Size-dependent Optical Response of Magneto-plasmonic Core-shell Nanoparticles

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ABSTRACT

Functional properties tunability prospect of multi-layered and multi-metallic nanoparticles with respect to their emerging applications has always been the area of attraction. Magneto-plasmonic nanoparticles (MPNPs) present the possibility to exhibit their tuneable magnetic and optical properties with broad applications in magnetic resonance imaging (MRI), cancer therapy, photovoltaic solar cells, biological microscopy, optical imaging, and biosensing. Present studies found that the optical properties (absorption and scattering efficiencies) of MPNPs using Fe, Co and Ni as magnetic materials coated with single noble metal (Au or Ag) or double (both Au & Ag) coating layer, can be effectively tuned with the controlled size of core and shell layers. It is found that absorption and scattering LSPR peaks occur in Ultraviolet (UV) and visible region of EM for single core-shell MNP@Ag nanoparticles respectively. The absorption LSPR are found in the visible region, and scattering LSPR in NIR region and also get blue shifted for single core-shell of MNP@Au nanospheres. In case of double core-shell layers of MNP@Ag@Au nanoparticles, the absorption and scattering LSPR peaks show spectra in visible and NIR region of EM spectrum and the absorption LSPR peak shifts from visible to UV region and scattering LSPR are found in NIR region with broadening peaks of EM spectrum. It is concluded that the LSPR peak depends upon the thickness of the shell, size of core and material. The shifting of optical peaks towards the longer wavelengths depends on gold shell thickness whereas optical efficiencies depend on silver (as core or shell) thickness.

Keywords: Magneto-plasmonic nanoparticles (MPNPs), Core-shell thickness, Absorption and scattering efficiency.

1 Introduction

Nanoparticles (NPs) with dimensions of 1 nm to 100 nm have attracted significant interest due to their fascinating properties viz. optical, magnetic and optoelectronic that are superior to their bulk counterparts because of large surface area to volume ratio [1, 2]. Nanoparticles have interesting applications in the fields of catalysis, bio-imaging, drug targeting and sensing. Significant progress has also been made in the synthesis of nanoparticles from any kinds of materials with the desired quality, quantity, and yield required for the systematic investigation of their peculiar properties and applications. Many metals (e.g., alkali metals and to some extent noble metals viz. Au and Ag) can be treated as free-electron systems whose electronic and
optical properties are determined by the conduction electrons alone. The collective oscillations of free conduction electrons (LSPR) of metallic NPs, (e.g., Ag and Au) are observed when illuminated with the light of plasmon wavelength and is called localized surface plasmon resonance (LSPR) [3]. The features of an LSPR depend upon the dispersive property of the material and its interaction with the surroundings. Metallic nanoparticles exhibiting localized surface plasmon resonances can show strong optical resonance for visible and near-infrared (NIR) wavelengths. Noble metal NPs have received considerable interest during the last few decades in the field of plasmonic, material sciences and biomedicine, due to their unique and intense optical, electronic, catalytic and tunable plasmonic properties [4-9]. On the other hand, magnetic nanoparticles (MNPs) have lesser free electron contribution to the optical properties as compared to plasmonic NPs, due to which they do not exhibit LSPR, but these MNPs possess spontaneous magnetization. Magnetic nanoparticles provide target ability or control over a motion which makes their use in a biosensor, magnetic fluids, information storage, magnetic bioseparation, and catalysis [10-16]. Recently, a great interest has arisen towards the hybrid structures of NPs to investigate their optical properties. The hybrid NPs can present enhanced novel features unavailable from single-element NPs. The advantage of hybrid NPs, compared to their single component counterparts, can be achieved in two ways. First, they can offer multiple functionalities (by coating) and secondly through coupling between their different components (alloying) by combining different classes of physical properties related to their constituents [17-19]. Corporate examples are binary (Au-Co, Ag-Co, Ag-Ni) or ternary (Fe-Ag-Si) magneto-plasmonic NPs with unique magneto-optical properties for various applications. Magnetic nanoparticles are often used in diagnostic [20] or drug delivery purposes [21], however, they can’t be used directly for in-vivo biological applications because they are sensitive to agglomeration and are corrosive. Magnetic nanoparticles can be protected by using noble metal coating, leading to core-shell nanoparticles called magneto-plasmonic nanoparticles (MPNPs). The coating not only protects the magnetic nanoparticles from oxidation, thus enhancing their stability and compatibility, but also provides a new platform for further functionalization of nanoparticles for optical properties [22-23]. The main advantage with MPNPs is to use the magnetic and plasmonic properties with an external magnetic field to provide boost to the local EM field and thus enhancing magneto-optic effects. The available experimental methods to coat magnetic nanoparticles are by directly depositing noble metals onto the core surface or by using a chelating material between the core and the shell. The optical properties of MPNPs are varied and can be tuned by changing their size, shell thickness, shapes and dielectric environment of particles [24]. Thus, the calculated optical properties (scattering and absorption efficiencies) are the relevant quantities for real applications of MPNPs. This paper highlights the absorption and scattering efficiencies of MPNPs of single core-shell and multiple core-shell nanospheres considering Fe, Co, and Ni as magnetic materials at core and Au and Ag noble metal nanoparticles as coatings (shells). We found that the absorption and scattering efficiencies of considered single and multi-core-shell MPNPs depend on core-shell ratio and thus can be tuned. The calculated spectra of efficiencies are found in visible and near-infrared (NIR) region of electromagnetic spectrum.

