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Superchiral hot-spots in “real” chiral plasmonic structures†

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Light scattering from chiral plasmonic structures can create near fields with an asymmetry greater than the equivalent circularly polarised light, a property sometimes referred to as superchirality. These near fields with enhanced chiral asymmetries can be exploited for ultrasensitive detection of chiral (bio)molecules. In this combined experimental and numerical modelling study, we demonstrate that superchiral hot-spots are created around structural heterogeneities, such as protrusions and indentations, possessed by all real metal structures. These superchiral hot-spots, have chiral asymmetries greater than what would be expected from an idealised perfect structure. Our work indicates that surface morphology could play a role in determining the efficacy of a chiral structure for sensing.

Introduction

Using the tools of modern nanofabrication, periodic arrays of complex nanostructures of the same design can be routinely manufactured. Although, derived from the same idealised design, the apparently identical individual nanostructures have unique geometric variations, caused by intrinsic surface roughness or structural defects. The presence of these structural irregularities causes highly localised enhancements of EM fields within the overall near field region.¹ It has been accepted that when ensembles of nanostructures are considered within an array, the resulting linear optical response (*e.g.* reflection/transmission) is dependent upon the average of the individual nanostructures’ morphologies. Consequently, an array of real structures can be considered a broadened version of that from an array of idealised structures.² However, this spatially averaging argument breaks down when one considers Raman scattering and non-linear optical responses, where observed spectroscopic responses are dominated by contributions from localised hot-spots associated with the geometric roughness and defects.³ In addition, for complex structures that consist of multiple elements, surface roughness can alter the level of inductive coupling between elements, and hence modify optical response.

In this study we have investigated how surface roughness influences the level of the chiral asymmetries of near fields

created by the optical excitation of chiral plasmonic structures. Near fields generated by light scattering from nanostructures can, in localised regions of space, possess a greater level of chiral asymmetry than comparable circularly polarised light (CPL), a property sometimes referred to as superchirality. Such fields with enhanced chiral asymmetry can be exploited for ultrasensitive detection of chiral (bio)molecules.^{4–7} Numerical simulations used in previous studies to understand the chiral asymmetries of these field have relied on idealised models of the chiral structure.^{6–8} In this study we have attempted to understand the influence of surface roughness by using a “real” model for the chiral nanostructure, a gammadion, directly derived from atomic force microscopy images, in periodic numerical simulations. The simulations are validated, by comparison with experimental circular dichroism spectra data. Simulated spectra obtained from the real model are in better agreement with the experimental data. Our study reveals that the idealised model underestimates the level of hybridisation, mediated by inductive coupling, between the arms of the gammadion structure. Significantly, surface roughness results in localised regions, close to the walls of the structure, with chiral asymmetries up to 2–3 times greater than those obtained from the idealised model. This suggests that surface roughness plays a role in determining the effectiveness of a chiral plasmonic structure for bio-detection applications.

Results

AFM characterisation of structures

The gammadion structure studied has been chosen because it most closely matches those used in previous chiral sensing

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studies.^{4,6} Representative atomic force microscopy (AFM) images of the structures are shown in Fig. 1(A and B). Key parameters have been calculated by taking the mean of 5 measured values from the micrograph, these are shown on a plane view schematic of the structures in Fig. 1(C). The gammadion consists of a central cross whose arms are 130 nm wide and 425 nm long and so the structure fits within a 425×425 nm square. The arms connected to the centre cross of the structure are thinner at 110 nm. The spacing between the nanostructures is 375 nm.

The surface roughness of a given area is parameterised by the root-mean-square roughness (R_q).⁹ For the idealised gammadion nanostructures the R_q will be 0 as the surfaces are planar. For the real gammadion, the R_q calculated at its top face is approximately 3.40 nm. Compare this with the glass substrate with a $R_q \sim 0.45$ nm.

Real and ideal models

The idealised model (left handed, LH), which simplifies the geometry and morphology of the structures, is shown in Fig. 2(A). Most evident simplifications are that the faces and the vertical profiles of the gammadion are made planar: they do not account for surface roughness, sloping or other morphological defects which occur in the fabrication of the nanostructure arrays.

The real model, shown in Fig. 2(B) is constructed directly from an AFM micrograph of a single gammadion structure. The micrograph of the entire array, it is cropped so it contains only one structure, selected arbitrarily, and the glass substrate is removed. The file is converted to a format that can be read by the numerical modelling software. This structure includes morphological defects and so it is described here as the 'real' structure.

