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Filtering of matter symmetry properties by circularly polarized nonlinear optics

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We propose a direct readout of symmetry information in matter using nonlinear optics. From combinations of circularly and longitudinally polarized optical fields, we construct irreducible spherical field tensors for second- and third-order nonlinear processes. The coupling of these field tensors to the matter susceptibility tensors allows filtering out of the susceptibility symmetries independently of the sample orientation in the laboratory frame. Experimental demonstrations are conducted on microcrystals, in a microscopy configuration compatible with symmetry order imaging.

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I. INTRODUCTION

Acquiring structural information in an assembly of atoms and molecules requires the knowledge of its symmetry elements, which are fundamental building blocks of matter. Using symmetry as an observable appears therefore as an optimal way to probe the structural properties of complex materials and biological processes, which would be moreover interesting to obtain at high spatial and temporal resolution. Reading out of symmetry properties of matter is possible using nonlinear optics, taking advantage of the intrinsic relation between symmetry and nonlinear susceptibilities [1–3].

Second-order harmonic generation (SHG), for instance, is known to be allowed only in materials without symmetry inversion, whereas third-order nonlinearities, such as third-order harmonic generation (THG), four-wave mixing (FWM), or coherent Raman scattering (CRS) can occur in materials belonging to isotropic or centrosymmetric crystallographic groups [4]. Moreover, nonlinear optics offers the possibility of probing high-order symmetry information, thanks to the multiple fields’ coupling mechanisms involving a large number of degrees of freedom in terms of light polarization states [5].

A large amount of work has been carried out to relate formally the probed nonlinear susceptibility tensors with the symmetry properties of matter [1,6,7]. Nonlinear Cartesian tensors can be expanded in their spherical representation in such a way that their components transform under proper rotations similarly to spherical harmonics of symmetry degree l and order m [2,8–10]. Following this spherical representation, harmonic light scattering signals in liquids have been used to extract the symmetry components of molecular nonlinear hyperpolarizability tensors [1,3,11], allowing the development of an original molecular engineering strategy that has led to materials with optimized nonlinear responses [12,13]. While successful for understanding molecular-scale properties, this approach has apparently not yet been applied to molecular assemblies: second-order [14,15] and third-order [16–18] Cartesian susceptibility tensors of molecular materials and biological samples are usually probed by sets of linear polarization states, which is time consuming and necessitates disentangling complex Cartesian expressions to retrieve information on molecular orientations and symmetry orders.

II. THEORY

A. Filtering process

The electric nonlinear polarization is defined by the constitutive relationship

\[ \vec{P} = \epsilon_0 \vec{\chi}^{(n)} : \vec{E}_1 \otimes \cdots \otimes \vec{E}_n, \]

where \( \vec{\chi}^{(n)} \) denotes the nonlinear susceptibility tensor of order \( n \), \( \vec{E}_1,\ldots,n \) are the vectorial fields involved in the nonlinear interaction, and \( \otimes \) denotes the tensorial product. Exhibiting the norm of \( \vec{P} \) and the complex unitary vectors \( \hat{e}_i \) such as \( \vec{P} = P_0 \hat{e}_0 \) and \( \vec{E}_i = E_i \hat{e}_i \), Eq. (1) becomes

\[ P_0 = \epsilon_0 (E_1, \ldots, E_n) \vec{\chi}^{(n)} \cdot [\hat{e}_0^* \otimes \hat{e}_1 \otimes \cdots \otimes \hat{e}_n], \]

where \( \cdot \) denotes the tensorial scalar product and \( \hat{e}_0^* \) is the complex conjugate of the dipole polarization. Here all the polarization dependencies are collected in a unique “polarization tensor” [10,11]

\[ \vec{F}^{(n)} = \hat{e}_0^* \otimes \hat{e}_1 \otimes \cdots \otimes \hat{e}_n, \]

showing that \( P_0 \) is finally the projection of the susceptibility tensor \( \vec{\chi}^{(n)} \) on the polarization tensor \( \vec{F}^{(n)} \).

In what follows, we show how this projection leads to a filtering out of the information on matter symmetry. Since symmetry must be decoupled from any orientation information, we use the irreducible spherical formalism where

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tensors are expanded on an orthonormal basis exhibiting rotational invariance. This representation allows identification of tensorial quantities based on pure symmetry considerations rather than a complex mixture of orientation and symmetry, such as is found in a Cartesian basis \([1, 8]\). An irreducible spherical \(n\)-order tensor is labeled by its symmetry order \(l \leq n + 1\) and possesses \(m = 2l + 1\) independent components, related to the spherical harmonics \(Y_{lm}\) [8, 20]. The coefficient \(m\) can moreover be interpreted as the symmetry order of the tensor into the sample plane \((x, y)\), \(z\) being the direction of propagation of the excitation and detected fields. In such a representation, \(\tilde{\chi}^{(n)}\) is decomposed on spherical-basis-like components \(\tilde{C}_m\), as introduced in [2, 21], with amplitudes \(\tilde{m}_m\).

