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Design of engineered nanoparticles for biomedical applications by computational modeling

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Engineered nanoparticles exhibit superior physicochemical, antibacterial, optical, and sensing properties compared to their bulk counterparts, rendering them attractive for biomedical applications. However, given that nanoparticle properties are sensitive to their nanostructural characteristics and their chemical stability is largely affected by physiological conditions, nanoparticle behavior can be unpredictable *in vivo*, requiring careful surface modification to ensure biocompatibility, prevent rapid aggregation, and maintain functionality under biological environments. Therefore, understanding the mechanisms of nanoparticle formation and macroscopic behavior in physiological media is essential for the development of structure–property relationships and, their rational design for biomedical applications. Computational simulations provide insight into nanoscale phenomena and nanoparticle dynamics, expediting material discovery and innovation. This review provides an overview of the process design and characterization of metallic and metal oxide nanoparticles with an emphasis on atomistic and mesoscale simulations for their application in bionanomedicine.

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1 Introduction

Nanotechnology has been a growing field of research due to proven applications of novel nanomaterials in drug or gene delivery, bioimaging, theranostics, tissue engineering, and antimicrobial materials to name a few. In the scale range between 1 to 100 nm, nanoparticles (NPs) possess distinct magnetization, and superior catalytic, optical, and antibac-

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terial activity,⁹ different from bulk materials owing to their high surface-to-volume ratio.¹⁰

The widespread implementation of engineered nanomaterials is contingent on advancing efficient and sustainable manufacturing methods that enable their synthesis at industrial scale. For example, liquid-fed flame reactors¹¹ are distinguished for their ability to manufacture sophisticated functional nanomaterials at scale with diverse compositions and morphologies,¹² finding applications in gas sensing, bioimaging, and nanomedicine. These reactors have been used to manufacture gas sensors for selective detection of analytes,¹³ phosphors for cell bioimaging (e.g., SiO₂-coated Y₂O₃:Tb³⁺ (ref.



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14)), dental materials, 15 radioenhancers for nanotherapeutics, 16 food supplements, 17 and nanomaterials for theranostic applications, such as aggregated Nd3+-doped BiVO4 luminescent nanothermometers, 18 luminescent CeO2:Eu3+ for realtime H₂O₂ biosensing,¹⁹ plasmonic SiO₂-coated Au/Fe₂O₃ nanoaggregates for photothermal killing of human breast cancer cells20 and SnO2 breath sensors for lung cancer diagnosis. 21 Table 1 shows a list of metallic nanoparticles, as well as metal, ceramic, and mixed oxides, highlighting their key properties and potential market applications.

Despite remarkable advancements in the production of sophisticated materials through gas-phase processes, commercialization efforts have not mirrored the same pace. Notable exceptions include the successful commercialization of nanosilver (HeiQ), amine-functionalized carbon-coated cobalt nanomagnets (TurboBeads AG), and flame-made carbon-coated iron carbide nanomagnets that are polymer-decorated with affinity binders²² for magnetic blood purification (Hemotune AG). This delay in commercialization stems from the difficulty in controlling NP characteristics during scale-up, underscoring the need for bridging the gap between academic advancements and successful commercial applications. At the same time, the broad range of materials and compositions explored experimentally is not reflected in computational nanoscience and nano-biomedicine, which typically focuses on pristine metallic nanoparticles (most commonly gold) or a few metal oxides (Table 1). In addition, toxicological concerns must be addressed before integrating engineered NPs into biomedical devices.23 For instance, while platinum has been used in the manufacture of silicone breast implants, questions about its toxic effects remain unresolved. 24,25