Fig. 2(C) shows the profile of the 'real' gammadion model at 0, 50 and 100 nm above the glass substrate. The structure shows significant sloping as the height of the structure increases. Tip convolution can overestimate the lateral dimensions of the nanostructures, making protrusions from the surface appear larger and making holes appear smaller.¹⁰

(A) Ideal



(B) Real



(C)

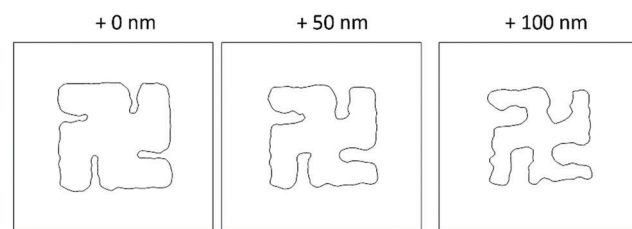


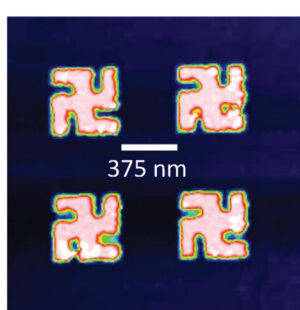
Fig. 2 A plan and 3-D view of the (A) ideal and (B) real models of LH gammadion nanostructures used in numerical simulations. (C) Cut slice profile of the 'real' structure 0, 50 and 100 nm above the glass substrate. The structure shows significant sloping.

Therefore, it is likely that the micrograph overestimates the extent of the sloping.

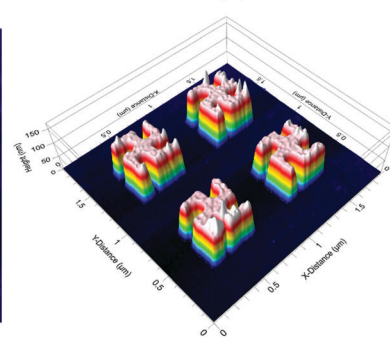
Experimental and simulated CD spectra

Both experimental and simulated spectra for both gammadion enantiomorphs are displayed in Fig. 3. As expected, both measured and simulated, spectra for LH and RH structures

(A)



(B)



(C)

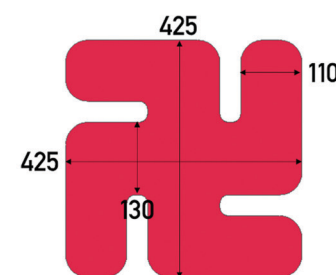


Fig. 1 AFM images of a LH gammadion displayed in (A) plane and (B) 3-D topological views; (C) is a schematic of the structures which displays relevant dimensions.



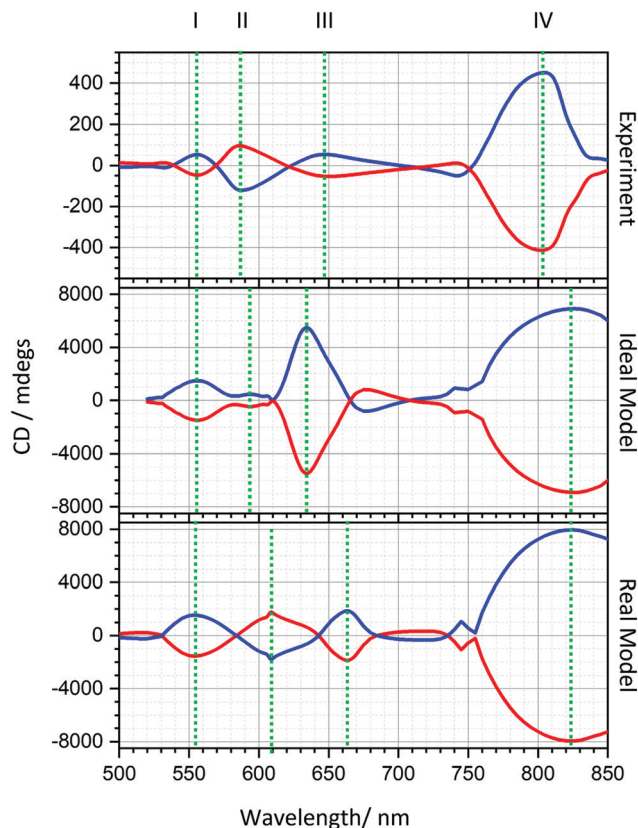


Fig. 3 Comparison of the experimental CD spectra (top) for LH (red) and RH (blue) gammadia, with spectra derived from the ideal (middle) and real models (bottom). The positions of modes I–IV are highlighted in the spectra with dotted green lines.

are equal in magnitude but opposite in sign: they are mirrored around the 0 millidegree line of the plot. The experimental data has four pronounced resonances, labelled I–IV in order of increasing wavelength, that have corresponding features in the simulated spectra derived from both models. The positions of these band for measured and simulated spectra are given in Table 1.