\[
\tilde{\chi}^{(n)} = \sum_{lm} \tilde{m}_m \tilde{C}_m \tag{4}
\]

**B. Irreducible field tensor**

Similarly to the \(\tilde{\chi}^{(n)}\) tensor, the field tensor \(\tilde{F}^{(n)}\) can be expanded under this irreducible spherical formalism. To calculate the spherical components of \(\tilde{F}^{(n)}\), each individual field polarization \(\hat{e}_l\) of Eq. (3) must be represented in a spherical basis preserving the rotational invariance. The development below details the buildup of irreducible spherical representations for second- and third-order processes. For \(l = 1\), the three independent vectors \(\hat{e}_m\) with \(m = (-1, 0, 1)\) are expressed in the Cartesian frame as [1]

\[
\begin{align*}
\hat{e}_{m=+1} &= \frac{1}{\sqrt{2}} (\hat{e}_x + i \hat{e}_y), \\
\hat{e}_{m=-1} &= \frac{1}{\sqrt{2}} (\hat{e}_x - i \hat{e}_y), \\
\hat{e}_{m=0} &= -i \hat{e}_z.
\end{align*}
\]

These field vector components coincide with the in-plane left circularly polarized field \((m = +1)\), right circularly polarized field \((m = -1)\), and axial polarization field along \(z\) \((m = 0)\), which are all invariant through rotations around the \(z\) axis. The irreducible representation of higher-order field tensors, built from this rank-1 expression in Eq. (3), have been described in the context of nonlinear optics by Makek [1]. The first step to expressing the rank-3 polarization tensor is to couple three irreducible tensors of rank one using the Clebsch-Gordan coefficients. A second step is required to take into account the different way of coupling the fields with respect to the symmetry of index permutation, using the \(\tilde{\gamma}_j\) Wigner coefficients. We limit this study to symmetric tensors, in which all the index permutations are allowed. Following this operation, the spherical field tensor basis components involved in symmetric nonlinear second-order processes (rank 3) are given by

\[
\tilde{F}_m^{(3)} = \sum_{m_0, m_1, m_2} \tilde{U}_l \tilde{m}_0 \tilde{m}_1 \tilde{m}_2 \left( \hat{e}_{m_0} \otimes \hat{e}_{m_1} \otimes \hat{e}_{m_2} \right)
\]

\[
= \sum_{m_0, m_1, m_2} \mathcal{M}_{m_0 m_1 m_2}^{l} \left( \hat{e}_{m_0} \otimes \hat{e}_{m_1} \otimes \hat{e}_{m_2} \right) \tag{6}
\]

where \(\tilde{m}_m\) is related to \(Y_{lm}\) by a phase factor \(i^l\).

1 Using the convention of Eq. (5), \(\tilde{F}_m^{(3)}\) is related to \(Y_{lm}\) by a phase factor \(i^l\).
TABLE I. $\mathbb{M}_{m_{0}...m_{n}}$ values relating the field polarization tensorial products (rows) and the spherical irreducible tensors $F_{m}^{l}$ (columns). No value means zero. A square root sign is to be understood over every coefficient, e.g., $\mathbb{M}_{m_{0}=0,m_{1}=1,m_{2}=1}^{n} = \sqrt{17}$. Due to the symmetric construction of the irreducible spherical tensor, permutations are allowed and, for instance $\hat{e}_{1} \otimes \hat{e}_{0} \otimes \hat{e}_{1} = \hat{e}_{0} \otimes \hat{e}_{1} \otimes \hat{e}_{0} = \sqrt{4/15} F_{0}^{1} - \sqrt{1/15} F_{1}^{1}$. 

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that in the case of nonlinear rank-3 processes (SHG, etc.), two symmetry orders are accessible: $\xi_{1}$ (dipolar contribution) and $\xi_{3}$ (octupolar contribution) [3], which both exhibit non-centrosymmetry as expected. Rank-4 processes (THG, FWM, CRS, etc.) give access to isotropic $\xi_{0}$, second $\xi_{2}$, and fourth $\xi_{4}$ orders. Table I can be used to identify which circular field combination leads to these orders, paying attention to the presence of conjugated fields in the nonlinear processes, which imply a change of the $m_{i}$ sign (since $\hat{e}_{m_{i}} = -\hat{e}_{-m_{i}}$). For instance, the filtering process which extracts $\xi_{0}$ in Eq. (9) will require the set $\hat{e}_{-1} \hat{e}_{1} \hat{e}_{-1} \hat{e}_{1}$ according to Table I. But due to the complex conjugates of the emitted field $m_{0}$ and the Stokes field $m_{3}$ in FWM, the experimental set of polarizations used to probe the isotropic order $\xi_{0}$ must be $\hat{e}_{1} \hat{e}_{1} \hat{e}_{1} \hat{e}_{1}$. Using the same experimental set, the third-order harmonic generation is sensitive to $\xi_{2}$ instead of $\xi_{0}$ [23–25].

