • lícules policristal • lines d'alguns compostos orgànics, o les mesures preliminars realitzades en seccions de meteorits.

## Appendix B.

## Computer software

The construction of a 2-MGE also involves the development a certain amount of computer software. All the theoretical descriptions of the instrument given in Part III of this thesis would not be of utility without an interface able to communicate with the different components of the 2-MGE and to handle several different routines of data acquisition and calibration. The 2-MGE was developed with the idea of offering a friendly and complete GUI (Graphic User Interface) that was easy to handle for any eventual user of the equipment. The inspiration we used for the organization of this GUI was the GUI programmed by Gerald E. Jellison for the control of the 2-MGEs built by him and available at Oak Ridge National Laboratory facilities.

2-MGE Software has been developed in Labview, which is a visual programming language from National Instruments. One benefit of LabVIEW is the extensive support for accessing instrumentation hardware. Another interesting feature of Labview is the easiness to create attractive GUIs without the necessity of being a very skilled programmer. Moreover, despite Labview is sometimes regarded as a limited language, only convenient for controlling instrumentation, we have been surprised by its versatility and it has suited all our demands.

Fig. B. 1 shows the front panel of the 2-MGE Software. Buttons on the left column give access to the basic functionalities of the 2-MGE when using mechanical rotators for the PSG and PSA: "calibration" for a complete calibration of the instruments, "oscilloscope" for a real time visualization of the signal, "transmission measurement" for a partial or complete determination of the transmissive Mueller matrix, "reflection mesurement" for a partial or complete determination of the reflective Mueller matrix using the NSC notation and "scattering measurement" for a partial or complete determination of the scattering Mueller matrix (this is still a prototype routine in light source has to be switched to a laser). Buttons on the right column of Fig. B. 2 give access to different functions that involve the quartz assisted method introduced in chapter 6 . Prior the selection any of these options the thickness of the set of quartz plates needs to be choosed: "thick" for measurements in the VIS and "thin" for measurements in the UV. After that users can choose between two types of calibrations and two different modes of spectrocopic Mueller matrix transmission measurements. The button "Mueller mapping" is used to obtain spatially resolved measurements of


Figure B.1.: Front panel of the 2-MGE Software. It gives access to every function of the program.).
the Mueller matrix. As this appendix is intended to give a general vision of the software developed we do not find it necessary to explain the particularities and characteristics of every acquisition type.

The 2-MGE Software is structured in a very decentralized manner. Each of the enumerated acquisition types is mostly independent from the others. They only share the configuration parameters of the 2-MGE which are accessible in every acquisition type and saved in a computer file. Fig. B. 2 shows the configuration windows available in the 2-MGE Software. Three of them are used to configure the mechanical elements of our 2-MGE: rotator stages of the PSG and PSA, the $\mathrm{X}-\mathrm{Y}$ translation units used to scan the sample and tilting mechanism used to set the angle of incidence. One more window is used to set the parameters of the digitizer and the remaining panel is used to define the wavelengths of measurement and the configuration of the monochromator.

Figs. B.3, B.4, B. 5 and B. 6 show various screenshots of the 2-MGE Software while performing different functions. At the present status the software is mostly useable in all their most significative functions, however we are aware that it contains many bugs and several functions could be still greatly improved. Despite the user interface looks quite accessible, the software still needs to be handled by a trained user and with some knowledge about the contents of this thesis.


Figure B.2.: Configuration panels of the 2-MGE. The "rotators", "monochromator" and "digitizer" windows are available for all acquisition routines. The "tilting" and "mapping" windows are exclusive of certain acquisition types.

Figure B.3.: Oscilloscope panel included in the 2-MGE Software. It permits a real time visualization of the digitized signal and the eight independent parameters $I_{X 0}, I_{Y 0}, I_{X 1}, I_{Y 1}, I_{X 0 X 1}, I_{X 0 Y 1}, I_{Y 0 X 1}$ and $I_{Y 0 Y 1}$ that the 2-MGE measures. The instrument was configured here in the straight-through configuration with PSG and PSA oriented at $\left(0^{\circ}, 0^{\circ}\right)$.

Figure B.4.: Panel shown during the spectroscopic calibration of the PEMs and the quartz rotators. Circles depict experimental data points while solid lines are calibration fitting. In this calibration panel we show the static strain of the modulators, the Bessel voltage needed to set the amplitude of modulation to 2.4048 rad , the electronic frequency-independent gain factor, the static strains of the quartz plates and their CB (in this case for plates of 1 mm thick)


Figure B.5.: Example of the panel window shown during a spectroscopic Mueller matrix measurement. The Mueller matrix is presented on screen in real time during the data acquisition. Mueller matrix elements measured in different configurations are presented in different colours.
Figure B.6.: Spatially resolved measurements are obtained using this panel. The intensity mappings corresponding to each one of the four measurements configurations accessed by quartz plates are given. Each Mueller matrix scanning takes 4 measurement rounds, and through these 4 intensity mappings representations the user can follow the status of its measurements.

## List of publications

Refereed submissions made during the course of this thesis

1. O. Arteaga, A. Canillas, J. Crusats, Z. El-Hachemi, J. Llorens, E. Sacristan and J. M. Ribó "Emergence of supramolecular chirality by flows", Submitted to Angew. Chem., (2010).
2. O. Arteaga, Z. El-Hachemi, A. Canillas and J. M. Ribó "Transmission Mueller Matrix Ellipsometry of Chirality Switching Phenomena", Submitted to Thin Solid Films, (2010).
3. O. Arteaga, Z. El-Hachemi, A. Canillas and J. M. Ribó "Transmission Mueller Matrix Ellipsometry of Chirality Switching Phenomena", Submitted to Thin Solid Films, (2010).
4. O. Arteaga, "On the existence of Jones birefringence and Jones dichroism", Opt. Lett. 35, 1359-1360 (2010).
5. O. Arteaga and A. Canillas, "Analytic inversion of the Mueller-Jones polarization matrices for homogeneous media", Opt. Lett. 35, 559-561 (2010).
6. O. Arteaga, A. Canillas, and J. Jellison, "Determination of the components of the gyration tensor of quartz by oblique incidence transmission twomodulator generalized ellipsometry", Appl. Opt. 48, 5307-5317 (2009).
7. O. Arteaga, A. Canillas, J. Crusats, Z. El-Hachemi, G. Jellison, J. Llorca, and J. M. Ribó, "Chiral biases in solids by effect of shear gradients: A speculation on the deterministic origin of biological homochirality", Origins of Life and Evolution of Biospheres 40, 27-40 (2010).
8. O. Arteaga, A. Canillas, R. Purrello, and J. M. Ribó, "Evidence of induced chirality in stirred solutions of supramolecular nanofibers", Opt. Lett. 34, 2177-2179 (2009).
9. O. Arteaga, C. Escudero, G. Oncins, Z. El-Hachemi, J. Llorens, J. Crusats, A. Canillas, and J. M. Ribó, "Reversible mechanical induction of optical activity in solutions of soft-matter nanophases", Chemistry - An Asian Journal 4, 1687-1696 (2009).
10. O. Arteaga and A. Canillas, "Pseudopolar decomposition of the Jones and Mueller-Jones exponential polarization matrices", J. Opt. Soc. Am. A 26, 783-793 (2009).
11. Z. El-Hachemi, C. Escudero, O. Arteaga, A. Canillas, J. Crusats, G. Mancini, R. Purrello, A. Sorrenti, A. D'Urso, and J. M. Ribo, "Chiral sign selection on the J-aggregates of diprotonated tetrakis-(4-sulfonatophenyl)porphyrin by traces of unidentified chiral contaminants present in the ultra-pure water used as solvent", Chirality 21, 408-412 (2009).
12. O. Arteaga, Z. El-Hachemi, and A. Canillas, "Application of transmission ellipsometry to the determination of cd spectra of porphyrin J-aggregates", physica status solidi (a) 205, 797-801 (2008).
13. S. Portal, M. Vallve, O. Arteaga, J. Ignes Mullol, A. Canillas, and E. Bertran, "Optical characterization of colloidal crystals based on dissymmetric metalcoated oxide submicrospheres", Thin Solid Films 517, 1053-1057 (2008).
14. Z. El-Hachemi, O. Arteaga, A. Canillas, J. Crusats, C. Escudero, R. Kuroda, T. Harada, M. Rosa, and J. M. Ribó , "On the mechano-chiral effect of vortical flows on the dichroic spectra of 5 -phenyl-10,15,20-tris(4-sulfonatophenyl) porphyrin J-aggregates", Chemistry - A European Journal 14, 6438-6443 (2008).