2 Calculation Methods

To investigate the optical properties (absorption and scattering) of nanoparticles with light (electromagnetic radiation) interaction, one has to obtain a solution to Maxwell's equations with corresponding boundary conditions on the spherical surface and the solution to the scattering problem for homogeneous sphere was proposed by Gustav Mie in 1908. Mie theory is the first analytical solution for the Maxwell equations to describe the scattering of light by spherical particles made of arbitrary materials. Mie assumed that optical constants of metals, which were obtained for bulk materials, are also applicable to nanoparticles which were at least one order of magnitude smaller than the wavelength of light, and luckily this was almost right for the size, wavelength and the material he considered. Mie theory takes less time, is cost effective and gives the same results as compared to other methods. However, for complex geometries there are other methods such as BEM, FEM, DDA and FDTD which are frequency and time domain numerical method. For the
spherical Au and Ag nanoparticles, \( Q_{\text{abs}} \) and \( Q_{\text{sca}} \) were calculated on the basis of Mie theory for homogeneous spheres. The Mie total extinction (\( Q_{\text{ext}} \)) and scattering efficiency (\( Q_{\text{sca}} \)) for a homogeneous sphere are expressed as infinite series:

\[
Q_{\text{ext}} = \frac{2}{\lambda^2} \sum_{n=1}^{\infty} (2n + 1) \text{Re}[a_n + b_n]
\]

\[
Q_{\text{sca}} = \frac{2}{\lambda^2} \sum_{n=1}^{\infty} (2n + 1)[a_n^2 + b_n^2]
\]

\[
Q_{\text{abs}} = Q_{\text{ext}} - Q_{\text{sca}}
\]

\[
a_n = \frac{m \varphi_n(mx)\varphi'_n(x) - \varphi_n(x)\varphi'_n(mx)}{m \varphi_n(mx)\varepsilon_n(x) - m\varepsilon_n(x)\varphi_n(mx)}
\]

\[
b_n = \frac{\varphi_n(mx)\varphi'_n(x) - m\varphi_n(x)\varphi'_n(mx)}{\varphi_n(mx)\varepsilon'_n(x) - m\varepsilon_n(x)\varphi_n(mx)}
\]

where \( m \) is the ratio of refractive index of the sphere \( n \) to that of the surrounding medium \( n_m \), \( x \) is the size parameter given as \( 2\pi n_m R/\lambda \), \( \varphi_n \) and \( \varepsilon_n \) are the Riccati-Bessel functions, and the prime represents first differentiation with respect to the argument in parentheses. Numerical calculations of the Mie series are performed at discrete points in the wavelength range from 200 to 1200 nm.

