Three-dimensional Random Dielectric Colloid Metamaterial with Giant Isotropic Optical Activity

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Motivated by the theoretical observation that isotropic chirality can exist even in completely random systems, a dielectric metamaterial consisting of a random colloid of meta-atoms is designed, which exhibits unprecedentedly high isotropic optical activity. Each meta-atom is composed of a helically arranged cluster of silicon nanospheres. Such clusters can be fabricated by large-scale DNA self-assembly techniques. It is demonstrated that the use of a high concentration of the meta-atoms in the colloid provides significant suppressions of incoherent scattering losses. As a result, the proposed system shows three orders of magnitude improvement of isotropic optical activity as compared with the previous metamaterial designs. This work highlights the significant potential of completely random systems, which are commonly produced in colloidal sciences, for applications as metamaterials toward novel photonic effects and devices.

1. Introduction

The development of metamaterials opens numerous new opportunities for manipulating electromagnetic waves. Most metamaterial structures consist of regular array of meta-atoms.\textsuperscript{[1]} As an alternative, one may wonder whether or how one can achieve non-trivial metamaterial functionalities in complete random or disordered systems. This is important from a fundamental perspective in the context of recent efforts in exploring the effects of disorders in photonic systems.\textsuperscript{[2–7]} It is also of significance from a practical point of view, since many large-scale fabrication techniques, such as self-assembly techniques, readily produce random array of meta-atoms.\textsuperscript{[8–13]}

For a linear metamaterial consisting of a completely random array of meta-atoms, after homogenization, its electromagnetic properties can be described by an effective medium with the following constitutive relations: [14, p. 28]

\[
\begin{align*}
D &= \varepsilon \cdot E + \xi \cdot H, \\
B &= \mu \cdot H + \zeta \cdot E
\end{align*}
\]

Here, \(D\), \(B\), \(E\), and \(H\) are the electric and magnetic displacement vectors and fields, \(\varepsilon\) and \(\mu\) are the permittivity and permeability tensors, respectively, and \(\xi\) and \(\zeta\) are the bianisotropic tensors. If one further assumes that the effective medium is lossless, then \(\xi = \xi^\dagger\). [14, § 2.7] Since the array is completely random, the effective medium must be rotationally invariant. Consequently, \(\varepsilon\), \(\mu\), and \(\xi\) must all be scalar. In the lossless case, the imaginary part of \(\xi\) describes effect of isotropic chirality (natural optical activity), which is a reciprocal effect, while the real part of \(\xi\) describes the effect of isotropic magneto-electric coupling (alternatively called Tellegen effect), which is a nonreciprocal effect. Therefore, one should be able to observe isotropic chirality or isotropic magneto-electric coupling even in a system that is completely random.

Motivated by the theoretical considerations above, in this letter, we introduce a dielectric metamaterial consisting of a random colloid of specifically designed silicon nanoclusters in order to achieve giant isotropic chirality. We demonstrate that due to the absence of absorption, the concentration of the nanoclusters in the colloid can be very high, which, in turn, provides a significant suppression of incoherent scatterings. Our metamaterial colloid exhibits isotropic chirality that is three orders of magnitude higher than those of previously reported isotropic chiral materials. Moreover, we demonstrate that similar superiority is achieved in terms of the figure of merit given by the ratio of the chirality parameter and the average extinction coefficient (such figure of merit is introduced in analogy to the magneto-optical figure of merit [15, § 9.6.5]). Since we consider in this contribution only isotropic chirality effect and neglect the nonreciprocal magneto-electric effect, in what follows, we use the conventional electromagnetic notation according to which scalar parameters \(\xi = ik/c\) and \(\zeta = -ik/c\) are expressed in terms of the so-called chirality parameter \(\kappa\) [14, p. 28] \((c\) is the speed of light). Time harmonic oscillations in the form \(e^{i\omega t}\) are assumed throughout the letter.

