Article

Selection rules for all-optical magnetic recording in iron garnet

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Rapid growth of the area of ultrafast magnetism has allowed to achieve a substantial progress in all-optical magnetic recording with femtosecond laser pulses and triggered intense discussions about microscopic mechanisms responsible for this phenomenon. The typically used metallic medium nevertheless considerably limits the applications because of the unavoidable heat dissipation. In contrast, the recently demonstrated photo-magnetic recording in transparent dielectric garnet for all practical purposes is dissipation-free. This discovery raised question about selection rules, i.e. the optimal wavelength and the polarization of light, for such a recording. Here we report the computationally and experimentally identified workspace of parameters allowing photo-magnetic recording in Co-doped iron garnet using femtosecond laser pulses. The revealed selection rules indicate that the excitations responsible for the coupling of light to spins are d-d electron transitions in octahedral and tetrahedral Co-sublattices, respectively.

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Ultrafast control of the electron spin in the matter with the help of ultrashort light pulses has evolved into one of the hottest topics of the fundamental magnetism and generated significant interest in the area of information technology. In particular, the fact that magnetization can be reversed by femtosecond laser pulses may result in new concepts of storage devices with unprecedented bit rates. Speaking in general, moreover, finding an electronic transition a subtle excitation of which can launch dramatic changes of electric, optical or magnetic properties of media is one of the long-standing dreams in the field of photo-induced phase transitions. This field got a great boost due to the development of new femtosecond laser sources allowing to explore photo-induced phase transitions in a broad spectral range and on a time scale of elementary processes in condensed matter such as electron–electron, electron–lattice, spin–lattice interactions, period of oscillations of lattice atoms, or spins. The interest to magnetic photo-induced phase transitions has been continuously fueled due to its relevance to magnetic recording technology and the discovery of the magnetization switching by a femtosecond laser pulse alone triggered intense discussions about mechanisms responsible for these laser-induced changes. For a long time practically all research on all-optical magnetic switching was limited to metals and relied on mechanisms directly related to laser-induced heating close to the Curie temperature. Ultrafast light–matter interactions in this case are therefore explained with the help of a simplistic temperature model, where features of the electronic structure hardly play any role. However, the most intriguing pathways to change the magnetization with a laser pulse are those that otherwise leave the entropy of the whole system almost intact. Such precisely directed impact has the potential to reduce energy waste and also to provide novel possibilities for ultrafast transfer of recorded data in the optical telecommunication systems using ultrashort laser pulses. Recently it has indeed been shown that using a single 50 fs laser pulse one can permanently switch the magnetization in Co-doped yttrium iron garnet thin film (YIG:Co) with (001) plane of the sample. In this material, the Co-dopants are responsible for strong magnetocrystalline anisotropy, making cube diagonals <111> to be eight easy directions. It was experimentally demonstrated in our previous work that optical excitation of localized d–d transition in Co-ions at 0.95 eV with light polarized along [1 00] or [0 10] axis was able to modify the magnetic anisotropy and thus to switch the magnetization between the metastable states. The presented results raise questions about selection rules e.g., the feasibility of magnetic recording using light of other photon energies and light polarizations.

In this paper, we demonstrate that contrary to previous reports, all-optical magnetic recording under both single-pulse and multiple-pulse sequences can be achieved, in narrow spectral ranges not with one but with two pairs of light polarization, either along [1 10] and [1−10] or [100] and [010] crystallographic axes of the garnet. Depending on the pump wavelength, different pairs of polarizations must be used to record and erase a magnetic domain. Such a possibility has not been either predicted or anticipated before. Note that the knowledge of the mechanisms, i.e., electronic transitions, responsible for the effect is the main obstacle preventing any quantitative modeling of light–matter interactions. Here, we reveal these transitions and thus open up the way for theoretical and computational studies of all-optical magnetic recording and ultrafast photo-magnetic phase transitions in dielectrics.