Calculations of the optical properties such as absorption and scattering efficiency of considered core-shell nanoparticles were performed by using a computer code employing Mie scattering for multi-layered spherical geometry developed by Ovidio Pena-Rodriguez team [25]. Mielab offers tools to perform several tasks related to the calculation of optical properties of nanoparticles and particles ensembles. Input parameters are the thickness of each layer, real (n) and imaginary (k) parts of the materials complex refractive index corresponding to energy in electron volt (eV), and the last parameter is the refractive index of the surrounding medium. The complex refractive index of water is 1.33+0i chosen as a surrounding medium to MPNPs because it is the main component of environments encountered in biomedical applications of MPNPs. The complex dielectric constant of magnetic and plasmonic (noble) material is obtained from Johnson and Christy [26-27]. The thickness or size of core-shell layers are not represented by the single radius, but are defined by the total radius (\( R_{\text{total}} \)) for the structure, and given as the sum of the radii of the magnetic core (\( R_{\text{con}} \)) and noble metal shell (\( R_{\text{shell}} \)). Plasmonics properties of core-shell nanoparticles are calculated for two sets of the nanostructure, firstly: by fixing the core and varying shell thickness for the total size and then changing the size of the core and varying the shell size also in total size. Secondly, by changing the core size and shell size as in the first case and fixing the size of the outer shell for total size.

### 3 Results and Discussion

Presently, optical properties are represented by scattering (\( Q_{\text{sca}} \)) and absorption (\( Q_{\text{abs}} \)) efficiencies. Since different optical properties of MPNPs are only associated with scattering and absorption efficiencies, therefore, these are the relevant representative parameters for real applications of MPNPs. Therefore, we have calculated the optical spectra for these important parameters (\( Q_{\text{sca}} \) and \( Q_{\text{abs}} \)) of core-shell magnetic-plasmonic nanoparticles with the different core (Fe, Co, Ni)-shell (Ag, Au) materials and thickness, considering nanospheres of various sizes between 20 nm to 70 nm.
3.1 Single Core-shell MPNPs

In the single core-shell study of nanospheres, we studied the magnetic nanospheres of Fe, Co and Ni coated with Ag or Au or both plasmonic materials. The optical properties were calculated as a function of wavelength, starting with the smallest size of 10 nm of the core and then changing the shell thickness from 10 nm to 60 nm. The core size was changed from 10 nm to 60 nm and the thickness of the shell was varied to complete the total size of core-shell up to 70 nm of considered nanospheres. We considered the maximum-height and -broadening of peak to optimize the shell thickness. In this way, we optimize the shell thickness for each considered core size of nanospheres.

3.1.1 MNP@Ag Nanoparticles

Figure 1(a) represents the LSPR peaks for Fe@Ag core shells and only the maximum peaks for varying core size and shell thickness have been shown. The core size is considered to be increasing from 10 nm to 60 nm with an increment of 10 nm. The LSPR peak is found at 413 nm for 10 nm core with 20 nm shell thickness and other peaks are at 382 nm, 362 nm, 361 nm, 365 nm and 360 nm for 20-60 nm core size with 10 nm shell thickness for each core size. As the size of shell increases for each considered core size, the peak position shift towards the higher wavelength with a decrease in efficiency. With the increase in core size the peak position shifts to lower wavelength from 382 nm to 360 nm as shown in Figure 1(a). Thus, the LSPR absorption peaks show the spectra near Ultraviolet (UV) regime in electromagnetic (EM) spectrum.

Figure 1(b) shows absorption spectra representing various peaks of Co@Ag at wavelength 414 nm for 10 nm core sizes with 20 nm shell thickness, and 396 nm, 366 nm, 367 nm, 374 nm, 370 nm for 10 nm shell thickness of remaining 20 to 60 nm core size respectively. As the size of shell increases for each considered core size, the peak position shifts toward the higher wavelength with a decrease in efficiency. With an increase in core size, the peak position shifts to a lower wavelength from 414 nm to 366 nm. Thus, the LSPR absorption peaks show the spectra near the UV regime in EM spectrum. Similarly, for Ni@Ag core-shell, we get various peaks at wavelength 415 nm for 20 nm shell thickness, and 405 nm, 438 nm, 369 nm, 377 nm, 393 nm for 10 nm shell thickness of each considered 10 to 60 nm core size respectively. As the shell thickness increases for each considered core size, the peak position shifts toward the higher wavelength with decreasing efficiency. With the increase in core size, the peak position shifts to a lower wavelength between 415 nm and 369 nm as shown in Figure 1(e). Thus, the overall LSPR absorption peaks for MPNPs show the spectra near the UV regime on EM spectrum.