Chirality phenomenon, first observed more than two centuries ago,\textsuperscript{[16]} is of current importance for a diversity of applications such as in organic and inorganic chemistry, pharmaceutical engineering, optical communications, displays, spectroscopy,
and sensors.\textsuperscript{[17–20]} The chiral effects in naturally occurring materials are relatively weak. Therefore, there is significant recent interest in designing metamaterials\textsuperscript{[1,21–28]} and colloids\textsuperscript{[11–13,26,29–31]} with enhanced chiral properties. In particular, there is an extensive literature on the chiral properties of metasurfaces.\textsuperscript{[12–35]} The chiral response of a three-dimensional (3D) metamaterial, however, can be qualitatively different. Isotropic chiral response, where the same chiral effects occur independently of the directions of light propagation, is a 3D effect that is difficult to achieve with metasurfaces. Such an isotropic chiral response is potentially important for topological photonics\textsuperscript{[36,37]} and stereochemistry.\textsuperscript{[19,38]} Moreover, isotropic chiral materials provide an attractive route toward achieving the phenomenon of negative refraction in bulk.\textsuperscript{[25–39]} In the optical frequency range, the existing designs of 3D chiral metamaterials rely upon metallic structures with complex geometries.\textsuperscript{[1,26–28]} The use of metal introduces significant inherent loss, whereas the use of complex geometries represents major challenges for 3D nanofabrication. Our proposed chiral metamaterial colloid overcomes all the above-mentioned drawbacks.

2. Results

2.1. Meta-Atom Design

In an isotropic chiral material, the right (RCP) and left (LCP) circularly polarized light are eigenstates for all propagation directions. The chirality parameter of a material is defined as $\kappa = (n_+ - n_-)/2$, where $n_+$ and $n_-$ are the refractive indices for right and left circularly polarized light, respectively.\textsuperscript{[20]} The isotropic chirality results in the optical rotation $\Delta \theta$ and the circular dichroism $\eta$ effects for waves propagating over distance $L$ in the material, which is expressed as $\Delta \theta + \eta \omega L/c$. \textsuperscript{[17, § 5.2]} For a metamaterial, the chirality parameter is nonzero only when the constituent meta-atoms have no mirror symmetry. For low-concentration colloids, the macroscopic chirality parameter and the properties of the individual meta-atoms are related by \textsuperscript{[14, § 2.10.1]}

$$\kappa = \frac{N}{3} \text{tr}(\overline{\alpha}_{\text{em}})$$

(2)

where $N$ is the volume concentration of the constituent meta-atoms in the colloid and $\overline{\alpha}_{\text{em}}$ is the magnetoelectric polarizability tensor of each meta-atom. This relation is equivalent to the Rosenfeld equation in chemistry literature. \textsuperscript{[17, Eq. 5.2.2]} For high constituent concentrations, the chirality parameter is obtained using the Maxwell Garnett mixing rule. \textsuperscript{[14, § 7.2.2]} Relation (2) is rotationally invariant since the trace of a matrix does not change under rotations.

As is seen from (2), when choosing the constituent meta-atom for our colloid, we need to maximize the trace of its magnetoelectric polarizability tensor $\overline{\alpha}_{\text{em}}$. Next, we suggest a simple extraction technique of this trace for an arbitrary meta-atom. The technique is based on a previously proposed method based on the far-field scattering.\textsuperscript{[40]} Consider an arbitrary meta-atom in free space.

We illuminate it alternately with RCP and LCP monochromatic light along the $-x$, $-y$, and $-z$ directions, as shown with gray arrows in Figure 1. The upper and bottom signs in the electric field amplitudes $|E_{\pm}| = \sqrt{2}E_0$ correspond to the right and left circular polarizations, respectively. Next, we can arrive at the following expression for the trace of the polarizability tensor $\overline{\alpha}_{\text{em}}$ (see Supporting Information [41, § 1])

$$\text{tr}(\overline{\alpha}_{\text{em}}) = \frac{\pi n c^2}{\omega^2 E_0^2} \sum_{m=n,+} \left( E_{m}^{(n)} - E_{m}^{(n)} \right)$$