Results

Model of ultrafast photo-magnetic switching. Aiming to define the workspace of the key parameters allowing the switching, we developed a phenomenological model of the photo-induced magnetization dynamics in YIG:Co film based on the Landau–Lifshitz–Gilbert (LLG) equation:

$$\mathbf{M} = \gamma \left( \mathbf{M} \cdot \frac{\partial \Phi}{\partial \mathbf{M}} + \alpha \left( \mathbf{M} \cdot \frac{\partial \Phi}{\partial \mathbf{M}} \right) \right) + T_{\text{eff}}(\mathbf{M}, E),$$  \hspace{1cm} (1)

where $\gamma$ is the gyromagnetic ratio, $\alpha$ is the Gilbert damping parameter, $M_s$ is the saturation magnetization, $E$ is the electric field vector, and $\Phi$ is the thermodynamic potential of the unperturbed magnet in the form of free energy.

The last term in the equation stands for optically induced spin-torque $T_{\text{eff}}(\mathbf{M}, E)$. The latter may be represented in the following way in the Cartesian coordinate system:

$$T_{\text{eff}}(\mathbf{M}, E) = -\gamma |\mathbf{M}| \mathbf{H}_{\text{eff}}(\mathbf{M}), \quad \mathbf{H}_{\text{eff}}(\mathbf{M}) = -\beta_{ijkl} M_i E_j E_k.$$  \hspace{1cm} (2)

where $\beta_{ijkl}$ is the photo-magnetic tensor. The number of independent tensor components can be found taking into account the 4mm point group for (001) oriented YIG:Co film and the fact that the tensor $\beta_{ijkl}$ must be invariant with respect to permutations of the last two indices. The material parameters and coefficients of the photo-magnetic tensor $a, b, c$ (and methods) were obtained by fitting the theoretical results to the experimental data from ref.

There are four easy magnetization axes near <111>− type directions and 8 ground states for magnetization vector, taking into account two possible orientations along each axis. We denote these metastable magnetization states with numbers 1−8 (see Fig. 1a). This model allows to simulate the trajectories of the switching (see Fig. 1b). Earlier studies allowed to estimate that the life-time of the photo-magnetic anisotropy is about 20 ps. The model accounts for this by introducing a temporal relaxation of the tensor components $\beta_{ijkl}$. Therefore, during the first 20 ps after the optical excitation the motion of spins will mainly be defined by the field of the photo-induced magnetic anisotropy and afterwards the motion will be driven by the effective field of the steady-state magnetocrystalline anisotropy.

We demonstrate the modeling of the photo-magnetic switching between possible magnetization axes (phases), e.g., 1−4, 1−5 and 1−8 (see Fig. 1b). Taking 1 as the initial magnetization state we performed the simulations of the trajectories of the magnetization vector varying the intensity of light and magnetic damping $a$ (see Methods). Figure 1c, d summarizes the phase diagram of the simulations showing the final states to which the magnetization relaxes after being excited by light of two polarizations ($E || [100]$ and $E || [110]$). The switching can be observed for these polarizations in a wide range of light intensities. This model demonstrates different trajectories of the switching (see Fig. 1e) between: (i) only in-plane magnetization components (between [110] and [1−10] axes, see 1−4 trajectory); (ii) only perpendicular magnetization components (between [001] and [00−1] axes, see 1−5 trajectory); and (iii) simultaneously in-plane and perpendicular magnetization components (between [111] and [1−1−1] axes, see 1−8 trajectory). This model allows to simulate the behavior of a single spin in a crystal lattice with the symmetry of the iron garnet, where the Gilbert damping is taken as a constant. The damping was treated as a fitting parameter. The best agreement of our simulations with the experimental data reported here was achieved for $a = 0.28$. Note that the previously mentioned experimental value of $a = 0.210,13$ was measured using the technique of ferromagnetic resonance, where the amplitudes of magnetization oscillations are small. It is well known, however, that the damping increases with the amplitude of the magnetization oscillations.

Hence the simulations suggest that in addition to the photo-magnetic recording reported in ref. 10, the symmetry of the garnet
femtosecond laser pulses, across the sample surface scanning a polarized laser beam, consisting of series of most conventional way to illustrate the switching is based on is shown in Fig. 2. The domain structure in YIG:Co thin pattern as observed in a polarized light microscope (see Methods) has been studied and reported in detail in refs. 22 of additional peaks associated with Co-ions 15. The most electronic transitions with other selection rules. mechanism of the switching must rely on excitation of other light polarizations than \([100]\), for which the phase diagrams to help illustrate the switching. Trajectories of the switching from state 1 to 8 was performed for \(\phi = 0^\circ\). The trajectories calculations are performed for various damping and fluence, based on the phase diagrams to help illustrate the switching. Trajectories of the switching from state 1 to 8 was performed for \(\alpha = 0.28\). The switching paths from 1 to 4 and from 1 to 5 were simulated for \(\alpha = 0.12\). Fit parameters for all simulations are \(b = -\alpha/3\), \(c = \alpha/2\) (see Eq. 4 in Methods).