Final product properties and their macroscopic manifestations of biological, environmental, and technological interest depend on NP size, morphology, crystallinity, state of agglomeration, and surface coating, which are largely determined by the process conditions during their synthesis. 26,27 As NP size decreases, the surface-to-volume ratio increases, enhancing electrical and thermal conductivity, while variations in NP composition, such as alloying, improves thermal stability compared to monometallic counterparts.²⁸ Superparamagnetism is also size-dependent, resulting in high magnetization only when NPs are exposed to a magnetic field, which disappears once the field is removed. The NP shape affects superparamagnetic properties. For example, cubic CoFe₂O₄ NPs exhibit lower coercivity than spherical ones.29 The composition and crystallite size in NPs play a critical role in determining magnetic properties. Metallic NPs (Fe, Ni, Co, icosahedral Au)³⁰ exhibit different magnetic properties from metal oxides (Fe₃O₄, CeO₂) or nanocomposites. ³¹ Additionally, larger crystallite sizes enhance ferrimagnetic properties of Fe₃O₄ NPs, while smaller sizes may transition from ferrimagnetic to superparamagnetic states, facilitating their use in biomedical imaging, 32,33 smart magnetic nanocarriers,³⁴ and hyperthermia therapy.³⁵

Nanoscale characteristics can significantly affect NP plasmonic and optical properties. Variation in size and length of Au nanorods can shift the local surface plasmonic resonance, 36 while bell-shaped Au nanostructures can enhance the local electromagnetic field.³⁷ Surface treatment can aid in controlling NP optical properties,³⁸ tuning their performance as imaging agents^{39,40} or in photothermal or photodynamic disease therapies.41

In drug delivery, nanoparticle size determines cellular uptake efficiency, 117 with NPs between 10-100 nm favoring targeted drug release. 118 The NP size, surface charge, and presence of ligands on NP surfaces influence the stability of colloidal suspensions and solubility, and hence their biological availability. 119 Large particles, for example, exhibit poor solubility, and reduced toxicity but have increased tendency to aggregate. 120 Furthermore, variations in the NP size, surface chemistry, and shape significantly affect their biodistribution and pharmacokinetics, thereby altering their toxicological profiles.121

Computer simulations can significantly aid in material optimization and commercialization of NPs by unravelling the mechanisms that govern nanoscale interactions during synthesis and by exploring NP reactivity with the environment. 122 These tools complement and accelerate experimental NP design¹²³ by exploring the effect of nanoparticle characteristics on their physicochemical properties and connecting them to performance factors for biomedical applications. A key challenge in nanomaterial modelling is their multiscale nature, ranging from 10 and 15 orders of magnitude in length and time. Quantum mechanics (QM), density functional theory (DFT), classic or reactive molecular dynamics simulations (MD), mesoscale and continuum models (Fig. 1), have been commonly used to predict NP reactivity and NP macroscopic behavior in biological media. 124 Depending on the computational model, different predictions can be made at multiple scales about NP stability, cellular interactions, and drug release kinetics. OM methods can predict Gibbs energies of solvation, chemisorption, and protein-ligand interactions. 125,126 MD and coarse-grained MD (CGMD) simulations can elucidate binding affinities, 127 protein conformational changes upon NP attachment, 128,129 ligand-protein interactions for functionalized nanomaterials, 130 drug release131 and NP interactions with lipid membrane proteins¹³² or other relevant biomolecules. 133 The effect of external electromagnetic fields on biologically relevant systems can be elucidated by MD simulations, which aid in the design of stimuli-responsive nanomaterials, 134 with applications in tissue engineering and wound healing 135 or radiofrequency cancer therapies. 136 Larger length and time scale models can be used to correlate NP characteristics with the performance in end-use applications. For example, the interaction of NPs with thermal 137,138 and electric field distributions 239 can be described by finite element method, allowing for the prediction of nanomaterial performance in sensing applications or photothermal therapies. Numerical simulations 140 and computational fluid dynamics141 can be used to link the agglomeration state of magnetic NPs with their performance in drug delivery, hyperthermia therapies, or as magnetic resonance agents.