(3)

where $E_{+}$ and $E_{-}$ are the RCP forward-scattered field from the meta-atom when illuminated by RCP light and the LCP forward-scattered field from the meta-atom when illuminated by LCP light, respectively. Subscript $m$ denotes the direction of the incident light. The scattered light can be measured using full-wave simulations at a distance $r$ from the meta-atom. Expression (3) allows simple calculation of the trace of the magnetoelectric tensor of an unknown meta-atom from six scattering calculations. As one can see, the trace as well as optical activity (according to relation (2)) are maximized when the meta-atom exhibits the highest contrast in forward scattering of incident RCP and LCP light for the three orthogonal directions. It is worth mentioning that the derived expression (3) to some extent resembles the conventional dimensionless circular intensity difference parameter defined as $\Delta = (I_+ - I_-)/(I_+ + I_-)$. \textsuperscript{[17, p. 162]} where $I_+$ and $I_-$ are the scattering intensities of the meta-atom excited by the incident RCP and LCP light, respectively.

Next, we design the constituent meta-atom in the form of a cluster of dielectric nanospheres. In order to maximize the interaction between the meta-atoms and incident light, we need to use high-permittivity dielectric. For the chosen operational frequency of 400 THz ($\lambda_0 = 750$ nm wavelength in vacuum or $\lambda = 564$ nm...
in water where the meta-atoms are dispersed), a good candidate is crystalline silicon.\(^{(42,43)}\) Fabrication of both crystalline and amorphous silicon nanosphere colloids have been demonstrated in the literature.\(^{(44–46)}\) The nanospheres in the colloid can be coated by DNA origami belts and hybridized to form a cluster with the desired chiral shape, as was demonstrated recently in.\(^{(47)}\) Some alternative fabrication techniques for such a colloid metamaterial can be found in refs. \((48–54)\). We choose the maximum size of the cluster meta-atom \(D_{\text{max}} = 130\) nm so that it remains subwavelength in water at wavelength \(\lambda = 564\) nm. Such choice, although not unique, was made in order to avoid generation of higher multipoles in the meta-atoms and diffraction effects, as well as to have the possibility of strong suppression of light scattering in the colloid metamaterial, as discussed below. With such a choice of sizes, the designed meta-atom is nonresonant: The first Mie resonance for a quasispherical nanoparticle at the chosen operating frequency occurs when its size is \(D_{\text{Mie}} \approx 202\) nm,\(^{(51)}\) which is twice larger than our chosen size \(D_{\text{max}}\).

Using the expression (3) and calculating the six scattered far-fields \(\mathbf{E}_s\) and \(\mathbf{E}_r\) via ANSYS HFSS (a finite element method solver), we have found that large value of the trace of \(\mathbf{\alpha}_{\text{em}}\) tensor (for the given maximum size of the occupied volume \(\pi D^3_{\text{max}}/6\)) is achieved for a helically shaped cluster of nanospheres (helical shape was chosen as it is considered optimal for metallic chiral shapes, as was demonstrated recently in.\(^{(47)}\) Some alternative fabrication techniques for such a colloid metamaterial can be found in refs. \((48–54)\). We choose the maximum size of the cluster meta-atom \(D_{\text{max}} = 130\) nm so that it remains subwavelength in water at wavelength \(\lambda = 564\) nm. Such choice, although not unique, was made in order to avoid generation of higher multipoles in the meta-atoms and diffraction effects, as well as to have the possibility of strong suppression of light scattering in the colloid metamaterial, as discussed below. With such a choice of sizes, the designed meta-atom is nonresonant: The first Mie resonance for a quasispherical nanoparticle at the chosen operating frequency occurs when its size is \(D_{\text{Mie}} \approx 202\) nm,\(^{(51)}\) which is twice larger than our chosen size \(D_{\text{max}}\).

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Table 1 summarizes the traces of the dipolar electric, magnetic, and magnetoelectric polarizability tensors extracted using (3) and technique reported in ref. \(^{(57)}\).