Films supports the possibility of the magnetic switching using other light polarizations than \([100]\) and \([010]\), for which the mechanism of the switching must rely on excitation of other electronic transitions with other selection rules. Compared to pure YIG, the absorption spectra of YIG:Co in the range of photon energies from 0.5 eV to 2 eV show a number of additional peaks associated with Co-ions 15. The most pronounced peaks are observed around 1.13 eV (the wavelength is 1.1 μm) and 0.95 eV (the wavelength is 1.3 μm) 16-18. So far the most conventional way to illustrate the switching is based on scanning a polarized laser beam, consisting of series of femtosecond laser pulses, across the sample surface 6-8. To verify the feasibility of the magnetic switching at other photon energies and other polarizations of light, we employed this method for two polarizations of light \([110]\) and \([1-10]\). The experiments were performed at several photon energies in the range from 0.83 eV to 1.24 eV.

**Multiple-pulse magnetic switching.** The initial magnetic domain pattern as observed in a polarized light microscope (see Methods) is shown in Fig. 2. The domain structure in YIG:Co thin films has been studied and reported in detail in refs. 22-24 together with a description of the experimental procedure of defining the magnetization in the domains (see Methods). We rotated the linear polarization over 45° with respect to \([100]\) axis (\(\phi\) is the angle between the electric field of light and \([100]\) axis) and searched for the wavelength allowing the switching. After scanning the beam with \(\phi = 45^\circ\) and photon energy 1.1 eV (\(\lambda = 1140 \text{ nm}\)) the large black domains are turned into white ones. The orthogonal polarization along \([1-10]\) crystallographic axis does the opposite. Both polarizations are different from those used for the switching in ref. 10 indicating that here it most likely proceeds along different microscopic mechanism, i.e., relies on excitation of different electronic transitions than those reported earlier.

Figure 3 shows how the switched area increases with an increase of number of pump pulses \(N\) using quasi-static magneto-optical imaging in Faraday geometry within 10 ms frame after excitation with a single pump pulse (see Methods) (see Fig. 3a). By increasing the pump pulse fluence, the number of pulses required for the switching can be brought down to one. Note that the appearance of a domain after a certain number of pulses is a very reproducible effect that is determined by a balance between the repetition rate of the pulses and magnetostatic field driven domain wall motion (for more details see Methods). The process of domain formation using multiple pulses for 105 mJ cm\(^{-2}\) is following: after first pump pulse the magnetization was switched, however the switched area was not stable enough. The next pulses resulted in an expansion and thus stabilization of the recorded
1000 pulses per second. The beam contains 50 fs laser pulses with the stable domain. The time period between the pulses is 1 ms (repetition rate domain) to the area of pump laser spot was calculated as the ratio of the recorded domain area (the black large and the pump polarization is along \([110]\). The normalized switched area before and after the laser excitation lines are guides to the eye. The inset shows the magnetic domain patterns as a matter of fact, the switched domains have been clearly and reproducibly seen after \(N = 5\) (see Figure 3).

**Selectivity of photo-magnetic recording in Co-doped YIG.** Figure 4 shows a summary of the switching efficiencies as functions of light wavelength, polarization angle, and pulse fluence. The spectral dependence shown in Fig. 4a reveals two resonant features around 1.1 eV and 0.95 eV, respectively. Figure 4b shows that the most efficient photo-magnetic switching with photon energy of 0.95 eV is observed for \([100]\) polarization. If the pumping is at photon energy of 1.1 eV, the switching has the maximum efficiency for \([110]\) polarization. This exactly corresponds to the modelling results revealing different threshold intensities for the switching with different light polarizations (see phase diagrams in Fig. 1). Finally, pumping at 0.95 eV is characterized by three times lower threshold intensity compared to the case of pumping at 1.1 eV (Fig. 4c).