## Patent

1. O. Arteaga, A. Canillas, and J. M. Ribó, "Method and system for determining a plurality of elements of a Mueller matrix", Patent application submitted, May 2010.

## Bibliography

[1] R. C. Jones, "A new calculus for the treatment of optical systems," J. Opt. Soc. Am. 31, 488-493 (1941).
[2] H. Mueller, "The foundations of optics," Journal of the Optical Society of America 38, 661-662 (1948).
[3] R. M. A. Azzam and N. M. Bashara, Ellipsometry and Polarized Light (North Holland, 1988).
[4] R. Barakat, "Bilinear constraints between elements of the $4 \times 4$ Mueller-Jones transfer matrix of polarization theory," Optics Communications 38, 159-161 (1981).
[5] J. W. Hovenier, "Structure of a general pure Mueller matrix," Appl. Opt. 33, 8318-8324 (1994).
[6] J. J. Gil, "Polarimetric characterization of light and media," The European Physical Journal Applied Physics 40, 1-47 (2007).
[7] O. S. O. America, Handbook of Optics, Vol. 2: Devices, Measurements, and Properties, Second Edition (McGraw-Hill Professional, 1994), 2nd ed.
[8] C. Brosseau, Fundamentals of Polarized Light: A Statistical Optics Approach (Wiley-Interscience, 1998).
[9] D. Goldstein, Polarized Light, Revised and Expanded (Optical Science and Engineering) (CRC, 2003), 2nd ed.
[10] J. J. Gil and E. Bernabeu, "A depolarization criterion in Mueller matrices," Journal of Modern Optics pp. 259-261 (1985).
[11] J. J. Gil and E. Bernabeu, "Depolarization and polarization indices of an optical system," Journal of Modern Optics 33, 185-189 (1986).
[12] S. R. Cloude and E. Pottier, "Concept of polarization entropy in optical scattering," Optical Engineering 34, 1599-1610 (1995).
[13] R. Ossikovski, "Alternative depolarization criteria for Mueller matrices," J. Opt. Soc. Am. A 27, 808-814 (2010).
[14] D. F. Arago, "Sur une modification remarquable qu' éprouvent les rayons lumineux dans leur passage à travers certains corps diaphanes, et sur quelques autres nouveaux phénomènes d'optique," Mem. Inst. 1, 93-134 (1811).
[15] J. B. Biot, "Mémoire sur un nouveau genre d'oscillations que les molécules de la lumière éprouvent, en traversant certains cristaux," Mem. Inst. 1, 1-372 (1812).
[16] Fresnel, "Ueber das Licht," Annalen der Physik und Chemie 81, 223-256 (1825).
[17] W. Haidinger, "Beobachtung der licht-polarisations-büschel auf flächen, welche das licht in zwei senkrecht auf einander stehenden richtungen polarisiren," Annalen der Physik und Chemie 144, 305-319 (1846).
[18] A. Cotton, "Absorption inégale des rayons circulaires droit et gauche dans certains corps actifs," C. R. Acad. Sci. 120, 989-991 (1895).
[19] T. M. Lowry, Optical rotatory power (Longman Scientific and Technical, 1935).
[20] S. Chandrasekhar, "Simple model for optical activity," American Journal of Physics 24, 503-506 (1956).
[21] P. L. Polavarapu, A. G. Petrovic, and P. Zhang, "Kramers-Kronig transformation of experimental electronic circular dichroism: Application to the analysis of optical rotatory dispersion in dimethyl-L-tartrate," Chirality 18, 723-732 (2006).
[22] M. Krykunov, M. D. Kundrat, and J. Autschbach, "Calculation of circular dichroism spectra from optical rotatory dispersion, and vice versa, as complementary tools for theoretical studies of optical activity using time-dependent density functional theory," The Journal of Chemical Physics 125, 194110+ (2006).
[23] J. Frenkel, "On the transformation of light into heat in solids. I," Physical Review Online Archive (Prola) 37, 17-44 (1931).
[24] V. Agranovich, Excitations in Organic Solids (International Series of Monographs on Physics) (Oxford University Press, USA, 2009).
[25] C. A. Schalley, ed., Analytical Methods in Supramolecular Chemistry (Wiley-VCH, 2007), 1st ed.
[26] S. E. Boiadjiev and D. A. Lightner, "Exciton chirality. (A) Origins of and (B) applications from strongly fluorescent dipyrrinone chromophores," Monatshefte für Chemie / Chemical Monthly 136, 489-508 (2005).
[27] E. E. Jelley, "Spectral absorption and fluorescence of dyes in the molecular state," Nature 138, 1009-1010 (1936).
[28] G. Scheibe, "Über die veränderlichkeit der absorptionsspektren in lösungen und die nebenvalenzen als ihre ursache," Angewandte Chemie 50, 212-219 (1937).
[29] V. Avetisov, "Question 4: Short remarks about the origin of homochirality," Origins of Life and Evolution of Biospheres 37, 367-370 (2007).
[30] A. Guijarro and M. Yus, The origin of chirality in the molecules of life: A revision from awareness to the current theories and perspectives of this unsolved problem (Royal Society of Chemistry, 2008).
[31] I. Weissbuch, L. Leiserowitz, and M. Lahav, ""stochastic mirror symmetry breaking" via self-assembly, reactivity and amplification of chirality: Relevance to abiotic conditions," in "Prebiotic Chemistry," , vol. 259 of Topics in Current Chemistry, P. Walde, ed. (Springer-Verlag, Berlin/Heidelberg, 2005), chap. 137067, pp. 123165.
[32] D. K. Kondepudi and K. Asakura, "Chiral autocatalysis, spontaneous symmetry breaking, and stochastic behavior," Accounts of Chemical Research 34, 946-954 (2001).
[33] P. L. Luisi, The emergence of life: from chemical origins to synthetic biology (Cambridge University Press, 2006).
[34] F. C. Frank, "On spontaneous asymmetric synthesis." Biochimica et biophysica acta 11, 459-463 (1953).
[35] K. Soai, T. Shibata, H. Morioka, and K. Choji, "Asymmetric autocatalysis and amplification of enantiomeric excess of a chiral molecule," Nature 378, 767-768 (1995).
[36] S. Pizzarello, "The chemistry of life's origin: A carbonaceous meteorite perspective," Accounts of Chemical Research 39, 231-237 (2006).
[37] J. R. Cronin and S. Pizzarello, "Enantiomeric excesses in meteoritic amino acids," Science 275, 951-955 (1997).
[38] R. L. Disch and D. I. Sverdlik, "Apparent circular dichroism of oriented systems," Analytical Chemistry 41, 82-86 (1969).
[39] Y. Shindo, M. Nakagawa, and Y. Ohmi, "On the problems of CD spectropolarimeters. II: Artifacts in CD spectrometers," Applied Spectroscopy pp. 860-868 (1985).
[40] N. Berova, K. Nakanishi, and R. Woody, Circular dichroism: principles and applications (Wiley-VCH, 2000).
[41] J. Schellman and H. P. Jensen, "Optical spectroscopy of oriented molecules," Chemical Reviews 87, 1359-1399 (1987).
[42] H. P. Jensen, J. A. Schellman, and T. Troxell, "Modulation techniques in polarization spectroscopy," Appl. Spectrosc. 32, 192-200 (1978).
[43] J. A. Schellman, "Circular dichroism and optical rotation," Chemical Reviews 75, 323-331 (1975).
[44] Y. Shindo and M. Nakagawa, "On the artifacts in circularly polarized emission spectroscopy," Applied Spectroscopy pp. 32-38 (1985).
[45] Y. Shindo, "On the problems of CD spectropolarimeter (IV) artifacts due to the light scattering by small particles," Applied Spectroscopy 39, 713-715 (1985).
[46] Y. Shindo and Y. Ohmi, "New polarization-modulation spectrometer for simultaneous circular dichroism and optical rotary dispersion measurements (I): Instrument design, analysis, and evaluation," Review of Scientific Instruments 56, 2237-2242 (1985).