As is seen, the largest isotropic response arises from the electric dipole excitation, exceeding by one or several orders of magnitude those of the magnetic dipole excitation and magnetoelectric coupling. Such unbalanced polarization response indicates that the designed cluster is still far from the theoretical maximum chirality bound which occurs when \(\alpha_{\text{em}} = \alpha_{\text{em}} = \alpha_{\text{mm}}\).\(^{(56,58,59)}\) Nevertheless, this theoretical bound can be achieved only with strong field localization, like in plasmonic meta-atoms. However, the lossy nature of plasmonic clusters is inconsistent with the goal of the present work. The imaginary part of the trace of the polarizability takes into account both the scattering of light away from the direction of incidence, which becomes the incoherent scattering loss for the random metamaterial, as well as material absorption loss since the operating frequency is above the band gap of crystalline silicon. For the proposed meta-atom, the scattering loss dominates over the absorption loss by two orders of magnitude. Additionally, we have analyzed a pyramidal cluster of four touching nanospheres (a cluster of three arbitrary nanospheres always has the inversion symmetry). The nanospheres were chosen with the same radii of 30 nm, while their permittivities were all different to break inversion symmetry: \(\epsilon_1 = 2, \epsilon_2 = 6, \epsilon_3 = 10,\) and \(\epsilon_4 = 13.8\). For this cluster, the absolute values of the polarizability traces are \(\text{tr}(\mathbf{\alpha}_{\text{em}})/3 = 8.2 \times 10^{-22} \text{m}^3\), \(\text{tr}(\mathbf{\alpha}_{\text{mm}})/3 = 2.0 \times 10^{-25} \text{m}^3\), and \(\text{tr}(\mathbf{\alpha}_{\text{mm}})/3 = 5.2 \times 10^{-23} \text{m}^3\). Comparing the obtained value of the magneto-electric polarizability trace to that in Table 1, one can see that the pyramidal cluster exhibits four times weaker chiral properties than the helical cluster, which is in agreement with the results in ref. \(^{(11)}\).

### 2.2. Scattering Inside a Random Metamaterial and Supercell Approximation

Next, we study optical response of the random distribution of the designed dielectric constituent meta-atoms, as shown in Figure 1, dissolved in water. From (2), since the strength of the isotropic chirality parameter increases as the volume concentration \(N\) increases, we are interested in the regime with the high volume concentration. As a dimensionless measure of the volume concentration, we define \(N_\nu = N V_{\text{circ}}\), where \(V_{\text{circ}} = 1.15 \times 10^{-21} \text{m}^3\) is the volume of the sphere circumscribed around the meta-atom shown in Figure 1. For given volume concentration, we analytically calculate the relative permittivity \(\mathbf{\tilde{\varepsilon}}\), permeability \(\mathbf{\tilde{\mu}}\), and chirality parameter \(\mathbf{\tilde{\eta}}\) of the metamaterial using the Maxwell Garnett mixing rule.\(^{(14, \text{§ 7.2.2})}\) Whose applicability is not limited to low concentrations. Effective refractive indices for RCP and LCP light propagating in the colloid metamaterial are expressed as \(n_{\nu} = \sqrt{\mathbf{\tilde{\varepsilon}}\mathbf{\tilde{\mu}}} \approx x\).\(^{(60, \text{§ 2.2.1})}\) In order to calculate the figure of merit, we consider the attenuation coefficient \(\beta = 2k_n\text{Im}(n)\) of the colloid metamaterial which shows how much light is attenuated over propagation distance \(L\), that is, \(I(L) = I(0)e^{-\beta L}\). As mentioned above, the primary origin of the attenuation is from incoherent light scattering, which arises from the irregular distribution of the constituent meta-atoms in the metamaterial (fluctuations of density of scatterers). The incoherent scattering occurs even in the situations when the average distance between the meta-atoms is much smaller than the wavelength and the medium can be considered as uniform.\(^{(61)}\)