The level of CD observed in the simulated spectra are approximately an order of magnitude larger than those observed experimentally. A reduction by a factor of 2 can be attributed to the ‘chessboard’ fabrication strategy, which reduces the writing time of the lithography tool by patterning only half of the substrate (see ESI†). Further reductions must be due to fabrication defects such as missing nanostructures, or missing parts of the structure which are not accounted for in either the ideal or real models.

Table 1 The (average) experimental positions of modes I–IV are compared with those derived from real and ideal models

	Wavelength position/nm			
	I	II	III	IV
Experiment (average LH & RH)	556.7	588.2	649.4	794.9
Ideal model	556.0	596.4	634.0	826.0
Real model	555.0	608.9	662.9	824.2

It is readily apparent that, predictably, the simulations based on the real model provide the best qualitative agreement with the experimental data. Although both the real and ideal models replicate modes I and IV equally well, there is divergence in how effectively they reproduce modes II and III. For both models, modes II and III show shifts relative to the experimental data. However, more significantly in contrast to the real model, the ideal model does not replicate the relative intensities (resonance III is much larger), nor the signs (resonance III have opposite signs) of the experimentally observed resonances. Finally, the separations between modes I and III are underestimated by the ideal, and over-estimated by the real models.

Simulated 3-D field plots

To rationalise the differences between ideal and real models requires an understanding of the origins of the resonances, which can in part be gained from maps of electric field magnitude $|E|$ and chiral asymmetries. The chiral asymmetry of a field can be conveniently parameterised using optical chirality density (C)¹¹

$$C \equiv \frac{\epsilon_0}{2} \mathbf{E} \cdot \nabla \times \mathbf{E} + \frac{1}{2\mu_0} \mathbf{B} \cdot \nabla \times \mathbf{B}, \quad (1)$$

where \mathbf{E} and \mathbf{B} the time harmonic electric and magnetic fields and ϵ_0 and μ_0 are electric permittivity and magnetic permeability.

Three dimensional plots of the $|E|$ and C have been generated to aid interpretation. Mapping the entire ranges for the $|E|$ and C is ineffective for 3-D visualisation, as then only the field at the boundaries of the simulation are visible. By limiting the mapped fields to only those above a certain threshold, allows localised ‘hot-spot’ regions of both $|E|$ and C to be easily observed. Analogous 2-D plots are shown in ESI†.

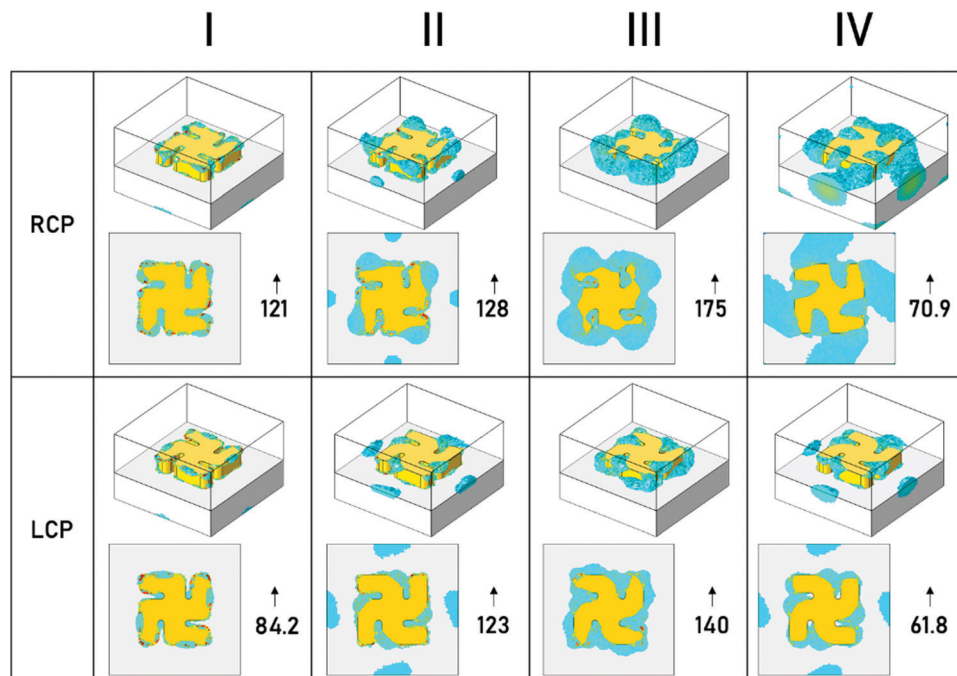
3-D field plots for $|E|$ and C have been generated for both incident left and right circularly polarised light (LCP and RCP) at the wavelengths of modes I–IV. C values have been normalised against that of RCP at the wavelength of interest. For $|E|$, maps are produced at two thresholds, 10 and 20 V m⁻¹, Fig. 4 and 5, while a threshold of 2.5 was set for the C maps, Fig. 6. For C map the highest and lowest values are given to provide guide to the level of the dynamic range. While the largest value is given for the $|E|$, maps.