The scattering efficiencies of MNPs@Ag core-shell have also been calculated, and we get various peaks at wavelengths 580 nm, 580 nm, 580 nm, 583 nm, 583 nm, 580 nm for 10 to 60 nm core-shells respectively for Fe@Ag magneto-plasmonic nanoparticles as shown in Figure 1(d). With the increase in shell thickness for considered core size, the peak position shifts toward the higher wavelength with a decrease in efficiency. Thus, the LSPR scattering peak shows the spectra in the visible region of electromagnetic spectrum. Similarly, Figure 1(e) shows various peaks of Co@Ag at wavelengths 580 nm, 580 nm, 580 nm, 583 nm, 583 nm, 580 nm for 10 to 60 nm core sizes with shell thickness 60 nm to 10 nm respectively. As the shell thickness increases for considered core size, the peak position shifts toward the higher wavelength with a decrease in efficiency. With the increase in core size, the peak position shifts to higher wavelength from 580 nm to 583 nm. Thus, the LSPR scattering spectra is found in the visible region of electromagnetic spectrum. Further, for Ni@Ag core-shell, we get various peaks at wavelengths 580 nm, 580 nm, 583 nm, 583 nm, 586 nm, and 590 nm for 10 nm to 60 nm core sizes with shell thickness 60 nm to 10 nm respectively. As the shell thickness increases for each considered core size, the peak position shifts toward the higher wavelength with a decrease in efficiency. With the increase in core size and decrease in shell thickness the peak position shifts to higher wavelength from 580 nm to 590 nm as shown in Figure 1(f). Thus, the LSPR scattering peak shows the spectra in the visible region of EM spectrum. Furthermore, the Figure 1(g, h) give relation between core size and LSPR wavelength of single and multi core-shell.
Figure 1: Calculated absorption efficiencies (a) Fe@Ag (b) Co@Ag (c) Ni@Ag and scattering efficiencies (d) Fe@Ag (e) Co@Ag (f) Ni@Ag and size dependent LSPR wavelength (g, h) for single core-shell nanospheres of magneto-plasmonics.
nanostructures for absorption (Figure 1g) and scattering (Figure 1h) to represent distinct LSPR tunability with considered core-shell thickness. Further analysed, there is an occurrence of two LSPR peaks in Figure 1; the first peak appears at lower wavelength and the second at higher wavelength which can be accounted towards the dipole and quadrupole resonant mode contribution in the LSPR spectra. Since, the interaction between incident energy with particles produces the external driving field which is not uniform. Therefore, dynamic polarization happens in the particle and second order LSPR such as quadrupole resonant mode will contribute to LSPR spectra.