According to the classical independent-scattering formulation of the radiative transfer theory, the total scattering in a colloid is proportional to the volume concentration of its constituent meta-atom.\(^{(62–66)}\) Under this assumption, the scatterings from all the meta-atoms are uncorrelated and \(\beta_{\text{uncorr}}(N) = 2k_n\text{Im}(n) = N\sigma\), where \(\sigma\) is the extinction cross section of individual meta-atom. However, as it was demonstrated in the early works of Twerisky et al.\(^{(62,67,68)}\) this assumption is accurate only for media with low volume fraction \(N_\nu\) (volume fraction is defined as a product of the volume of a single meta-atom and the volume concentration)
of the colloid, i.e., \( N_y = V_y N \). In dense colloids, there exist correlations in positions of the meta-atoms which lead to the modified attenuation coefficient

\[
\beta_{\text{corr}}(N, N_y) = N \sigma W(N_y) = 2k_y \text{Im}(n_{\text{corr}})
\]  

(4)

where \( W(N_y) = (1 - N_y)/(1 + 2N_y) \) is the so-called packing factor for spherical-shape constituent meta-atoms \([62]\) and \( \text{Im}(n_{\text{corr}}) = \text{Im}(n) W(N_y) \) is the effective refractive index of the colloid metamaterial whose imaginary part is scaled by the packing factor. The factor \( W(N_y) \) monotonically decreases with increasing volume fraction, leading to the fact that attenuation due to incoherent scattering in dense colloid metamaterials can be several orders of magnitude smaller than that predicted by the independent-scattering theory. Such scattering reduction can be intuitively understood by increased regularity of the colloid due to the loss of available space for its constituent meta-atoms. Equation (4), which was sometimes referred to as the quasicrystalline approximation (QCA), was subsequently confirmed for dense media experimentally \([63,66]\) and numerically \([65]\).

In our study, we exploit the effect of scattering loss reduction to design a chiral dielectric colloid metamaterial with high figure of merit, that is \( \text{FOM} = |\kappa|/\text{Im}(n_{\text{corr}}) \). This FOM is introduced in analogy to the magneto-optical FOM, where the latter is defined as the specific Faraday rotation divided by the attenuation constant. \([15, \S 9.6.5]\) For our system, the maximum volume fraction of the colloid is approximately \( N_y^{\text{max}} = 0.63 \), which is the measured value of \( N_y \) for densely packed amorphous solids of identical spheres \([69]\). This volume fraction corresponds to that of the mean of the cubic and hexagonal packings.

Full-wave simulations of a macroscopic random colloid metamaterial are not practical due to the large size of the simulation domain which must be large enough to accommodate the entire structure. Therefore, we implement an approach based on the supercell approximation \([70-72]\). The idea of the approach is depicted in Figure 2. The electromagnetic response of an infinite random metamaterial is approximated with that of an infinite periodic metamaterial whose supercell is enough large to include many randomly oriented meta-atoms. Consider first a periodic metamaterial with a supercell of dimensions \( L \times \Delta_x \times \Delta_y \) (along the \( x \), \( y \), and \( z \) axes, respectively) shown in Figure 2a. The different colors in the figure are employed only for visual clarity of periodicity. The meta-atoms are distributed within the supercell with completely random positions and orientations without touching one another. The supercell is periodically repeated in the \( y \) and \( z \) directions. Due to the periodicity, this metamaterial slab of thickness \( L \) will diffract some part of normally incident light into several diffraction orders. According to the supercell approximation, the amount of diffracted light in the periodic metamaterial provides an approximation of the incoherent scattering loss in a corresponding random metamaterial (of the same size and with the same meta-atom concentration) which is shown in Figure 2c. When the size of the supercell \( \Delta_y \) increases (see Figure 2b), the number of diffraction orders also increases, and the approximation becomes more accurate. For our simulations, we choose \( \Delta_y(N_y) > \lambda \) such that there are two diffraction harmonics in reflection and two diffraction harmonics in transmission. The periodicity along the \( y \) direction is chosen to be sub-wavelength to reduce the overall simulation domain. Thus, the scattering analysed in this way occurs only in the plane of incidence.