The $|E|$ field plots for both the real and idealised models display qualitatively similar behaviour. The fields for resonances I–III are all localised on the nanostructure, while for LH (RH) structures under RCP (LCP) illuminations fields are observed to connect neighbouring structures. This observation is consistent with previous work which has shown that resonances I–III are localised modes, associated with inductive coupling between the constituent elements of the gammadiom structure, and resonance IV is a lattice (or Bloch) mode arising from the periodicity of the structures.¹²

Additionally, for both ideal and real structures the most intense fields are associated predominately with side walls of the nanostructure. For the real structure there are localised hot-spots associated with protrusions on the top rough Au



(A) Ideal Model



(B) Real Model

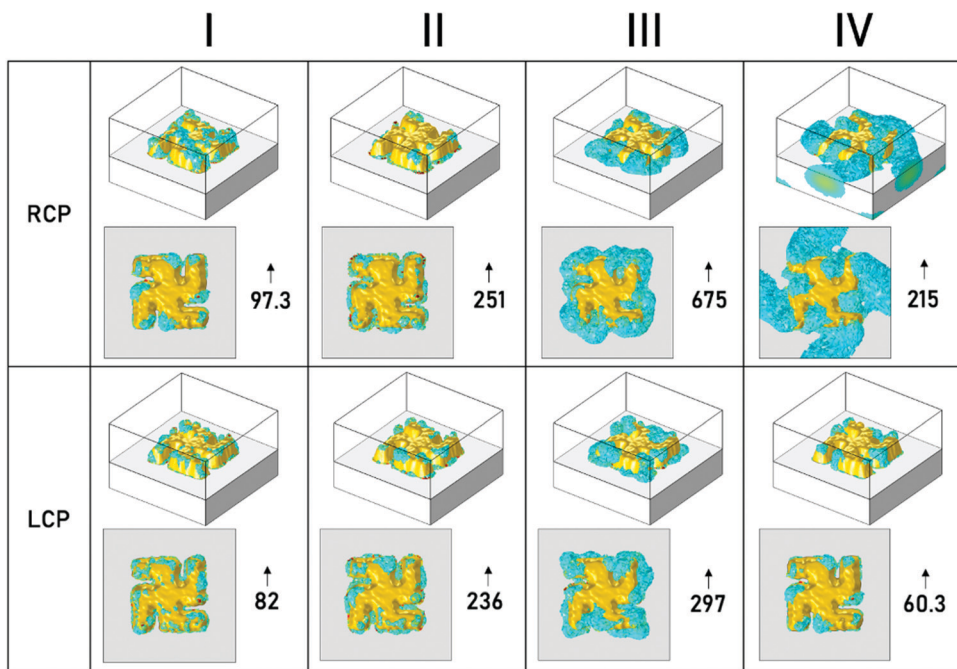


Fig. 4 3-D electric field plots for incident LCP and RCP light, with a 10 V m^{-1} threshold, generated from (A) real (B) idealised models. The largest value of $|E|$ is given with an up arrow in each inset.

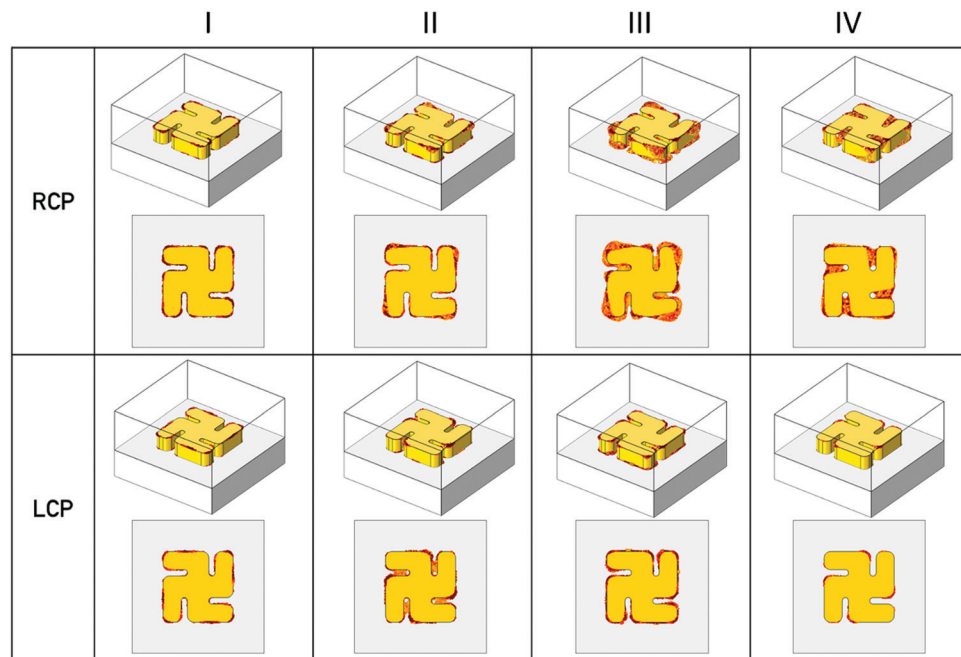
surface, which are not present for flat top surface of the idealised model.