The revealed selection rules shown in Fig. 4a, b allow us to identify the microscopic mechanisms responsible for the photo-magnetic recording. According to refs. 17,18 a sharp peak around 1.1 eV is due to \(\text{^4T}_1 \rightarrow \text{^4T}_2\) electronic transition in Co\(^{2+}\) ions in octahedral sites. A broader peak at 0.95 eV corresponds to \(\text{^5E} \rightarrow \text{^5T}_2\) transitions in Co\(^{3+}\) ions and \(\text{^4A}_2 \rightarrow \text{^4T}_1\) transitions in Co\(^{2+}\) ions in tetrahedral sites\(^{17}\). In both experiment and simulations the threshold fluence for the switching with light polarization along \([110]\) axis is about three times larger than the threshold for light polarization along \([100]\) axis (see Figs. 1c, d and 4c). The reason for this is possibly related to the fact that, as a general crystallographic property, the oscillator strength of low-symmetry tetrahedral sites is much stronger than high-symmetry octahedral sites\(^{16,19,20}\). As a result, the response of octahedral ions becomes clearly noticeable only when sufficient pump fluence is applied. The contribution to the magnetic anisotropy at room temperature is also stronger from Co-ions in the tetrahedral sites\(^{21}\).

**Discussion**

Finally, it is interesting to note the remarkable efficiency and selectivity of the switching by pumping different electronic transitions in Co-dopants. Apart from purely fundamental interest and impact on further development of theory of photomagnetism, the discovered multi-wavelength magnetic recording in iron garnet may spur applications of photo-magnetism in information processing technology. Firstly, the possibility of recording at several wavelength shows that the wavelength for the most efficient photo-magnetic recording is not fixed but can be tuned in the range relevant to optical telecommunication systems. Secondly, our work also shows that information transferred from light to spins can be encoded not only in polarization state, but also in the wavelength and the intensity of the recording beams. In fact, our work shows that due to the additional degree of freedom one can increase the intensity of data traffic from light to spins. For instance, different spectral components of light can carry information, which due to dispersion will be focused and recorded on different domains of a storage medium. Thirdly, the discovered selection rules allowed us to reveal the origin of the photo-magnetic centers and thus to perform quantitative estimates of the efficiency of photo-magnetic recording. In the studied films only one of forty Fe ions is substituted by Co\(^{2+}\)-Co\(^{3+}\). It means that single-photon excitation of Co ion must be, in principle, sufficient to control the spins in a rather large volume of 28 nm\(^3\) (see Methods). Fourthly, the selection rules reported in this paper reveal the possibility of an independent control of spins coupled to octahedral and tetrahedral photo-magnetic centers. Employing the discovered degrees of freedom, polarization pulse
shaping and the principles of coherent control, one can anticipate new opportunities for ultrafast optical manipulation of coherent spin waves far from the center of the Brillouin zone.

This work shows that tuning the wavelength, magnetic damping, polarization and fluence of light it is possible to find different regimes of the switching in a wide class of materials, which can possibly be even more efficient than those demonstrated before. The examples of suitable materials are ferrites (garnet, spinel, ortho- and hexaferrites, ferric borates, and magnetite)\(^2\), perovskites, spin glasses. For instance, a strong contribution to single ion magnetic anisotropy is typical for \(3d\) (e.g., Mn\(^2+\), Mn\(^3+\), Cr\(^3+\), Cr\(^2+\), Fe\(^{2+}\), Co\(^{2+}\), Co\(^{3+}\), Ni\(^2+\), Ni\(^3+\)), \(4d\) (Ru\(^{3+}\), Ru\(^{4+}\)), \(5d\) (Ir\(^{3+}\), Ir\(^{4+}\)), and \(4f\) (Ce\(^{3+}\), Tb\(^{3+}\)) elements. Finding novel families of magnetic dielectrics switchable with laser pulses is crucial for future applications of photo-magnetism.