Next, according to the supercell approximation, we analyze the attenuation coefficient \( \beta \) in a random metamaterial by calculating diffraction scattering intensity for light propagating through the corresponding periodic metamaterial of the same concentration. In what follows, we choose the periodical configuration with four diffraction orders shown in Figure 2a. The supercells dimensions were chosen as \( \Delta_y = (V_{\text{corr}}/N_y)^{1/3} \), \( L = 3 \Delta_y \), and \( \Delta_z = p \Delta_y \), where \( p \) is the smallest integer for which the periodic metamaterial supports four diffraction orders. For each of the seven discrete values of \( N_y \), we performed 15 scattering simulations of the metamaterial supercell. In each of the 15 simulations, the supercell had the same dimensions but different random distributions and orientations of the chiral meta-atoms. The simulated total diffracted intensity \( I_{\text{diff}} \) normalized by the incident intensity
I\textsubscript{inc} is plotted in Figure 3 versus the volume fraction $N_V$. Furthermore, we additionally plot in the same Figure the incoherent scattering intensity loss calculated based on two analytical fitting models for a truly random metamaterial. The first model is based on the aforementioned QCA theory and implies $I\textsubscript{scat}/I\textsubscript{inc} = \frac{[I(0) - I(L)]}{I(0)} = 1 - e^{-\rho\textsubscript{inc}L} \approx \rho\textsubscript{scat}L$. According to the supercell approximation, the ratios $I\textsubscript{scat}/I\textsubscript{inc}$ and $I\textsubscript{diff}/I\textsubscript{inc}$ should be proportional to one another with some unknown proportionality coefficient $\psi$. Substituting $\rho\textsubscript{scat}$ from (4), we chose $\psi = 6.67 \times 10^{-23}$ m$^4$ which ensures the best fitting for the simulated curve in Figure 3. As is seen, the simulated diffraction loss in the periodic metamaterial follows the same dependence on the volume concentration $N_V$ as the theoretical orange fitting curve predicted by the QCA theory for the random metamaterial. The corresponding incoherent scattering loss for the random metamaterial predicted by the independent-scattering theory is plotted by the orange dashed line. Summarizing, one can see that incoherent scattering in the designed colloid metamaterial is in fact suppressed by the factor $1/\text{W}(N_v)$ in accordance with the QCA theory. Thus, for high concentrations of the meta-atoms in the colloid metamaterial, the scattering loss drastically decreases when the volume concentration increases.

2.3. Giant Isotropic Chirality

Next, we analyze the chirality parameter versus the volume fractions for random metamaterial shown in Figure 2a. First, the transmission and reflection data from the metamaterial slab obtained by the full-wave simulations were extracted. Using these data and the theoretical formulation for wave propagation through a chiral slab [60, p. 83] (including multiple reflections at the interfaces), the effective chirality parameter was calculated. For each of the seven discrete values of $N_v$, we performed 50 simulations of random metamaterial supercells. Figure 4a shows the calculated mean values and the standard deviations of the chirality parameter and the specific rotation of the colloid (calculated as $\Delta \theta/L = \Re(\kappa_0\theta/c)$). Only the real part of parameter $\kappa$ is plotted since the imaginary part is approximately ten times smaller due to very low absorption loss in silicon at the considered wavelengths. Therefore, the circular dichroism $\eta = \Im(\kappa_0\theta L/c)$ is much smaller than the optical rotation $\Delta \theta$. Additionally, the figure depicts the analytically calculated isotropic chirality parameter based on the polarizabilities of individual meta-atoms (see Table 1) and the Maxwell–Garnett mixing rule. [14, § 7.2.2] The theoretical curve is in good agreement with the statistical data from the full-wave simulations. In sharp contrast to the scattering loss, the chirality parameter monotonically increases with $N_V$.

Next, it is important to examine whether the periodicity of the metamaterial contributes to the magnitude of the chirality parameter. Figure 4b plots the real part of parameter $\kappa$ for four different supercell sizes $\Delta_z$. The volumetric concentration of the meta-atoms in all four cases was fixed to $N_v = 0.4$. As is seen, there is no strong correlation between the chirality parameter and the supercell size. Thus, we deduce that the strong chirality is due to the optimized meta-atoms and not due to their periodical arrangement.