[47] Y. Shindo and M. Nakagawa, "Circular dichroism measurements. I. calibration of a circular dichroism spectrometer," Review of Scientific Instruments 56, 32-39 (1985).
[48] R. Kuroda, T. Harada, and Y. Shindo, "A solid-state dedicated circular dichroism spectrophotometer: Development and application," Review of Scientific Instruments 72, 3802-3810 (2001).
[49] R. Kuroda and T. Honma, "CD spectra of solid-state samples," Chirality 12, 269277 (2000).
[50] R. Kuroda, Chiral photochemistry, Molecular and supramolecular photochemistry (Marcel Dekker, 2004).
[51] T. Harada, H. Hayakawa, and R. Kuroda, "Vertical-type chiroptical spectrophotometer (I): Instrumentation and application to diffuse reflectance circular dichroism measurement," Review of Scientific Instruments 79, 073103+ (2008).
[52] T. Harada, Y. Miyoshi, and R. Kuroda, "High performance diffuse reflectance circular dichroism spectrophotometer," Review of Scientific Instruments 80, 046101+ (2009).
[53] E. Castiglioni, P. Biscarini, and S. Abbate, "Experimental aspects of solid state circular dichroism," Chirality 21, E28-E36 (2009).
[54] J. Kobayashi and Y. Uesu, "A new optical method and apparatus "HAUP" for measuring simultaneously optical activity and birefringence of crystals. I. Principles and construction," Journal of Applied Crystallography 16, 204-211 (1983).
[55] J. Kobayashi, T. Asahi, S. Takahashi, and A. M. Glazer, "Evaluation of the systematic errors of polarimetric measurements: application to measurements of the gyration tensors of $\alpha$-quartz by the HAUP," Journal of Applied Crystallography 21, 479-484 (1988).
[56] J. R. L. Moxon and A. R. Renshaw, "The simultaneous measurement of optical activity and circular dichroism in birefringent linearly dichroic crystal sections. I. Introduction and description of the method," Journal of Physics: Condensed Matter 2, 6807-6836 (1990).
[57] P. Gomez and C. Hernandez, "High-accuracy universal polarimeter measurement of optical activity and birefringence of $\alpha$-quartz in the presence of multiple reflections," J. Opt. Soc. Am. B 15, 1147-1154 (1998).
[58] K. Claborn, J. Herreros Cedres, C. Isborn, A. Zozulya, E. Weckert, W. Kaminsky, and B. Kahr, "Optical rotation of achiral pentaerythritol," Journal of the American Chemical Society 128, 14746-14747 (2006).
[59] W. Kaminsky, M. A. Geday, J. Herreros-Cedres, and B. Kahr, "Optical rotatory and circular dichroic scattering," The Journal of Physical Chemistry A 107, 28002807 (2003).
[60] K. Claborn, E. Puklin-Faucher, M. Kurimoto, W. Kaminsky, and B. Kahr, "Circular dichroism imaging microscopy: Application to enantiomorphous twinning in biaxial crystals of 1,8-dihydroxyanthraquinone," Journal of the American Chemical Society 125, 14825-14831 (2003).
[61] J. H. Cedrés, C. H. Rodriguez, and W. Kaminsky, "Absolute optical rotation of $\mathrm{CsLiB}_{6} \mathrm{O}_{1} 0$," Journal of Applied Crystallography 38, 544-554 (2005).
[62] W. Kaminsky, E. Weckert, H. Kutzke, A. M. Glazer, and H. Klapper, "Non-linear optical properties and absolute structure of metastable 4-methyl benzophenone," Zeitschrift f?r Kristallographie 221, 294-299 (2006).
[63] K. Claborn, C. Isborn, W. Kaminsky, and B. Kahr, "Optical rotation of achiral compounds," Angewandte Chemie International Edition 47, 5706-5717 (2008).
[64] W. Kaminsky, "Experimental and phenomenological aspects of circular birefringence and related properties in transparent crystals," Reports on Progress in Physics 63, 1575-1640 (2000).
[65] M. F. Maestre and J. E. Katz, "A circular dichroism microspectrophotometer," Biopolymers 21, 1899-1908 (1982).
[66] W. Kaminsky, K. Claborn, and B. Kahr, "Polarimetric imaging of crystals," Chem. Soc. Rev. 33, 514-525 (2004).
[67] J. H. Freudenthal, E. Hollis, and B. Kahr, "Imaging chiroptical artifacts," Chirality 21, S20-S27 (2009).
[68] B. Kahr, J. Freudenthal, and E. Gunn, "Crystals in light $\dagger$," Accounts of Chemical Research 43, 684-692 (2010).
[69] D. W. Berreman, "Optics in stratified and anisotropic media: $4 \times 4$-matrix formulation," J. Opt. Soc. Am. 62, 502-510 (1972).
[70] M. Schubert, "Polarization-dependent optical parameters of arbitrarily anisotropic homogeneous layered systems," Phys. Rev. B 53, 4265-4274 (1996).
[71] E. Georgieva, "Reflection and refraction at the surface of an isotropic chiral medium: eigenvalue eigenvector solution using a $4 \times 4$ matrix method," Journal of the Optical Society of America A 12, 2203-2211 (1995).
[72] E. U. Condon, "Theory of optical rotatory power," Reviews of modern physics $\mathbf{9}$, 432-457 (1937).
[73] R. M. A. Azzam, "Photopolarimetric measurement of the Mueller matrix by Fourier analysis of a single detected signal," Opt. Lett. 2, 148-150 (1978).
[74] H. Wöhler, G. Haas, M. Fritsch, and D. A. Mlynski, "Faster $4 \times 4$ matrix method for uniaxial inhomogeneous media," J. Opt. Soc. Am. A 5, 1554-1557 (1988).
[75] C. R. Jones, "New calculus for the treatment of optical systems. VII. Properties of the N-matrices," J. Opt. Soc. Am. 38, 671-685 (1948).
[76] R. M. A. Azzam, "Propagation of partially polarized light through anisotropic media with or without depolarization: A differential $4 \times 4$ matrix calculus," J. Opt. Soc. Am. 68, 1756-1767 (1978).
[77] Complex Component Characterization Based on Elementary Matrices, OSA Technical Digest (CD) (Optical Society of America, 2009).
[78] R. Barakat, "Exponential versions of the Jones and Mueller-Jones polarization matrices," J. Opt. Soc. Am. A 13, 158-163 (1996).
[79] R. C. Jones, "New calculus for the treatment of optical systems. VIII. Electromagnetic theory," J. Opt. Soc. Am. 46, 126-131 (1956).
[80] O. Arteaga, "Mesura de dicroisme circular i activitat òptica molecular mitjançant el-lipsometria generalitzada." Master's thesis, University of Barcelona (2007).
[81] O. Arteaga, Z. El-Hachemi, and A. Canillas, "Application of transmission ellipsometry to the determination of CD spectra of porphyrin J-aggregates," physica status solidi (a) 205, 797-801 (2008).
[82] S.-Y. Lu and R. A. Chipman, "Interpretation of Mueller matrices based on polar decomposition," J. Opt. Soc. Am. A 13, 1106-1113 (1996).
[83] G. E. Jellison, J. D. Hunn, and C. M. Rouleau, "Normal-incidence generalized ellipsometry using the two-modulator generalized ellipsometry microscope," Appl. Opt. 45, 5479-5488 (2006).
[84] E. B. Graham and R. E. Raab, "On the Jones birefringence," Proceedings of the Royal Society of London. Series A, Mathematical and Physical Sciences 390, 73-90 (1983).
[85] C. Graham and R. E. Raab, "Eigenvector approach to the evaluation of the Jones N matrices of nonabsorbing crystalline media," J. Opt. Soc. Am. A 11, 2137-1944 (1994).
[86] R. E. Raab, "Some unmeasured crystal properties," Crystal Research and Technology 38, 202-214 (2003).
[87] T. Roth and G. L. J. A. Rikken, "Observation of magnetoelectric Jones birefringence," Physical Review Letters 85, 4478-4481 (2000).