3.1.2 MNP@Au Nanoparticles

The spectra of absorption and scattering efficiency have been calculated for gold (Au) coated on Fe, Co and Ni spherical nanostructures. The absorption peaks for Fe@Au core-shell nanospheres have been calculated from 10 to 60 nm size of core at wavelengths 541 nm of 30 nm shell thickness, 534 nm of 20 nm shell thickness, 532 nm of 20 nm shell thickness, and 521 nm, 517 nm, 507 nm of 10 nm shell thickness respectively shown in Figure 2(a). As the size of core increases with the combination of shell thickness, the peak position shifts to lower wavelength from 541 nm to 507 nm. Thus, LSPR absorption peaks are found in the visible region of EM spectrum. Figure 2(b) shows various peaks of Co@Au at wavelengths 541 nm for 10 and 30 nm, 534 nm for 20 and 20 nm, 532 nm for 30 and 20 nm, and 519 nm, 510 nm, 501 nm for 40 nm to 60 nm size of core with shell thickness 10 nm respectively. As the size of core increases with the combination of shell thickness, the peak position shifts from 541 nm to 507 nm. Thus, LSPR absorption peaks are found in the visible region of EM spectrum. For Ni@Au core-shell, we get various peaks at wavelengths 542 nm for 10 and 30 nm, 536 nm for 20 and 20 nm, 538 nm for 30 and 20 nm, and 525 nm, 516 nm, 505 nm for 40 nm to 60 nm size of the core with shell thickness 10 nm respectively. As the size of core increases with the combination of shell thickness, the peak position shifts from 542 nm to 505 nm as shown in Figure 2(c). Thus, LSPR absorption peaks are found in the visible region of EM spectrum. We get the scattering peaks for Fe@Au at wavelengths 636 nm, 636 nm, 636 nm, 630 nm, and 613 nm for 10 nm to 60 nm core sizes with shell thickness 60 nm to 10 nm respectively shown in Figure 2(d). As the thickness of shell increases for each considered core size, the peaks position shifts to the higher wavelength with greater efficiency. The scattering peaks are found in NIR of EM spectrum, and peaks are shifted from 636 nm to 613 nm with increase in core size. Thus, LSPR peaks get blue shifted for Fe@Au core-shell nanospheres. For Co@Au core-shell, the various peaks are found at wavelengths 636 nm, 636 nm, 636 nm, 629 nm, 614 nm for 10 nm to 60 nm core sizes with shell thickness 60 nm to 10 nm respectively. As the thickness of shell increases for each considered core size, the peak position shifts to higher wavelength with greater efficiency as shown in Figure 2(e). With the increase in core size, peaks shift from 636 nm to 614 nm and the spectra are found in NIR on EM spectrum. Thus, LSPR peaks get blue shifted for Co@Au core-shell nanospheres. Figure 2(f) shows various peaks of Ni@Au at wavelengths 636 nm, 636 nm, 636 nm, 623 nm, and 625 nm for 10 nm to 60 nm core sizes with decrease in shell thickness from 60 nm to 10 nm respectively. As the thickness of shell increases for each considered core size, the peak position shifts to higher wavelength with greater efficiency. The peak is shifting from 636 nm to 625 nm with an increase in core size and spectra are found in NIR region on EM spectrum. Thus, LSPR peaks get blue shifted for Ni@Au core-shell nanospheres. In addition, the Figure 2(g, h) give relation between core size and LSPR wavelength of single and multi core-shell nanostructures for absorption (Figure 2g) and scattering (Figure 2h) to represent distinct LSPR tunability with considered core-shell thickness.
Figure 2: Calculated absorption efficiencies (a) Fe@Au (b) Co@Au (c) Ni@Au and scattering efficiencies (d) Fe@Au (e) Co@Au (f) Ni@Au and size dependent LSPR wavelength (g, h) for single core-shell nanospheres of magneto-plasmonics.
3.2 Multi-layer Core-shell MPNPs

We coated magnetic nanoparticles of Fe, Co and Ni firstly with Ag and then with Au nanoparticles and vice versa. We used the optimized core-shell size of single core-shell nanospheres and fixed the outer shell with 10 nm thickness for multi core-shell. Thus, we calculated the absorption (left side) and scattering (right side) efficiencies of 80 nm total size of multiple core-shell nanospheres as shown in Figures 3 and 4.