 Table 1
 Properties and biomedical applications of nanoparticles

NPs	Biomedical application	NP properties
Ag	Antibacterial activity: Ag ⁺ ions released from oxidized Ag NPs cause cell membrane damage. Used in medical devices and textile protective articles ⁴²	NP size and oxidation, 43 NP shape 44 and functionalization 45 affects $\mathrm{Ag}^{^{+}}$ release
	Anticoagulant activity: drug-loaded Ag NPs release phenindione molecules to improve therapeutic coagulant activity ⁴⁶	Size influence drug effect as molar ratio determines optimal drug release
	Antidiabetic activity: Ag NPs can reduce blood glucose and increase insulin levels; ⁴⁷ used in surgical dressings to reduce infections and	Sub 100 nm Ag NPs should be used to optimize Ag ⁺ use in surgical dressings ⁴⁸
	promote healing in diabetic foot treatment ⁴⁸ <i>Bioimaging</i> : efficient in absorbing and scattering light, used for	Surface functionalization ⁵¹ or coatings ^{49,52} on Ag NPs induces
	cancer cell bioimaging ⁴⁹ and biomarkers tracking; ⁵⁰ Ag NP optical activity supports potential photothermal therapy ⁴⁹	important changes in optical activity
	<i>Drug delivery</i> : due to their stability and biocompatibility, ⁵³ Ag NPs can cross the blood–brain barrier and deliver anti-seizure drugs ⁵⁴	Coatings on Ag NPs determine surface charges and aggregation affecting NP cytotoxicity ⁵⁵
Au	Anticancer activity: Au NPs can inhibit the viability of cancer cells. ⁵⁶ The efficiency of anticancer drugs can be improved by conjugating in Au NPs surfaces ^{57–59}	Cellular uptake of drug loaded Au NPs depends on NP size, shape and surface properties ⁵⁸
	Antioxidant activity: surface chemistry of Au NPs provides protection against reactive oxygen species, ⁶⁰ finding applications in Alzheimer's disease treatment ⁶¹	Sub 20 nm Au NPs can inhibit free radicals showing superior antioxidant and hepatoprotective effects ⁶⁰
	Bioimaging: the high biocompatibility and surface plasmon properties of Au NPs allows for their use in imaging applications such as cancer diagnosis ⁶²	Optical activity of Au NPs is controlled by functionalization and shape modification $^{\rm 62}$
	Biosensors: Au nanorods are used as ultrasensitive sensors based only on surface plasmon resonance, which are capable of detecting trace amounts of analytes, such as proteins and toxins ⁶³	Optical activity can be tuned by changing NP dimensions ³⁶
	Drug delivery: nanocarrier for drug, peptide, protein and gene delivery with high biocompability ⁶⁴	Negative charge on Au NPs leads to easy conjugation for multiple fragments release ⁶⁴
	Nanotherapy: deoxyribonucleic acid (DNA) conjugation of Au NPs provides DNA targeting in infectious diseases or gene malformation ⁶⁵	Increased stability and cell uptake efficiency by easy conjugation with AuNPs^{65}
Cu/ CuO	Antibacterial activity: low-cost antibacterial agents can be produced with copper-based NPs due to their ability to produce free radicals, change cell membrane integrity, and inactivate enzymes ⁶⁶	Smaller sizes and higher reactivity for some morphologies of Cu NPs increases their cytotoxicity 67