## Bibliography

[88] D. Budker and J. E. Stalnaker, "Magnetoelectric Jones dichroism in atoms," Physical Review Letters 91, 263901+ (2003).
[89] V. V. Chernushkin, P. V. Mironova, and V. D. Ovsiannikov, "Magnetoelectric Jones spectroscopy of alkali atoms," Journal of Physics B: Atomic, Molecular and Optical Physics 41, 115401+ (2008).
[90] P. V. Mironova, V. D. Ovsiannikov, and V. V. Chernushkin, "Magnetoelectric Jones birefringence and dichroism in a medium of free atoms," Journal of Physics B: Atomic, Molecular and Optical Physics 39, 4999-5011 (2006).
[91] A. Rizzo and S. Coriani, "Jones birefringence in gases: Ab initio electron correlated results for atoms and linear molecules," The Journal of Chemical Physics 119, $11064+(2003)$.
[92] M. Izdebski, W. Kucharczyk, and R. E. Raab, "Effect of beam divergence from the optic axis in an electro-optic experiment to measure an induced Jones birefringence," J. Opt. Soc. Am. A 18, 1393-1398 (2001).
[93] V. A. De Lorenci and G. P. Goulart, "Magnetoelectric birefringence revisited," Physical Review D 78, 045015+ (2008).
[94] D. L. Andrews, G. J. Daniels, and G. E. Stedman, "Magnetoelectric Jones dichroism," Journal of Physics B: Atomic, Molecular and Optical Physics 36, L357-L364 (2003).
[95] L. D. Barron, Molecular light scattering and optical activity (Cambridge University Press, 2004).
[96] O. Arteaga, "On the existence of Jones birefringence and Jones dichroism," Opt. Lett. 35, 1359-1360 (2010).
[97] O. Arteaga, A. Canillas, R. Purrello, and J. M. Ribó, "Evidence of induced chirality in stirred solutions of supramolecular nanofibers," Opt. Lett. 34, 2177-2179 (2009).
[98] Chandrasekhar, Liquid Crystals (Cambridge University Press, 1993), 2nd ed.
[99] O. Arteaga and A. Canillas, "Analytic inversion of the Mueller-Jones polarization matrices for homogeneous media," Opt. Lett. 35, 559-561 (2010).
[100] S. R. Cloude, "Conditions for physical realizability of matrix operators in polarimetry," in "Society of Photo-Optical Instrumentation Engineers (SPIE) Conference Series," , vol. 1166, R. ${ }^{\sim}$ A. ${ }^{\sim}$ Chipman, ed. (1990), vol. 1166, pp. 177-185.
[101] A. A. Kochanovskij, Light scattering reviews 4: single light scattering and radiative transfer (Springer, 2008).
[102] S.-Y. Lu and R. A. Chipman, "Homogeneous and inhomogeneous Jones matrices," J. Opt. Soc. Am. A 11, 766+ (1994).
[103] C. Whitney, "Pauli-algebraic operators in polarization optics," J. Opt. Soc. Am. 61, 1207+ (1971).
[104] J. J. Gil and E. Bernabeu, "Obtainment of the polarizing and retardation parameters of nondepolarizing optical system from polar decomposition of its Mueller matrix," Optik 76, 67-71 (1987).
[105] R. Ossikovski, A. De Martino, and S. Guyot, "Forward and reverse product decompositions of depolarizing Mueller matrices," Opt. Lett. 32, 689-691 (2007).
[106] S. N. Savenkov, O. I. Sydoruk, and R. S. Muttiah, "Conditions for polarization elements to be dichroic and birefringent," J. Opt. Soc. Am. A 22, 1447-1452 (2005).
[107] M. K. Swami, S. Manhas, P. Buddhiwant, N. Ghosh, A. Uppal, and P. K. Gupta, "Polar decomposition of $3 \times 3$ Mueller matrix: a tool for quantitative tissue polarimetry," Opt. Express 14, 9324-9337 (2006).
[108] N. Ghosh, M. F. G. Wood, and A. I. Vitkin, "Mueller matrix decomposition for extraction of individual polarization parameters from complex turbid media exhibiting multiple scattering, optical activity, and linear birefringence," Journal of Biomedical Optics 13 (2008).
[109] J. Chung, W. Jung, M. J. Hammer-Wilson, P. Wilder-Smith, and Z. Chen, "Use of polar decomposition for the diagnosis of oral precancer," Appl. Opt. 46, 3038-3045 (2007).
[110] C. Collet, J. Zallat, and Y. Takakura, "Clustering of Mueller matrix images for skeletonized structure detection," Opt. Express 12, 1271-1280 (2004).
[111] S. Manhas, M. K. Swami, P. Buddhiwant, N. Ghosh, P. K. Gupta, and J. Singh, "Mueller matrix approach for determination of optical rotation in chiral turbid media in backscattering geometry," Opt. Express 14, 190-202 (2006).
[112] R. M. Wilcox, "Exponential operators and parameter differentiation in quantum physics," Journal of Mathematical Physics 8, 962-982 (1967).
[113] D. Scholz and M. Weyrauch, "A note on the Zassenhaus product formula," Journal of Mathematical Physics 47 (2006).
[114] M. Suzuki, "On the convergence of exponential operators: the Zassenhaus formula, BCH formula and systematic approximants," Communications in Mathematical Physics 57, 193-200 (1977).
[115] R. Ossikovski, "Analysis of depolarizing Mueller matrices through a symmetric decomposition," J. Opt. Soc. Am. A pp. 1109-1118 (2009).
[116] C. Fallet, A. Pierangelo, R. Ossikovski, and A. De Martino, "Experimental validation of the symmetric decomposition of Mueller matrices," Opt. Express 18, 831-842 (2010).
[117] G. E. Jellison and F. A. Modine, "Two-modulator generalized ellipsometry: theory," Appl. Opt. 36, 8190-8198 (1997).
[118] G. E. Jellison and F. A. Modine, "Two-modulator generalized ellipsometry: experiment and calibration," Appl. Opt. 36, 8184-8189 (1997).
[119] G. E. Jellison and F. Modine, Two modulator generalized ellipsometer for complete Mueller matrix measurement (1999).