3.2.1 MNP@Ag@Au Nanoparticles

Figure 3(a) shows the absorption spectra for Fe@Ag@Au multi core-shell nanospheres at wavelengths 541 nm, 541 nm, 521 nm, 504 nm, 401 nm and 444 nm for 10 nm to 60 nm core sizes with the combination of shell thickness respectively. For a 10 nm core size, the peak position shifts toward higher wavelength with greater absorption efficiency for 10 and 20 nm shell thickness, and towards lower wavelength with lesser absorption efficiency for 30 to 60 nm shell thickness respectively. On increasing the size of core above 20 nm with a combination of shell thickness, spectra are found with lesser absorption efficiency at lower wavelength. Absorption LSPR peaks are found between 541 nm and 401 nm on EM spectrum. For Co@Ag@Au multi core-shell MPNPs, various peaks are found at wavelengths 518 nm, 518 nm, 515 nm, 502 nm, 415 nm, 442 nm for 10 nm to 60 nm core size respectively as shown in Figure 3(b), with the combination of shell thickness. As the core size increases, the LSPR peaks shift from 518 nm to 415 nm. Various absorption efficiency spectra of Ni@Ag@Au nanospheres at wavelengths 518 nm, 521 nm, 519 nm, 502 nm, 422 nm, 457 nm for 10 nm to 60 nm core sizes with the combination of shell thickness respectively are shown in Figure 3(c). The LSPR peaks are found between 521 nm and 422 nm when the size of shell increases. Moreover, there is shifting of absorption peak towards the higher wavelength with peak difference of 100 – 150 nm due to the Au coating on MNP@Ag core-shell nanospheres. The scattering efficiency spectra for Fe@Ag@Au multi core-shell nanospheres at wavelengths 657 nm, 657 nm, 657 nm, 661 nm, 663 nm and 665 nm for 10 nm to 60 nm core sizes with shell thickness 60 nm to 10 nm respectively are shown in Figure 3(d). LSPR peaks shift towards higher wavelength when the core size is less than 30 nm (10-30nm), and for core size greater than 40 nm, the LSPR peaks also move towards higher wavelength with an increase in scattering efficiency. As the core size increases with decrease in the shell thickness, LSPR peaks shift between 665 nm and 657 nm and spectra are found in NIR region of EM spectrum. Figure 3(e) shows the LSPR spectra for Co@Ag@Au nanospheres at wavelengths 657 nm, 657 nm, 657 nm, 657 nm, 662 nm and 665 nm for the considered multi core-shells. For core size between 10-40 nm, LSPR peaks shift towards higher wavelength, and also shift towards higher wavelength for 50 nm and 60 nm with an increase in efficiency. LSPR peaks shift between 665 nm and 657 nm and spectra are found in NIR region of EM spectrum when core size increases with decrease in shell thickness. In case of Ni@Ag@Au, the scattering peaks are found at wavelengths 657 nm, 657 nm, 657 nm, 659 nm, 661 nm and 664 nm for 10 nm to 60 nm core sizes with shell thickness 60 nm to 10 nm respectively as shown in Figure 3(f). For core size less than 40 nm (10-40nm), LSPR peaks shift toward higher wavelength and with core size greater than 50 nm, LSPR peaks shift towards higher wavelength with an increase in efficiency. As the core size increases with decrease in the shell thickness, the LSPR peaks shift between 664 nm and 657 nm, and spectra are found in NIR region of EM spectrum. Moreover, there is shifting of scattering peaks towards the higher wavelength with peak difference of 75 - 80 nm due to the Au coating on MNP@Ag core-shell nanospheres. Further, the Figure 3(g, h) give relation between core size and LSPR wavelength of single and multi core-shell nanostructures for absorption (Figure 3g) and scattering (Figure 3h) to represent distinct LSPR tunability with considered core-shell thickness.
Figure 3: Calculated absorption efficiencies (a) Fe@Ag@Au (b) Co@Ag@Au (c) Ni@Ag@Au and scattering efficiencies (d) Fe@Ag@Au (e) Co@Ag@Au (f) Ni@Ag@Au and size dependent LSPR wavelength (g, h) for multiple core-shell nanospheres of magneto-plasmonics.
3.2.2 MNP@Au@Ag Nanoparticles

Absorption and scattering spectra have also been calculated for MNP@Au@Ag core-shell nanospheres. The absorption peaks for Fe@Au@Ag at wavelengths 487 nm, 471 nm, 496 nm, 376 nm, 439 nm and 656 nm for 10 nm to 60 nm core size with the combination of shell thickness respectively are as shown in Figure 4(a). On increasing the core size for different combinations of shell thickness, the spectra are found between 656 nm and 376 nm, in the NIR to UV region of EM spectrum. For Co@Au@Ag, the absorption peaks are found at wavelengths 490 nm, 462 nm, 493 nm, 382 nm, 442 nm and 430 nm for 10 nm to 60 nm core size with the combination of considered shell thickness respectively. As the core size increases for various combinations of shell thickness, the spectra are found between 493 nm and 382 nm wavelength with broadening peaks as shown in Figure 4(b). Hence, the absorption LSPR is found from visible to UV region of EM spectrum. Figure 4(c) shows the peaks for Ni@Au@Ag nanospheres at wavelengths 487 nm, 469 nm, 496 nm, 382 nm, 442 nm and 430 nm for the considered multi core-shell. The LSPR spectra are found between 496 nm and 382 nm with broadening of the peak when the core size increases along with the various values of shell thickness. Thus, the LSPR absorption peaks shift from visible to UV region of EM spectrum. Lastly, it has to be mentioned that due to Ag coating on MNP@Au core-shell nanospheres, broadening in scattering peaks have been found.