Figure 4c depicts the frequency dispersion of the isotropic chirality parameter in the metamaterial with the volume fraction of meta-atoms $N_v = 0.6$. The dispersion of silicon was used from the experimental data in [42]. As is seen, in the frequency range from 200 to 600 THz, the colloid metamaterial is nonresonant, providing high isotropic chirality with very large bandwidth. At the higher frequencies, the meta-atoms sizes become comparable to the wavelength in water, and the colloid cannot be considered as a homogeneous metamaterial. Due to the low frequency dispersion and the dielectric nature of the meta-atoms, we expect the metamaterial to have relatively robust optical response in the presence of polydispersity of the silicon spheres or their imperfect assembly.

Let us compare the obtained chirality parameter with the previously reported values for random colloids suitable for large-scale fabrication. Table 2 presents such comparison for different optical characteristics. As is seen, the dielectric colloid metamaterial presented in this work exhibits three orders of magnitude increase for the chirality parameter and figure of merit (expressed as $\text{FOM} = |\kappa|/\text{Im}(\kappa)$) compared to previous publications. Such striking improvement is due to the two factors: The absence of plasmonic losses and scattering loss reduction due to the high-density colloidal arrangement of the dielectric meta-atoms. Interestingly, analogous three orders of magnitude difference is observed between magneto-optical FOMs of yttrium iron garnet and transition metal ferromagnets. [74, § 5.6.5] Due to the higher
magneto-optical FOM (lower insertion loss for the given isolation ratio), garnets rather than ferromagnets are usually employed as Faraday-rotation components in optical isolators. Likewise, chiral dielectric colloid proposed in this Letter has the advantage of low attenuation compared to conventional plasmonic colloids, which makes it a better candidate for multiple applications. It is also worth mentioning that the designed isotropic colloid metamaterial exhibits specific rotation more than two order of magnitude larger than that of natural substances like sugar \([75]\) and even one order of magnitude larger than that in anisotropic alpha-quartz crystal\([76]\).

One can notice that the isotropic polarizability \(|\text{tr}(\mathbf{\alpha}_{\text{em}})|/3\) of the individual meta-atoms proposed in this work is several orders of magnitude smaller than that from ref. \([73]\). This is due to the fact that in ref. \([73]\), the meta-atoms exhibit plasmonic resonances. In contrast, our metamaterial has nearly six orders of magnitude larger than that from ref. \([73]\). This is due to the fact that in ref. \([73]\), the meta-atoms exhibit plasmonic resonances which results in overall increase of the chirality parameter over that in the plasmonic colloid. At first glance, it may seem that by increasing the volume concentration \(N\) in the plasmonic colloid, one can achieve stronger chirality effect. However, doing so, while enhancing the chirality, also inevitably increases the absorption loss. The figure of merit introduced above takes into account this trade-off. Let us show that FOM does not depend on the volume concentration of the meta-atoms in sparse colloids like in ref. \([30,31,73]\). Since in this low-concentration case, chirality parameter is given by \((2)\) and the imaginary part of the refractive index \(\text{Im}(n) = N \text{Im}(\alpha_{\text{eff}})\) where \(\alpha_{\text{eff}} = \sqrt{\text{tr}(\mathbf{\alpha}_{\text{ee}})\text{tr}(\mathbf{\alpha}_{\text{mm}})/3}\), the figure of merit becomes

\[
\text{FOM} = \frac{\text{tr}(\mathbf{\alpha}_{\text{ee}})}{3 \text{Im}(\alpha_{\text{eff}})} \tag{5}
\]

Indeed, it depends only on the electrodynamic properties of the individual meta-atoms and not on the volume concentration of the colloid \(N\). To further demonstrate that the giant chirality effect reported in this work cannot be achieved with conventional plasmonic solutions, we take as an example colloid from ref. \([73]\) and calculate the transmittance of light \(I_r/I_{\text{inc}}\) propagating through it (reflections at interfaces are assumed negligible) for different volume concentrations \(N\)

\[
I_r(N) = I_{\text{inc}} \exp(-2k_0L \text{Im}(n(N))) \tag{6}
\]