**Methods**

**Materials.** The monocristalline cobalt-doped yttrium iron garnet (YIG:Co) thin film with composition \(\text{Y}_3\text{Fe}_5\text{O}_{12}\cdot\text{Co}\cdot(\text{GeO}_2)\) was deposited on gadolinium–gallium garnet (GGG) substrate\(^3\). The miscut angle of the GGG substrate was about \(4^\circ\). The dopant Co\(^{2+}\) and Co\(^{3+}\) ions substitute Fe\(^{3+}\) in both tetrahedral and octahedral sublattices, which are coupled antiferromagnetically. The Co-ions enhance the magneto-crystalline anisotropy and result in a strong photo-magnetic effect, allowing to manipulate the magnetization with light\(^2\). In this sample the magnetic anisotropy constant \(K_{\text{eff}} = 8.4 \times 10^6 \text{erg cm}^{-3}\) and uniaxial \(K_{\text{u}} = -2.5 \times 10^6 \text{erg cm}^{-3}\) were measured at room temperature. Incorporation of Co-ions also increases the Gilbert damping parameter to a large value of \(a = 0.2\), which was measured using ferromagnetic resonance technique. Both tetrahedral and octahedral Fe\(^{3+}\) ions are substituted by Ge\(^{4+}\) ions to decrease the saturation magnetization \(M_S = 7 \mu_B\). Co\(^{2+}\) dopants enter the dodecahedral sites and take care of charge compensation. The Curie temperature was \(455\) K.

As single ion contribution to magnetic anisotropy from Co-ions is orders of magnitude higher than that of Fe ions\(^1\), a single photon can control spins in a large garnet volume. For YIG:Co film with the thickness of \(7.5 \mu\text{m}\), the minimum pump intensity required for the switching was \(34 \text{ mJ cm}^{-2}\). YIG:Co film is a ferrimagnetic dielectric transparent in the near infrared spectral range\(^4\). For photon energy at \(0.95 \text{ eV}\) the total absorption in the film is about \(12\%\). It means that the density of absorbed photons is \(3.6 \times 10^{19} \text{ cm}^{-2}\) and a single-photon switches the magnetization in \(28 \text{ nm}^2\) of the garnet.

**Experimental technique for photo-magnetic imaging.** To investigate the magnetization switching induced by femtosecond linearly polarized laser pulses in YIG:Co thin film, we employed the technique of magneto-optical polarized microscopy in Faraday geometry. The magnetic contrast in polarizing microscope comes from the fact that magnetic domains with different out-of-plane magnetization component will result in different angles of the Faraday rotation. The light emitted from the LED source, passing through the linear polarizer \(\phi\) and quarter-wave plate \(\phi = 45^\circ\) will acquire different polarization rotation in different magnetic domains. The differences can be visualized with the help of an analyzer and a CCD camera. The obtained images visualize magnetic domains. The garnet film was excited with one or multiple (N) pulses. The duration of each pulse was about 50 fs and pulse-to-pulse separation \(t_\text{sep}\) was varied. The pump beam with the fluence below \(250 \text{ mJ cm}^{-2}\) was focused to a spot about \(130 \mu\text{m}\) in diameter. Further increasing the fluence or the number of pulses does not increase domain size considerably, but instead leads to heating of the sample. The wavelength of the pump pulses was tuned within the range between \(1000 \text{ nm}\) and \(1300 \text{ nm}\). The images of magnetic domains were taken before and after the pump excitation. Taking into account the differences between the images we deduced photo-induced effects on the magnetization in the garnet. In YIG:Co thin film with small miscut angle four types of magnetic domains were observed. Any of the four domains with magnetization states \(1, 4, 5,\) and \(8\) (orientations are shown in Fig. 1b) could be stabilized by applying and removing an in-plane field of about \(\mu_0H = 2 \text{ mT}\) directed along one of the four easy-axes of the garnet \(\text{<110>}\). The magneto-optical contrasts for 5 (or 4) and \(8\) domains are substituted by Ge\(^{4+}\) ions to decrease the saturation magnetization \(M_S = 7 \mu_B\). Co\(^{2+}\) dopants enter the dodecahedral sites and take care of charge compensation. The Curie temperature was \(455\) K.