	Biosensors: Cu-based NPs are cheap and susceptible sensors for pesticides ⁶⁸ and glucose ⁶⁹	Surface charge and size distribution of Cu NPs influences their interactions 68
Pt	Antibacterial activity: small (<20 nm) PtNP composites show high antibacterial activity against pathogens ⁷⁰ causing cell-membrane rupture induced by their negative zeta potential ⁷¹	Antibacterial properties depend on NP shape, size and charge. ⁷¹ Pt NPs (<3 nm) exhibit high cytotoxicity ⁷²
	Anticancer activity: Pt ions released from functionalized Pt NPs induce DNA damage inhibiting its replication in various cell lines of cancer. ⁷³ Pt NPs can potentially avoid side effects of Pt-based	Pt NPs show size dependent cytotoxicity in cancer treatments. ⁷⁵ Pt nanoflowers act as radioenhancers in radiation therapy ⁷⁶
	drugs while presenting similar reactivity during chemotherapy ⁷⁴ <i>Anti-inflammatory activity</i> : Pt NPs show scavenging activity against free radicals ⁷⁷ showing potential in therapies for diseases where oxidative stress is produced ⁷⁸	Versatile functionalization of Pt NPs can tune their antioxidant activity 77
	Antifungal activity: Pt-based NPs show remarkable antifungal activity against diverse phytopathogens ⁷⁹	Spherical and oval Pt NPs with sizes 10–50 nm possess antifungal properties ⁷⁹
	Bioimaging: the excellent magnetic imaging resonance and optical activity of Pt NPs have been exploited in cancer theranostics Drug delivery: multifunctional conjugated Pt NPs are able to deliver	Photosensitizers such as porphyrins on Pt NPs improve theranostic performance Sub 30 nm Pt NPs ⁸² and surface PEGylation ⁸¹ increases thera-
	small molecules, peptides, and antibodies in cancer treatment and diagnostics ⁸¹	peutic efficacy
	Neuroprotection: bare and conjugated Pt NPs can inhibit aggregation of toxic Aβ amyloid peptides in Alzheimer's disease by facilitating the binding of small molecules to the amyloid. 83 Biosynthesized Pt NPs possess antioxidant and neuroprotective activity, with potential	Functionalization of Pt NPs produces stronger binding affinity with A β which inhibits formation of toxic oligomers ⁸³
Ni/ NiO	application in Parkinson's disease monitoring ⁸⁴ and treatment ⁸⁵ Antibacterial activity: the interaction between NiO NPs and bacteria generates reactive oxygen species that induce lipid, protein, and DNA damage ⁸⁶	Smaller NiO NPs show greater antibacterial effect in infectious microorganisms $^{\rm 87}$
	Antifungal activity: the radical scavenging and enzyme inhibition produced from NiO NPs causes damage in bacterial and fungal strains ⁸⁸	The oxidative stress and intracellular release of Ni^{2^+} likely depend on NP size
	Antioxidant activity: moderate antioxidant activity is presented in Ni and Ni-starch NPs, which can be used to develop new therapy agents ⁸⁹	Surface conjugation is possible in Ni NPs by nanocapsulation with ${\rm starch}^{\rm 89}$
	<i>Biosensors</i> : laser-induced graphene/Ni nanoparticle electrode for enzyme-free glucose detection ⁹⁰	High surface area NPs allows glucose response with high sensitivity and stability 90