From the C maps it is apparent that regions with the highest level of chiral asymmetry (C) are located at the bottom of the

gammadiion structure, in the vicinity of the Au-quartz interface, for both the ideal and real models. The largest values of C for both models are observed for resonance III, which in previous work has been demonstrated to be the most sensitive for the



(A) Ideal Model



(B) Real Model

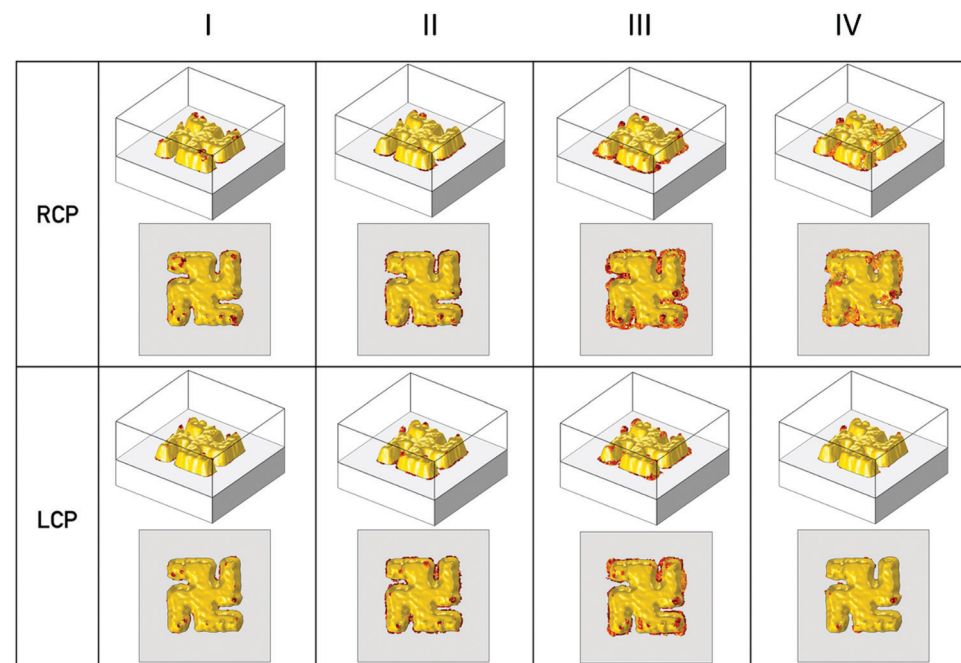


Fig. 5 3-D electric field plots for incident LCP and RCP light, with a 20 V m^{-1} threshold, generated from (A) real (B) idealised models.