[120] S. N. Jasperson and S. E. Schnatterly, "An improved method for high reflectivity ellipsometry based on a new polarization modulation technique," Review of Scientific Instruments 40, 761-767 (1969).
[121] B. Drévillon, J. Perrin, R. Marbot, A. Violet, and J. L. Dalby, "Fast polarization modulated ellipsometer using a microprocessor system for digital Fourier analysis," Review of Scientific Instruments 53, 969-977 (1982).
[122] O. Acher, E. Bigan, and B. Drévillon, "Improvements of phase-modulated ellipsometry," Review of Scientific Instruments 60, 65-77 (1989).
[123] A. Canillas, E. Pascual, and B. Drévillon, "An IR phase-modulated ellipsometer using a Fourier transform spectrometer for in situ applications," Thin Solid Films 234, 318-322 (1993).
[124] A. Canillas, E. Pascual, and E. Bertran, "Calibration improvement of Fourier transform infrared phase-modulated ellipsometry," J. Opt. Soc. Am. A 13, 24612467 (1996).
[125] P. Hauge, "Recent developments in instrumentation in ellipsometry," Surface Science 96, 108-140 (1980).
[126] J. C. Kemp, "Piezo-optical birefringence modulators: new use for a long-known effect," J. Opt. Soc. Am. 59, 950-953 (1969).
[127] J. C. Canit and J. Badoz, "New design for a photoelastic modulator," Appl. Opt. 22, 592-594 (1983).
[128] B. Wang, E. Hinds, and E. Krivoy, "Basic optical properties of the photoelastic modulator part II: residual birefringence in the optical element," (SPIE, 2009), vol. 7461, pp. 746110+.
[129] G. E. Jellison and F. A. Modine, "Accurate calibration of a photoelastic modulator in polarization modulation ellipsometry," in "Society of Photo-Optical Instrumentation Engineers (SPIE) Conference Series,", vol. 1166, R. A. Chipman, ed. (1990), vol. 1166, pp. 231-241.
[130] H. Tompkins and E. A. Irene, Handbook of Ellipsometry (Materials Science and Process Technology) (William Andrew, 2004).
[131] F. A. Modine, "Circuit for maintaining constant phototube current in polarization modulation spectroscopy," Review of Scientific Instruments 50, 386-387 (1979).
[132] H. K. Pak and B. M. Law, "2D imaging ellipsometric microscope," Review of Scientific Instruments 66, 4972-4976 (1995).
[133] A. De Martino, Y.-K. Kim, E. Garcia-Caurel, B. Laude, and B. Drévillon, "Optimized Mueller polarimeter with liquid crystals," Opt. Lett. 28, 616-618 (2003).
[134] P. S. Hauge, "Mueller matrix ellipsometry with imperfect compensators," J. Opt. Soc. Am. 68, 1519-1528 (1978).
[135] J. S. Baba, J.-R. R. Chung, A. H. DeLaughter, B. D. Cameron, and G. L. Coté, "Development and calibration of an automated Mueller matrix polarization imaging system." Journal of biomedical optics 7, 341-349 (2002).
[136] E. Compain and B. Drévillon, "Complete high-frequency measurement of Mueller matrices based on a new coupled-phase modulator," Review of Scientific Instruments 68, 2671-2680 (1997).
[137] R. W. Collins and J. Koh, "Dual rotating-compensator multichannel ellipsometer: instrument design for real-time Mueller matrix spectroscopy of surfaces and films," J. Opt. Soc. Am. A 16, 1997-2006 (1999).
[138] C. Chen, I. An, G. M. Ferreira, N. J. Podraza, J. A. Zapien, and R. W. Collins, "Multichannel Mueller matrix ellipsometer based on the dual rotating compensator principle," Thin Solid Films 455-456, 14-23 (2004).
[139] F. Delplancke, "Automated high-speed Mueller matrix scatterometer," Appl. Opt. 36, 5388-5395 (1997).
[140] E. Garcia-Caurel, A. De Martino, and B. Drévillon, "Spectroscopic Mueller polarimeter based on liquid crystal devices," Thin Solid Films 455-456, 120-123 (2004).
[141] A. De Martino, "General methods for optimized design and calibration of Mueller polarimeters," Thin Solid Films 455-456, 112-119 (2004).
[142] J. O. Stenflo, "Solar polarimetry with ZIMPOL. Plans for the future," Memorie della Societa Astronomica Italiana 78 (2007).
[143] R. M. A. Azzam, "Simulation of mechanical rotation by optical rotation: Application to the design of a new Fourier photopolarimeter," J. Opt. Soc. Am. 68, 518-521 (1978).
[144] O. Arteaga and et. al, "Chiral biases in solids by effect of shear gradients: A speculation on the deterministic origin of biological homochirality," Origins of Life (2009).
[145] J. F. Nye, Physical Properties of Crystals: Their Representation by Tensors and Matrices (Oxford University Press, USA, 1985).
[146] G. E. Jellison, "Windows in ellipsometry measurements." Applied optics 38, 47844789 (1999).
[147] S. Chandrasekhar, "Optical Rotatory Dispersion of Crystals," Proceedings of the Royal Society of London. Series A, Mathematical and Physical Sciences 259, 531553 (1961).
[148] T. Yoshizawa, ed., Handbook of Optical Metrology: Principles and Applications (CRC, 2009), 1st ed.
[149] C. Y. Han and Y. F. Chao, "Photoelastic modulated imaging ellipsometry by stroboscopic illumination technique," Review of Scientific Instruments 77, 023107+ (2006).
[150] P. Gleyzes, A. C. Boccara, and H. Saint-Jalmes, "Multichannel Nomarski microscope with polarization modulation: performance and applications," Opt. Lett. 22, 1529-1531 (1997).
[151] H.-M. Tsai and Y.-F. Chao, "Optimization of a four-temporal phase lock for photoelastic-modulated polarimetry," Opt. Lett. 34, 2279-2281 (2009).