Table 1 (Contd.)

NPs	Biomedical application	NP properties
TiO ₂	Antibacterial activity: owing to their high chemical stability, low cost, and strong oxidation propensity, TiO ₂ NPs can be used as antibacterial agents in coatings for medical devices and clothing ⁹¹ or in orthopedic implants ⁹² to prevent infection	High concentration of carboxyl groups on ${\rm TiO_2}$ promotes stronger attachment to textiles conserving photobactericidal activity 91
	Anticancer activity: the superior photo-catalytic activity of anatase TiO ₂ inhibits tumor growth under heat ⁹³ or ultrasonic treatment ⁹⁴ in chemotherapy and phototherapy ⁹⁵	Surface conjugation of ${\rm TiO_2}$ NPs improves their performance in photothermal and photodynamic therapies 93
	Drug delivery: the low cost, reduced toxicity, and photoactivity of small (<30 nm) TiO ₂ are desired in the formulation of nanocarriers for cancer treatments ^{96,97}	The conjugation of Au and TiO $_2$ NPs forms nanocomposites with enhanced the rapeutical effects 97
	Theranostics: ${\rm TiO_2}$ -based NPs multi-functional activity (photothermal and specific cell targeting) has been used in drug resistant pancreatic tumor therapy and diagnosis 98	Conjugation of ${\rm TiO_2}$ NPs with ${\rm Gd}^{3+}$ forms nanocomposites which can be loaded with receptors and drugs 98
Fe ₃ O ₄	Anticancer activity: stability, biocompatibility, magnetic hyperthermia, and high specificity make iron oxide NPs appropriate for potential cancer treatments ⁹⁹	Size and morphology dependent toxicity exhibit by $\mathrm{Fe_3O_4~NPs^{100}}$
	Antidiabetic activity: diabetes-inducing enzyme, α -amylase, can be inhibited by Fe ₃ O ₄ NPs that reduce the levels of glucose uptake ¹⁰¹	Surface loading of drugs on ${\rm Fe_3O_4}$ NPs produce antidiabetic effect 101
	<i>Biosensors</i> : the superior magnetic activity of Fe ₃ O ₄ -based NPs has been exploited in biosensoring. ¹⁰² Different coatings can be used to tune magnetic activity and improve biosensor efficiency ¹⁰³	Sub 15 nm magnetic NPs can reach difficult access regions of the body leading to identification of circulating tumor cells ¹⁰²
	Drug delivery: engineered Fe ₃ O ₄ -based NPs have been used as bio- degradable nanocarriers with good tissue penetration, low toxicity, and magneto-mechanical activity ¹⁰⁴	The shape of ${\rm Fe_3O_4}$ NPs influence the biodistribution of NPs in tumor tissues showing higher anti tumor activity 105
	Tissue engineering: the magnetic Fe_3O_4 NPs have been used in tissue disease detection or tissue regeneration by cell therapy ¹⁰⁶ Theranostics: the ability of Fe_3O_4 NPs to interact with multiple biomolecules and their good magnetic resonance imaging performance make them potential theranostic agents for cancer and other pathologies ¹⁰⁷	High magnetic forces are obtained at optimized sizes and compositions of nanocarriers for tissue regeneration 106 High near-infrared absorption for theranostic use is observed for core-shell Fe $_3$ O $_4$ -CuS nanoflower composites 107
CeO ₂	Antibacterial activity: CeO ₂ NPs exhibit antibacterial activity in three ways: they can penetrate the membrane cell and interact with enzymes and proteins, they increase toxicity by changing the intracellular pH, or they generate reactive radicals affecting the membrane ¹⁰⁸	Antibacterial mechanisms depend on NP size: <30 nm cellular uptake and denaturation of proteins, <50 nm toxicity in cell wall <100 nm induction of oxidative stress ¹⁰⁸
	Anticancer activity: CeO ₂ -based NPs can interfere with enzymatic activity and cause mitochondria and DNA damage, specifically targeting cancer cells without affecting normal ones ¹⁰⁹	Reactive oxygen species levels can be regulated by ${\rm CeO_2}$ NPs involved in antitumor mechanisms 109
	Antidiabetic activity: the high antioxidant activity of CeO ₂ NPs can reduce the oxidative stress which causes complications in diabetes and increases insulin levels ¹¹⁰	Ce oxidation states and oxygen vacancies suggest that ${\rm CeO_2~NPs}$ surfaces influence antioxidant effects 111
	Antifungal activity: the multidrug-resistant fungal species is a growing concern for public health. CeO ₂ show promising antifungal activity ¹¹²	Fungicidal activity is likely due to oxidative damage of cell membrane 112 generated by ${\rm CeO_2}$ NPs surfaces
	Drug delivery: CeO ₂ -based NPs have good stability and biodegradability making them suitable to design effective pH-sensitive drug delivery systems in cancer or neurodegenerative disease treatment ¹¹³	Neutral and positive surface charges of ${\rm CeO_2~NPs}$ at different pH conditions improve their multifunctional chemotherapeutic activity 113
	Neuroprotection: the aggregation of α -syn proteins generates amyloid filaments commonly observed in neurodegenerative diseases. These amyloids may generate radical species that damage nerve cells. Hybrid CeO ₂ NPs induces surface regeneration and antioxidant activity, ¹¹⁴ reducing toxicity after bonding to α -syn ¹¹⁵	The Ce ⁴⁺ /Ce ³⁺ pair formed on the surface of core–shell CeO ₂ hybrid NPs shows antioxidant features while conserving superparamagnetic properties
	Tissue regeneration: polymeric composites of CeO ₂ NPs have Ce ⁴⁺ and Ce ³⁺ valence states that affect its interaction with proteins. This is important in controlling cell proliferation and adhesion in biomedical materials ¹¹⁶	Surface charges and hydrophobicity in ${\rm Ce}^{4+}$ and ${\rm Ce}^{3+}$ rich region determines cell–NP interactions 116