Scattering peaks for Fe@Au@Ag multiple core-shell are found at wavelengths 656 nm, 656 nm, 656 nm, 657 nm, 658 nm and 656 nm for 10 nm to 60 nm core size with shell thickness 60 nm to 10 nm respectively as shown in Figure 4(d). For core size less than 30 nm (10-30 nm), LSPR peaks shift towards higher wavelength, and for core size more than 40 nm, LSPR peaks shift towards higher wavelength with an increase in efficiency. As the core size increases with decrease in shell thickness, LSPR peaks are found in NIR region of EM spectrum. Figure 4(e) shows the various peaks for Co@Au@Ag nanospheres at wavelengths 654 nm, 654 nm, 654 nm, 655 nm, 656 nm and 654 nm for 10 nm to 60 nm core sizes with Au shell thickness 60 nm to 10 nm respectively and fixed outer Ag shell thickness of 10 nm. For core size less than 30 nm (10-30 nm), LSPR peaks shift towards higher wavelength, and for core size greater than 40nm, LSPR peaks shift towards higher wavelength and efficiency increases. With the increasing core size and decreasing shell thickness, LSPR peaks are found in NIR region of EM spectrum. For Ni@Au@Ag MPNPs, the scattering peaks are shown in Figure 4(f) at wavelengths 655 nm, 655 nm, 655 nm, 655 nm, 657 nm, and 654 nm for the considered multi core-shell thickness. For the core size less than 30nm (10-30nm), LSPR peaks shift towards higher wavelength, and for core size greater than 40nm, LSPR peaks shift towards higher wavelength with an increase in efficiency. On increasing core size with decrease in shell thickness, LSPR peaks shift between 654 nm and 657 nm and LSPR spectra is found in NIR region of EM spectrum. Moreover, with Ag coating on MNP@Au core-shell nanospheres broadening of peaks has been observed along with shifting of peaks towards higher wavelength with peak difference of 15 – 20 nm. Similarly, the Figure 4(g, h) give relation between core size and LSPR wavelength of single and multi core-shell nanostructures for absorption (Figure 4g) and scattering (Figure 4h) to represent distinct LSPR tunability with considered core-shell thickness.
Figure 4: Calculated absorption efficiencies (a) Fe@Au@Ag (b) Co@Au@Ag (c) Ni@Au@Ag and scattering efficiencies (d) Fe@Au@Ag (e) Co@Au@Ag (f) Ni@Au@Ag and size dependent LSPR wavelength (g, h) for multiple core-shell nanospheres of magneto-plasmonics.
4 Conclusion

We calculated the optical properties (absorption and scattering) of single and multi core-shell magneto-plasmonic nanoparticles (MPNPs) with Fe, Co and Ni as core materials and Ag or/and Au as shell materials with water as a surrounding medium to control the LSPR peaks. It was found that absorption and scattering LSPR peaks are in UV and visible region of EM for core-shell MNP@Ag nanospheres respectively. The absorption LSPR peaks are found in visible region and scattering LSPR peaks in NIR region along with blue shift for MNP@Au nanospheres and these peaks show spectra in visible and NIR region of EM spectrum with a shifting of absorption peaks from visible to UV region and scattering LSPR are found in NIR region with broadening peaks on EM spectrum for multi core-shells of MNP@Au@Ag nanospheres. It is found that shifting of optical peaks towards higher wavelengths depends on gold shell thickness whereas optical efficiencies depend on silver (as core or shell) thickness. It is also observed that double-layered MPNPs show larger LSPR peaks for scattering and absorption as compared to single core-shell MPNPs. Thus, it is concluded that the LSPR peak depends upon the thickness of the shell, size of core and material. Tuning and controlling the optical properties of MPNPs in the UV to NIR region of the electromagnetic spectrum may be useful for the applications of MNPs in optical imaging and cancer therapy.

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