[152] D. J. Diner, A. Davis, B. Hancock, S. Geier, B. Rheingans, V. Jovanovic, M. Bull, D. M. Rider, R. A. Chipman, A.-B. Mahler, and S. C. McClain, "First results from a dual photoelastic-modulator-based polarimetric camera," Appl. Opt. 49, 2929-2946 (2010).
[153] T. M. Lowry, "Optical Rotatory Dispersion. Part I: The Natural and Magnetic Rotatory Dispersion in Quartz of Light in the Visible Region of the Spectrum," Royal Society of London Philosophical Transactions Series A 212, 261-297 (1913).
[154] C. Chou, Y.-C. Huang, and M. Chang, "Precise optical activity measurement of quartz plate by using a true phase-sensitive technique," Appl. Opt. 36, 3604-3609 (1997).
[155] D. Yogev-Einot and D. Avnir, "The temperature-dependent optical activity of quartz: from Le Châtelier to chirality measures," Tetrahedron: Asymmetry 17, 2723-2725 (2006).
[156] M. B. Myers and K. Vedam, "Effect of pressure on the optical rotatory power and dispersion of crystalline sodium chlorate," J. Opt. Soc. Am. 57, 1146-1148 (1967).
[157] J. R. L. Moxon, A. R. Renshaw, and I. J. Tebbutt, "The simultaneous measurement of optical activity and circular dichroism in birefringent linearly dichroic crystal sections. II. Description of apparatus and results for quartz, nickel sulphate hexahydrate and benzil," Journal of Physics D: Applied Physics 24, 1187-1192 (1991).
[158] C. Hernandez-Rodriguez and P. Gomez-Garrido, "Optical anisotropy of quartz in the presence of temperature-dependent multiple reflections using a high-accuracy universal polarimeter," Journal of Physics D: Applied Physics 33, 2985-2994 (2000).
[159] O. Arteaga, A. Canillas, and G. E. Jellison, "Determination of the components of the gyration tensor of quartz by oblique incidence transmission two-modulator generalized ellipsometry," Appl. Opt. 48, 5307-5317 (2009).
[160] G. L. Tan, M. F. Lemon, D. J. Jones, and R. H. French, "Optical properties and London dispersion interaction of amorphous and crystalline $\mathrm{SiO}_{2}$ determined by vacuum ultraviolet spectroscopy and spectroscopic ellipsometry," Physical Review B 72, 205117+ (2005).
[161] G. Szivessy and C. Münster, "Über die prüfung der gitteroptik bei aktiven kristallen," Annalen der Physik 412, 703-736 (1934).
[162] M. Born, Optik: Ein Lehrbuch der elektromagnetische Lichttheorie (Springer, 1933).
[163] A. Konstantinova, B. Nabatov, E. Evdishchenko, and K. Konstantinov, "Modern application packages for rigorous solution of problems of light propagation in anisotropic layered media: II. Optically active crystals," Crystallography Reports 47, 815-823 (2002).
[164] Konstantinova, A., Evdishchenko, E., Imangazieva, and K., "Manifestation of optical activity in crystals of different symmetry classes," Crystallography Reports 51, 998-1008 (2006).
[165] D. Eimerl, "Quantum electrodynamics of optical activity in birefringent crystals," J. Opt. Soc. Am. B 5, 1453-1461 (1988).
[166] G. Ghosh, "Dispersion-equation coefficients for the refractive index and birefringence of calcite and quartz crystals," Optics Communications 163, 95-102 (1999).
[167] T. Bradshaw and G. H. Livens, "The formula for the optical rotatory dispersion of quartz," Proceedings of the Royal Society of London. Series A, Containing Papers of a Mathematical and Physical Character 122, 245-250 (1929).
[168] S.-L. Lu and A. P. Loeber, "Depolarization of white light by a birefringent crystal," J. Opt. Soc. Am. 65, 248+ (1975).
[169] G. E. Jellison, C. O. Griffiths, D. E. Holcomb, and C. M. Rouleau, "Transmission two-modulator generalized ellipsometry measurements." Applied optics 41, 65556566 (2002).
[170] O. Arteaga and A. Canillas, "Pseudopolar decomposition of the Jones and MuellerJones exponential polarization matrices," J. Opt. Soc. Am. A 26, 783-793 (2009).
[171] G. E. Jellison and C. M. Rouleau, "Determination of optical birefringence by using off-axis transmission ellipsometry," Appl. Opt. 44, 3153-3159 (2005).
[172] J. Jerphagnon and D. S. Chemla, "Optical activity of crystals," The Journal of Chemical Physics 65, 1522-1529 (1976).
[173] A. Konstantinova, K. Rudoy, B. Nabatov, E. Evdishchenko, V. Stroganov, and O. Pikul', "The influence of optical activity on the intensity and polarization parameters of transmitted light in crystals," Crystallography Reports 48, 823-831 (2003).
[174] C. Honda and H. Hada, "Circular dichroism of poly-molecular associate, Jaggregate, of 1,12-diethyl-2,22-cyanine chloride by regular stirring of the solution," Tetrahedron Letters 17, 177-180 (1976).
[175] O. Ohno, Y. Kaizu, and H. Kobayashi, "J-aggregate formation of a water-soluble porphyrin in acidic aqueous media," The Journal of Chemical Physics 99, 4128+ (1993).
[176] J. M. Ribo, J. Crusats, F. Sagues, J. Claret, and R. Rubires, "Chiral sign induction by vortices during the formation of mesophases in stirred solutions," Science 292, 2063-2066 (2001).
[177] A. D'Urso, R. Randazzo, L. Lo Faro, and R. Purrello, "Vortexes and nanoscale chirality," Angewandte Chemie International Edition 49, 108-112 (2010).
[178] A. Tsuda, Md, T. Harada, T. Yamaguchi, N. Ishii, and T. Aida, "Spectroscopic visualization of vortex flows using dye-containing nanofibers," Angewandte Chemie International Edition 46, 8198-8202 (2007).