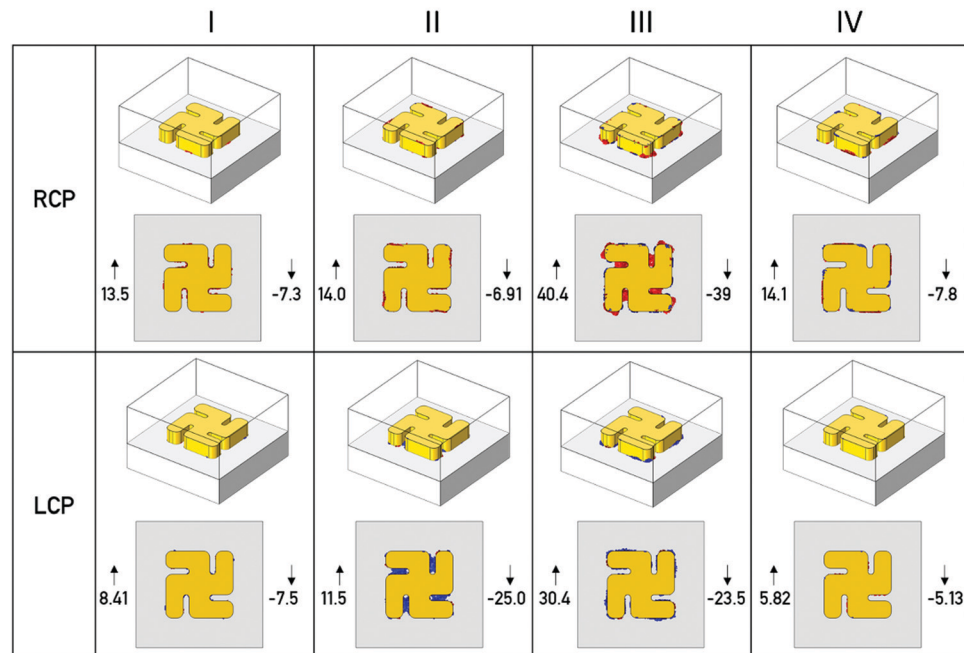
detection of chiral (bio)materials.⁸ In contrast to the $|E|$ maps, there are no analogous hot-spots of enhanced optical chirality associated with protrusions of the rough top surface. However, the maximum C values, for a particular enantiomorph and helicity of light, are up to factor of ~ 3 times greater for the real compared to the ideal structure.

Discussion

It is unsurprising that the simulations based on the real structure are in closer agreement with the overall measured experimental spectra. However, the ideal and real models do both replicate mode IV, the lattice mode, equally well. This can



(A) Ideal Model



(B) Real Model

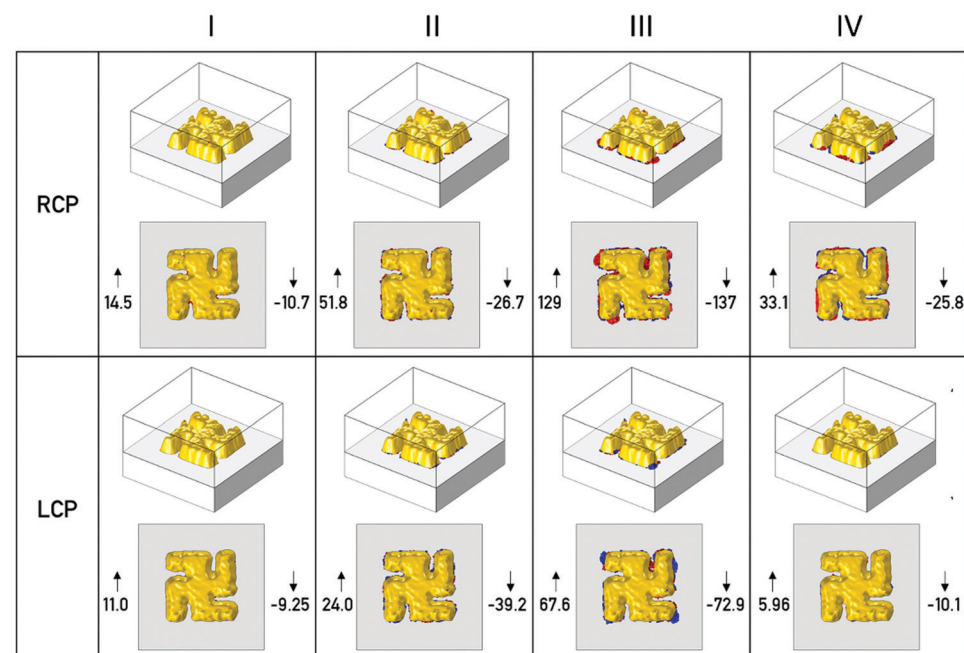


Fig. 6 Time averaged normalised (with RCP) optical chirality plots above a threshold of 2.5 (red) (−2.5 (blue)) for (A) real and (B) ideal models. The minimum (↓) and maximum (↑) values for each resonance and light handedness are included.

be attributed to the fact that the lattice mode will be predominately controlled by the periodicity and symmetry of the array. In contrast the localised modes, I–III, which arise through the inductive coupling of the individual rod elements of the gammadion structure would be sensitive to morphological heterogeneity of the real structures. The presence of structural

heterogeneity would be expected to influence the properties of the gammadion through two mechanisms. Firstly, the presence of randomly distributed protrusions and indentations on the surfaces of the gammadion act as a symmetry breaking perturbation. In isolation the ideal gammadion structure placed on a substrate belongs to the C_4 point group. The presence of