**Figure 4.** Selection rules of photo-magnetic switching on wavelength and polarization. For measurements 5 laser pulse pumps (\(N = 5\)) were used. a Normalized switched area as a function of the photon energy for two polarizations \(E \parallel \text{[110]}(\phi = 45^\circ)\) and \(E \parallel \text{[100]}(\phi = 0^\circ)\). Solid lines were fitted by Gaussian function. b Normalized switched area as a function of the angle \(\phi\) for the incoming polarization for two photon energies 0.935 eV \((\lambda = 1300 \text{ nm})\) and 1.1 eV \((\lambda = 1140 \text{ nm})\). Solid lines are guides to the eye. c Fluence dependence of the normalized switched area for two photon energies 0.95 eV and 1.1 eV. Dashed lines are guides to the eye. The measurements at different photon energies reveal different threshold fluences \(I_{\text{th}}\).
Fig. 5 Photo-magnetic recording in YIG:Co with delay between pump pulses. The study was performed by means of analysis of normalized switched area with N = 5 laser pump pulses as a function of the pump pulse-to-pulse separation time \( t_p \). The red solid line is a guide to the eye. The blue solid line was fitted using exponential function with the characteristic time \( t_0 = 7 \) ms.

Data availability

The data that support the findings of this study are available from the corresponding authors on reasonable request.

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Theoretical model of photo-magnetic switching. The expression for the photo-magnetic tensor in Voigt notation is:

\[
\begin{align*}
\beta_{ij} &= \begin{pmatrix}
0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0
\end{pmatrix},
\end{align*}
\]

(3)

We take the action of the laser pulse as a perturbation to the initial state of the magnetization in the film, obtained from the LLG-equation Eq. (1). Depending on the axis of linear polarization of light it gives:

\[
\begin{align*}
\theta(0) &= \theta_0 + A \frac{\pi}{180} (a \sin(2\phi_0) \sin(\theta_0) + \frac{1}{2} b \sin(2\theta_0)) \\
\phi(0) &= \phi_0 + A \frac{\pi}{180} (b \cos(\theta_0) \pm a \sin(2\phi_0)) \\
\theta(0) &= \theta_0 - \frac{\Delta}{2} \pi (b \cos(\theta_0) \pm a \sin(2\phi_0)) \\
\phi(0) &= \phi_0 + A \frac{\pi}{180} (b \cos(\theta_0) \pm a \sin(2\phi_0)) \cos(\theta_0)
\end{align*}
\]

We use the following values of the magnetic fields and magnetic moments of the g- and f-electrons.

\[
\begin{align*}
\theta(0) &= \theta_0 + A \frac{\pi}{180} (a \sin(2\phi_0) \sin(\theta_0) + \frac{1}{2} b \sin(2\theta_0)) \\
\phi(0) &= \phi_0 + A \frac{\pi}{180} (b \cos(\theta_0) \pm a \sin(2\phi_0)) \\
\theta(0) &= \theta_0 - \frac{\Delta}{2} \pi (b \cos(\theta_0) \pm a \sin(2\phi_0)) \\
\phi(0) &= \phi_0 + A \frac{\pi}{180} (b \cos(\theta_0) \pm a \sin(2\phi_0)) \cos(\theta_0)
\end{align*}
\]

where \( \phi_0 \) and \( \theta_0 \) denote the ground state angles of the initial domain with respect to [100] and [001] axes, correspondingly. \( a = a(\lambda) \), \( b = b(\lambda) \), and \( c = c(\lambda) \) are the parameters, which can be expressed through photo-magnetic tensor coefficients and \( A = \mu n(E^2) \Delta t \), where \( E^2 \) is the average intensity of the field in the pulse and \( \Delta t = 20 \) ps is the characteristic time of the emergence of the magnetic domain taken from the experiment. After time interval of the order of \( \Delta t \) due to relaxation of the photo-excited Co-electrons, the magnetization dynamics can be regarded as fast (see Fig. 1). As it is seen from Eq. (4), the initial perturbation strongly depends on the Gilbert damping, initial state and the intensity of light; these are parameters which are crucial for the switching. Wavelength dependence of the efficiency of switching implies wavelength dependence of the photo-magnetic tensor coefficients, namely \( a(\lambda) \), \( b(\lambda) \), and \( c(\lambda) \).
Author contributions
A.S. conceived the project with contributions from A.K. and A.V.K. The measurements were performed by K.S. and A.S. The model was designed and the simulations were performed by M.D.D, K.A.Z, and A.K.Z. A.S., A.K., and A.V.K. co-wrote the manuscript with contributions from K.S. and A.K.Z. The project was coordinated by A.S.

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