[179] M. Wolffs, S. J. George, Z. Tomović, S. C. Meskers, A. P. Schenning, and E. W. Meijer, "Macroscopic origin of circular dichroism effects by alignment of self-assembled fibers in solution." Angewandte Chemie (International ed. in English) 46, 82038205 (2007).
[180] O. Arteaga, C. Escudero, G. Oncins, Z. El-Hachemi, J. Llorens, J. Crusats, A. Canillas, and J. M. Ribó, "Reversible mechanical induction of optical activity in solutions of soft-matter nanophases," Chemistry - An Asian Journal 4, 1687-1696 (2009).
[181] C. Escudero, J. Crusats, I. Díez-Pérez, Z. El-Hachemi, and J. M. Ribó, "Folding and hydrodynamic forces in J-aggregates of 5 -phenyl-10,15,20-tris(4sulfophenyl)porphyrin13," Angewandte Chemie 118, 8200-8203 (2006).
[182] H. Fenniri, B.-L. L. Deng, and A. E. Ribbe, "Helical rosette nanotubes with tunable chiroptical properties." Journal of the American Chemical Society 124, 1106411072 (2002).
[183] J. G. Moralez, J. Raez, T. Yamazaki, R. K. Motkuri, A. Kovalenko, and H. Fenniri, "Helical rosette nanotubes with tunable stability and hierarchy," Journal of the American Chemical Society 127, 8307-8309 (2005).
[184] R. S. Johnson, T. Yamazaki, A. Kovalenko, and H. Fenniri, "Molecular basis for water-promoted supramolecular chirality inversion in helical rosette nanotubes," Journal of the American Chemical Society 129, 5735-5743 (2007).
[185] A. Y. Elezzabi and S. Sederberg, "Optical activity in an artificial chiral media: a terahertz time-domain investigation of Karl F. Lindman's 1920 pioneering experiment," Opt. Express 17, 6600-6612 (2009).
[186] K. F. Lindman, "Über eine durch ein isotropes system von spiralförmigen resonatoren erzeugte rotationspolarisation der elektromagnetischen wellen," Annalen der Physik 368, 621-644 (1920).
[187] G. S. Ranganath and S. Ramaseshan, "Piezo-rotatory coefficients and crystal symmetry," J. Opt. Soc. Am. 59, 1229-1232 (1969).
[188] D. K. Kondepudi, R. J. Kaufman, and N. Singh, "Chiral symmetry breaking in sodium chlorate crystallizaton," Science 250, 975-976 (1990).
[189] C. Viedma, "Chiral symmetry breaking during crystallization: Complete chiral purity induced by nonlinear autocatalysis and recycling," Physical Review Letters 94, 065504+ (2005).
[190] N. K. Chaudhuri and M. A. El-Sayed, "Molecular origin of the optical rotatory dispersion of the benzil crystal," J. Chem. Phys. 47, 1133-1143 (1967).
[191] J. Ríha, I. Vysín, and H. Lapanská, "Theory, measurement, and origin of optical activity in benzil crystal," Molecular Crystals and Liquid Crystals 442, 181-201 (2005).
[192] D. Hochberg and M. Zorzano, "Reaction-noise induced homochirality," Chemical Physics Letters 431, 185-189 (2006).
[193] A. Giaquinta and D. Hochberg, "Chiral symmetry breaking: (I) limited enantioselectivity and (II) mutual inhibition," Physica D: Nonlinear Phenomena 237, 2563-2576 (2008).
[194] O. Arteaga, A. Canillas, J. Crusats, Z. El-Hachemi, G. Jellison, J. Llorca, and J. Ribó, "Chiral biases in solids by effect of shear gradients: A speculation on the deterministic origin of biological homochirality," Origins of Life and Evolution of Biospheres 40, 27-40 (2010).
[195] K. Kvenvolden, J. Lawless, K. Pering, E. Peterson, J. Flores, C. Ponnamperuma, I. R. Kaplan, and C. Moore, "Evidence for extraterrestrial amino-acids and hydrocarbons in the Murchison meteorite," Nature 228, 923-926 (1970).
[196] G. Cooper, N. Kimmich, W. Belisle, J. Sarinana, K. Brabham, and L. Garrel, "Carbonaceous meteorites as a source of sugar-related organic compounds for the early Earth," Nature 414, 879-883 (2001).
[197] S. Pizzarello, Y. Huang, and M. Fuller, "The carbon isotopic distribution of Murchison amino acids," Geochimica et Cosmochimica Acta 68, 4963-4969 (2004).
[198] D. P. Glavin and J. P. Dworkin, "Enrichment of the amino acid l-isovaline by aqueous alteration on CI and CM meteorite parent bodies," Proceedings of the National Academy of Sciences 106, 5487-5492 (2009).
[199] T. Kawasaki, K. Hatase, Y. Fuj, K. Jo, K. Soai, and S. Pizzarello, "The distribution of chiral asymmetry in meteorites: An investigation using asymmetric autocatalytic chiral sensors," Geochimica et Cosmochimica Acta 70, 5395-5402 (2006).
[200] T. Zega, "Serpentine nanotubes in the Mighei CM chondrite," Earth and Planetary Science Letters 223, 141-146 (2004).
[201] G. E. Christidis, F. Dellisanti, G. Valdre, and P. Makri, "Structural modifications of smectites mechanically deformed under controlled conditions," Clay Minerals 40, 511-522 (2005).
[202] M. P. Silverman, J. Badoz, and B. Briat, "Chiral reflection from a naturally optically active medium," Opt. Lett. 17, 886-888 (1992).
[203] A. Ghosh and P. Fischer, "Chiral molecules split light: Reflection and refraction in a chiral liquid," Physical Review Letters 97, 173002+ (2006).
[204] D. Lara and C. Dainty, "Axially resolved complete Mueller matrix confocal microscopy," Appl. Opt. 45, 1917-1930 (2006).
[205] Z. El-Hachemi, O. Arteaga, A. Canillas, J. Crusats, C. Escudero, R. Kuroda, T. Harada, M. Rosa, and J. M. Ribó, "On the mechano-chiral effect of vortical flows on the dichroic spectra of 5 -phenyl-10,15,20-tris(4-sulfonatophenyl) porphyrin Jaggregates," Chemistry - A European Journal 14, 6438-6443 (2008).