Here, \(\text{Im}(n(N)) = \frac{N}{N_0} \text{Im}(n(N_0))\) and parameters \(N_0 = 6 \times 10^{21} \text{ m}^{-3}\) and \(\text{Im}(n(N_0)) = 9.5 \times 10^{-6}\) are listed in the first row of Table 2. Assuming \(\lambda = 750 \text{ nm}\) and \(L = 1 \text{ mm}\), we obtain the curve shown in Figure 5a. One can see that when the concentration of the plasmonic colloid is increased by merely two orders of magnitude, the transmittance drops by more than six orders of magnitude. Therefore, the colloid becomes impermeable at high volume concentrations. Note that expression \((6)\) does not explicitly include packing factor \(W(N_v)\) as in \((4)\), and \(I_r\) depends linearly on concentration \(N\). This assumption is valid in the range of small volume fraction \(N_v\) (\(N_v\) does not exceed \(2.5 \times 10^{-3}\) in the range plotted in Figure 5a, assuming that meta-atom volume is \(V_0 \approx 4.2 \times 10^{-21} \text{ m}^3[73]\)).

For comparison, Figure 5b depicts the light transmittance versus the volume concentration for the silicon colloid proposed in this letter. Since in this case the concentration is high (the maximum concentration \(N_{\text{max}} = 5.2 \times 10^{20} \text{ m}^{-3}\) is shown in Table 2), the transmittance intensity is now a function of the packing factor

\[
I_r(N) = I_{\text{inc}} \exp(-2k_0L \text{Im}(n(N))W(N_v)) \tag{7}
\]

Here, we assumed, likewise, that \(\lambda = 750 \text{ nm}\) and \(L = 1 \text{ mm}\). As is seen from Figure 5b, the attenuation does not monotonically increase with increasing the volume concentration but, on contrary, it starts decaying when \(N > 0.22N_{\text{max}}\). This is a consequence of the scattering suppression in the quasicrystalline regime. Note that for this metamaterial, the chirality parameter is monotonically increasing with \(N_v\), as seen in Figure 4a. Therefore, the silicon metamaterial provides several orders of magnitude chirality enhancement compared to the plasmonic colloids, assuming the same level of light attenuation.

**Figure 4.** a) The real part of the intrinsic (isotropic) chirality parameter and specific rotation of the metamaterial versus the volume fraction. The plotted data corresponds to the operational frequency \(f = 400 \text{ THz}\). b) The chirality parameter calculated for four different supercell sizes. In all the cases the volume fraction was fixed to \(N_v = 0.4\). c) Frequency dispersion of the isotropic chirality parameter and specific optical rotation of the random colloid metamaterial with the volume fraction of meta-atoms \(N_v = 0.6\).
3. Conclusion

In summary, we have designed a dielectric metamaterial consisting of a random colloid of specifically designed silicon nanoparticles which exhibits record-high isotropic chirality with significantly suppressed incoherent scattering. The suggested geometry of the chiral constituent meta-atoms favors large-scale mass production. This is in strong contrast with the previous designs of dielectric chiral metamaterials and colloids which either suffered from low writing speeds\cite{77} or were suitable only for low colloid concentrations (such colloids were achieved by dispersing nanoparticles from a 2D area into a bulk solution).\cite{33,34,78} The chirality can be further increased by optimizing the geometry of the meta-atoms using, for example, via inverse design.\cite{79} It should be mentioned that although in this work, we concentrated on sub-wavelength dielectric meta-atoms, it is possible to extend this study toward meta-atoms of larger sizes. In this case, the regime of strong Mie resonances can be achieved, which would further increase the isotropic chirality of the colloid by one or several orders of magnitude. The higher order multipole moments in isotropic samples do not contribute to chirality due to orientational averaging [17, p. 142, 227], and therefore, the theory presented in this letter can be still applied. Moreover, the QCA analysis demonstrated here can be extended to the case of meta-atoms with higher order multipole moments (see e.g., ref. [80]).

**Supporting Information**

Supporting Information is available from the Wiley Online Library or from the author.

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**Conflict of Interest**

The authors declare no conflict of interest.