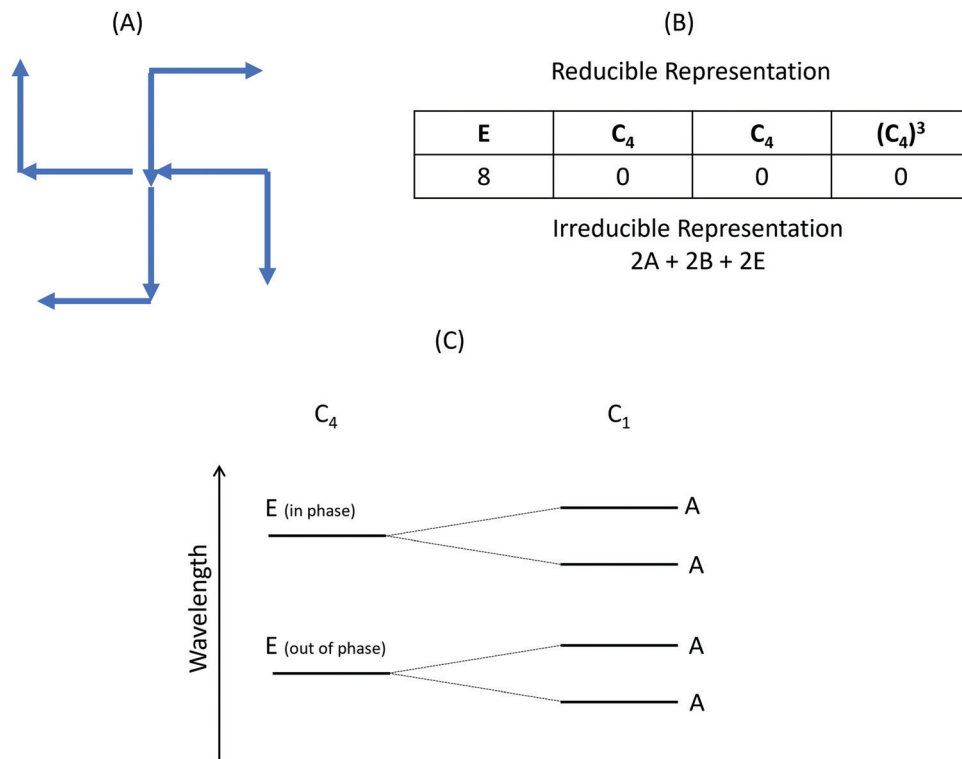


Fig. 7 (A) Basis set of vectors used in the symmetry analysis; (B) the reducible and irreducible representations produced by the basis set of vectors. (C) An energy level diagram showing the effects of a symmetry reducing perturbation.

random structural heterogeneity effectively reduces the symmetry to C_1 . Analogous to the procedure used in molecules, a symmetry analysis of the optical modes of the gammadiion structure can be carried out. The analysis involves a basis set of 8 vectors, mimicking dipoles, associated with the rods that make up the gammadiion structure, Fig. 7(A). For the C_4 point group this analysis gives four modes 2A, 2B and 2E, with E modes being doubly degenerate, (Fig. 7(B)). The two E modes represent symmetric and antisymmetric combinations of the dipoles associated with the rods. Only the E modes would be excited by the incident circularly polarised light and the B mode is optically dark. Under the symmetry breaking perturbation of the surface roughness the degeneracy of the E modes is lifted, causing a splitting in to two A modes, this is illustrated in an energy level diagram, Fig. 7(C). This symmetry reducing perturbation would be expected to cause a broadening of the CD resonances.

The differences in the positions and line shapes of modes I-III predicted by ideal and real models cannot be solely justified using the symmetry reducing perturbation argument. Specifically, since surface roughness modifies both the intensities and chiral asymmetries of the near fields, creating hot-spots, one would expect that this would alter the level of inductive coupling between the rod elements of the gammadiion structure. Intuitively, one would assume that the presence of hot-spots would enhance inductive coupling between the consistent rods. In previous work¹² it has been shown that the wavelength separation between modes I and III (subsequently

labelled S_{I-III}) scales with increasing coupling between neighbouring rod elements. The value of S_{I-III} derived from the ideal model (78 nm) is smaller than that observed experimentally (92.7 nm), implying that it underestimates the magnitude of the inductive coupling. In contrast the S_{I-III} obtained using the real model (107.9 nm) is larger than the experimental value, thus overestimating the level of coupling. These observations imply that the inclusion of “real” structural heterogeneity in numerical simulations are necessary to successfully mimic the level of inductive coupling observed experimentally. The over-estimate of coupling predicted by the real model has two potential origins. Firstly, the morphology of the chosen structure may display a higher level of structural heterogeneity than the average. Alternatively, the exaggerated level of slope of the side wall of the nanostructures observed in AFM images, caused by tip effects, could be the cause. Such tip convolution effects would result in the separation between the bases of the arms being closer than they are in reality, thus increasing the level of coupling.