[^0]:    ${ }^{1}$ Sometimes the denomination of ellipticity is given to $e= \pm \tan \epsilon$ where the + and - signs correspond to right- and left-handed polarizations, respectively. In these cases $\epsilon$ is referred as ellipticity angle.

[^1]:    ${ }^{2}$ The Hermitian adjoint of a matrix is the complex conjugate of the transpose of the matrix, thus $\mathbf{E}^{\dagger}$ is a row vector.

[^2]:    ${ }^{3}$ This definition of positive azimuthal rotation angles is not universal in optics. There are authors that use the opposite convention and, sometimes, this is not clearly noticed. If the opposite convention was used the Jones matrix of rotation would be $\mathbf{R}(\alpha)=\left[\begin{array}{cc}\cos \alpha & -\sin \alpha \\ \sin \alpha & \cos \alpha\end{array}\right]$.

[^3]:    ${ }^{4}$ See Refs. [4] and [5] for the explicit equations.

[^4]:    ${ }^{5}$ These two modes are sometimes referred to as eigenmodes, because radiation linearly polarized according to any of these two modes will not be changed when going through the medium. For any other direction of polarization, or for circular polarization, the polarization form will change as the radiation passes through a linearly birefringent medium

[^5]:    ${ }^{6}$ See section 3.2.3 for more details.

[^6]:    ${ }^{a} n$ stands for refractive index, $k$ for the extinction coefficient, $l$ for path length through the medium, and $\lambda$ for the vacuum wavelength of light. Subscripts specify the polarization of light as, $x, y, 45^{\circ}$ to the $x$ axis, $135^{\circ}$ to the $x$ axis, circular right + , or left.-

[^7]:    ${ }^{1}$ The form of the Drude equation can be directly derived from Eq. (1.51) if one asumes a Sellmeier dispersion for the refractive index: $n^{2}(\lambda)=1+\sum_{i} \frac{B_{i} \lambda^{2}}{\lambda^{2}-C_{i}} . B_{i}$ would be slightly different for left- and right-circularly polarized light.

[^8]:    ${ }^{2}$ Point groups are mathematical groups of geometric symmetries. In chemistry they are used to describe the symmetries of a molecule.
    ${ }^{3} \mathrm{~A}$ rotation-reflection axis $n>2$ means a rotation around an axis by $\frac{360^{\circ}}{n}$, followed by a reflection in a plane perpendicular to it. $n$ must be an even number $>2$.

[^9]:    ${ }^{4}$ There also exist aggregates which show a blue-shifted absorption spectrum (compared to the monomer), which is broader and less visible. These aggregates are called H-aggretates

[^10]:    ${ }^{5}$ See section 5.2 for more details about the characteristics of photoelastic modulators
    ${ }^{6}$ www.jascoinc.com

[^11]:    ${ }^{7}$ The influence that linear birefringence has in the measurement of CB is much more evident than in case of CD and it was a known topic since the first experiments about optical rotation of quartz in the XIX century. The measurement of CB was considered to be not possible in presence of linear birefringence.

[^12]:    ${ }^{1}$ A medium with optical activity is often referred in theoretical works as a gyrotropic medium
    or a medium with gyrotropy

[^13]:    ${ }^{2}$ See section 1.1 for an introductory discussion about the constitutive relations

[^14]:    ${ }^{3}$ Note that in the context of this work the words "diattenuation" and "dichroism" are synonyms.
    The same is also applicable to the pair "birefringence" and "retardation"

[^15]:    ${ }^{1}$ This sum factorization is sometimes referred to as spectral decomposition, see Refs. [6,100] for more details

[^16]:    ${ }^{2}$ Frequently depolarizers are described with diagonal Mueller matrices. However note that Lu-Chipman decomposition uses a depolarizer with depolarizance, which, in general, has a non-diagonal Mueller matrix.

[^17]:    ${ }^{3}$ The Frobenius norm of a $n \times n$ matrix $\mathbf{X}$ is defined as $\|\mathbf{X}\|=\left(\sum_{i=0}^{n-1} \sum_{j=0}^{n-1}\left|x_{i j}\right|^{2}\right)^{1 / 2}=$ $\left[\operatorname{Tr}\left(\mathbf{X}^{*} \mathbf{X}\right)\right]^{1 / 2}$

[^18]:    ${ }^{1} \sin \theta_{i}=\theta_{i}$ and $\cos \theta_{i}=1$.

[^19]:    ${ }^{2} \cos \theta \cos \varphi=[\cos (\theta-\varphi)+\cos (\theta+\varphi)] / 2$,
    $\sin \theta \sin \varphi=[\cos (\theta-\varphi)-\cos (\theta+\varphi)] / 2$,
    $\sin \theta \cos \varphi=[\sin (\theta+\varphi)+\sin (\theta-\varphi)] / 2$.

[^20]:    ${ }^{1}$ They are usually made of quartz and sold under the name "quartz rotators".

[^21]:    ${ }^{1}$ There are different notations for crystallograpic point group. Using the Schönflies quartz belongs to $D_{3}$ point group

[^22]:    ${ }^{1}$ The details about the preparation of the aggregate solutions can be checked in Refs. [180,181].

[^23]:    ${ }^{2}$ In section 2.2.2 we showed how the excitonic transitions between chromophores are responsible for the CD signals.

[^24]:    ${ }^{1}$ Note that in the experiments by Kondepudi and Viedma there are two components: $\mathrm{NaClO}_{3}$ and $\mathrm{H}_{2} \mathrm{O}$.

[^25]:    ${ }^{2}$ Carbonaceous chondrites are a class of chondritic meteorites. They include some of the most primitive known meteorites, but represent only a small proportion $\sim 4.6 \%$ of meteorite falls.
    ${ }^{3} \mathrm{CM} 2$ carbonaceous chondrites are composed of about $70 \%$ fine-grained material (matrix), and most have experienced extensive aqueous alteration

[^26]:    ${ }^{4}$ Poorly characterized phases are usually fine-grain ( $<1 \mu \mathrm{~m}$ in size) areas which are difficult to characterize with optical or X-ray techniques

[^27]:    ${ }^{1}$ En aquest resum en català hem optat per mantenir els mateixos acrònims d'origen anglès emprats al llarg de tots els capítols anteriors de la tesi.

[^28]:    ${ }^{2}$ Un medi que no introdueix despolarització és aquell pel qual la llum que incideix totalment polaritzada continua estan totalment polaritzada després després d'interactuar amb el medi. L'efecte d'un medi que despolaritzi no es pot descriure amb una matriu de Jones.

[^29]:    ${ }^{3}$ La matriu de Mueller corresponent a una matriu de Jones sovint s'anomena matriu de MuellerJones i és una matriu de Mueller que no introdueix despolarització. Tota matriu de Jones sempre té una matriu de Mueller equivalent, però l'invers no és cert.