III. EXPERIMENTAL RESULTS

A. Experimental setup

To experimentally illustrate nonlinear light-matter symmetry filtering, microscopy experiments were conducted on microcrystals using circular polarizations. Microcrystals of 1,3,5-tricyano-2,4,6-tris(p-diethyl-aminostyryl)bezene (TTB) [26] and NaCl were imaged using second-order (SHG) and third-order (FWM) nonlinear processes. The TTB crystals were obtained by evaporation [26] and NaCl crystals were prepared by spraying saturated salt water on a cover glass. The crystal samples were covered with oil for index-matching purposes. The setup for polarized SHG and FWM microscopy is based on previously reported works [14,17]. Briefly, the light is focused on the sample using an objective lens ($\times 40$ numerical aperture 0.6). Images are obtained by moving the sample with a piezoscanning stage and the emitted signal is collected in the forward direction. The circular excitation polarization states are implemented using a unique achromatic quarter-wave plate just before the objective, the control of its left- or right-circular nature being performed beforehand using appropriate half-wave plates in the excitation paths involved. The emitted light is analyzed using an achromatic quarter-wave plate oriented at 45° with respect to a Wollaston prism which separates the right and left circularly polarized light on two avalanche photodiodes (APDs). We denote by $I_{m}$ the intensity obtained by a set of excitation and detection circular polarizations such as $m = m_{0} + \cdots + m_{n}$.

B. Results

Figure 1 shows $I_{1}$ and $I_{3}$ images obtained with SHG on TTB microcrystals of needle shapes. These molecular crystals are known to be octupolar crystals oriented perpendicularly to their major axis [26]. Due to the crystal orientations, the planar octupolar symmetry is perpendicular to the propagation axis, which leads to unidirectional symmetry in the sample plane. From the intensities $I_{1}$ and $I_{3}$ the ratio $\xi_{3}/\xi_{1} = \sqrt{T_{3}/T_{1}}$ is calculated for each pixel of the image and summarized in a histogram; see Fig. 1(d). Due to the field-invariant rotation along the $z$ axis, the information extracted is also expected

FIG. 1. (Color online) (a) SHG raw images of TTB crystals with dipolar ($\hat{e}_{1} \hat{e}_{1} \hat{e}_{1}$) and octupolar ($\hat{e}_{-1} \hat{e}_{1} \hat{e}_{1}$) polarization sets. (b) White light image and selected area for polarization-resolved SHG images. (c) Ratio between third and first orders (see text) and (d) its histogram. Scale bars: 10 $\mu$m.
to be invariant with respect to the orientation of the sample in the sample plane. Except for some specific regions which could be attributed to crystal defects, the value of \( \xi_3/\xi_1 \) is indeed confirmed to be homogeneous over the whole crystal. Moreover, the \( \xi_3/\xi_1 \) ratio is close to 1, which leads to a similar weight for the dipolar and octupolar contributions, as expected for unidirectional crystals [11].

FWM images of NaCl salt microcrystals are depicted in Fig. 2. The different sets of circular polarizations lead to different information: the isotropic image \( I_0 \) shows a strong signal from the crystal and the oil. The \( I_2 \) image shows a very weak signal, two orders of magnitude smaller than \( I_0 \) (this residual signal is probably due to remaining imperfections in the circular excitation and detection states). Finally, the \( I_4 \) image shows a dominant, very strong signal in the NaCl crystal. This is expected from the fourfold symmetry of NaCl, whereas the oil is an isotropic liquid. As previously, the \( I_4 \) magnitude does not depend on the crystal orientation, the ratio \( \xi_4/\xi_0 = \sqrt{I_4/I_0} \) being the same for all crystals present in the image; see Fig. 2(c). The \( \xi_4/\xi_0 \) ratio gives, moreover, quantitative information about the relative weight of fourth order and zero (isotropic) orders and is shown to be a fundamental symmetry property of NaCl crystals [17]. Note finally that the fourth-order measurement \( (I_4) \) allows complete rejection of the NaCl isotropic contribution present in the crystals. This scheme, in addition to providing contrasted images, is a unique way to filter out symmetry orders without the need of postprocessing.

**IV. CONCLUSION**

We have applied an irreducible spherical formalism to both fields and matter to express their coupling in nonlinear optics processes. Proper combinations of circular polarization fields were exhibited in order to filter out individually the accessible in-plane symmetry orders present in the probed matter. This readout process reveals quantitative symmetry information regardless of the in-plane orientation of the sample in the measurement plane. This scheme requires finally single, well-defined excitation and detection schemes, which makes it compatible with fast imaging of dynamical processes.

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**APPENDIX: MATRICES U AND V**

Here we give details of the matrices \( U \) and \( V \) for the complete symmetric set \([1,22]\).

<table>
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<th>( I_4 )</th>
<th>( I_1 )</th>
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<tr>
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