To summarise the current study illustrates the advantages of using realistic structural models for the numerical modelling of the optical properties of metamaterials. Field maps derived from these “realistic” models reveal the presence of localised regions, or hot-spots, of enhanced $|E|$ and C which have a greater magnitude than the maximum values obtained from idealised models. An important implication of structural heterogeneity creating enhanced “superchiral” hot-spot in the vicinity of nanostructure, is that surfaces roughness could enhance the enantiomeric sensing ability of chiral nanostructures.



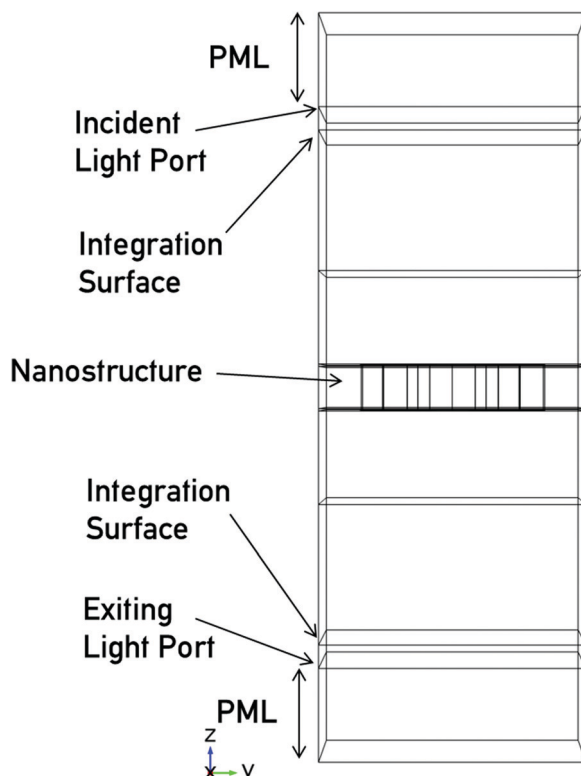


Fig. 8 Geometry of the simulation with PML, light ports and integration surfaces identified.

Methods

Numerical modelling

Numerical simulations of electromagnetic fields were performed using the COMSOL Multiphysics platform. The nanostructure is surrounded by a cuboid representing a unit cell, with the x - and y -dimensions defining the periodicity of the metamaterial, as calculated from AFM images. A schematic showing the simulation geometry is shown in Fig. 8. The z -dimensions of the cell are sufficiently large ($\geq \lambda_{\max}/2$) that near-fields generated by the nanostructures do not extend to any integration surfaces above and below, the total height of the cell is 1600 nm. The unit cell is split up into layers of varying thickness. The top 200 nm is a perfectly matched layer (PML) which absorbs all reflections from the nanoparticle. The surface at 200 nm is the excitation port, from where light originates and its polarisation is specified. 100 nm below the excitation port is an integration surface where reflected power is measured. The gammadion is positioned in the centre of the cuboid. 300 nm from the bottom is another integration surface where transmitted power is calculated. 200 nm from the bottom is the radiation exiting port, followed by a 200 nm PML layer. In order to simulate an array of gammadiions, Floquet periodic conditions are applied at the x - and y -boundaries.

The material properties of the structure can then be implemented. To replicate the experimental CD work from the previous section, the gammadion nanostructure is placed onto

the glass substrate with a refractive index equal to 1.5 then covered with water with refractive index 1.33.

Gammadion sample fabrication

The gammadiions structures were fabricated using an electron beam lithography process. Quartz glass slides were cleaned under ultrasonic agitation in acetone, methanol and isopropyl alcohol (AMI) for 5 minutes each, dried under N_2 flow and exposed to O_2 plasma for 5 minutes at 100 W. A PMMA resist bilayer (AllResist 632.12 50k in anisole and 649.04 200k in ethyl lactate) was then spun at 4000 rpm for 1 minute and baked at 180 °C for 5 minutes in between spins. A 10 nm aluminium conducting layer was evaporated on the substrates using a PLASSYS MEB 550s evaporator. Patterns were designed on the CAD software *L-Edit* and written by a Raith EBP-G 5200 electron beam tool operating at 100 kV. The resist was developed in 3:1 MIBK:IPA solution at 23.2 °C for 1 minute, rinsed in IPA (5 s) and water before drying under N_2 flow. A 5 nm nichrome adhesive layer was then evaporated below a 100 nm gold layer. The process was completed with a lift-off procedure in acetone at 50 °C overnight and then agitated to remove all remaining resist and excess metal.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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