This review aims to provide an overview of most recent advances in the use of computational simulations of engineered NPs in biomedical applications. Section 2 focuses on atomistic simulations that are routinely employed to explore the effect of nanoparticle characteristics, such as structure, crystallinity, and surface characteristics, on their reactivity, oxidation, dissolution, and interactions within biological environments. Such nanoscale properties and interactions are critical

for optimizing drug delivery systems and approaches for developing treatments for neurodegenerative diseases. Section 3 discusses the application of mesoscale and continuum models in nano-biomedical research. These models link nanostructural characteristics of NPs with their macroscopic manifestations, including settling behavior in nanotoxicology studies and properties such as optical activity and dissolution, crucial in biomedical research. The review concludes with a Conclusions

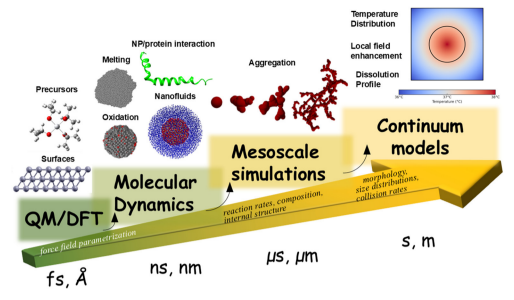


Fig. 1 Overview of time and length scales of density functional theory (DFT), molecular dynamics (MD), mesoscale, and continuum models for simulation of nanomaterials

and outlook section that highlights current challenges in computational nano-biomedicine and future directions that can contribute to the development of next-generation biomedical technologies.

2 Atomistic modeling of nanoparticles: density functional theory (DFT) and molecular dynamics (MD) simulations

Assessing NP chemical stability and reactivity in vivo is challenging, requiring detailed understanding across multiple scales. Atomistic models are extensively used to investigate these properties at the nanosecond and nanometer scale (Fig. 1). These are instrumental in describing early stages of NP formation, growth rates, as well as their physicochemical stability and reactivity. Quantum mechanics (QM) can fully describe the electronic structure of atoms and molecules, enabling the prediction of chemical properties (e.g., chemical reaction barriers, binding energies, and reaction pathways), which cannot be observed directly by experiments. 142 DFT is the most frequently applied QM approach, offering a balance between chemical accuracy and computational cost.143 The reliability of DFT predictions depends on the selection of appropriate functionals to describe electron interactions, which must be rigorously calibrated with reference data. 144,145

While DFT calculations focus on electronic structure which is useful for surface NP models, ¹⁴⁶ molecular dynamics (MD) simulate the dynamic behavior and long-term interactions of NPs with water and biological media, handling up to hundreds of thousands of atoms over microsecond timeframes (Fig. 1). Classical MD has been used traditionally to model NP stability

upon their formation and growth by different mechanisms (e.g., nucleation, 147 coalescence, 148 sintering 149), phase transitions (e.g., crystallization¹⁵⁰), and their interactions with polymers¹⁵¹ and proteins.¹⁵² Classical force fields, however, lack the ability to model chemical reactions due to the absence of charge transfer terms and their inability to describe bond formation and breaking. To overcome this limitation, reactive force fields (FF), such as ReaxFF¹⁵³ and the chargeoptimized many-body potential (COMB), 154 have been developed incorporating bond order principles from QM for the simulation of chemical reactions in a computationally feasible manner. In this section, in silico experiments of commonly studied metal and metal oxide NPs are discussed, highlighting the importance of their nanoscale characteristics (e.g., size, crystallinity) on properties such as oxidation and dissolution rates, and thermal performance. The section then extends to more complex systems, examining interactions within biological environments by atomistic or coarse-grained simulations, emphasizing on their effect on antibacterial activity, drugdelivery capabilities and disease treatment.

2.1 Nanoparticle oxidation

The oxidation state and ion release rate are crucial in determining the antimicrobial and antibacterial NP activity. ⁴³ The oxidation rate is affected by a range of factors, including temperature, nanomaterial purity, NP size, crystallinity, oxidation time, and oxygen concentration. ¹²² For example, nanosilver has remarkable antibacterial activity, finding applications in consumer products, including food containers, textiles, and cosmetics. ¹⁵⁵, ¹⁵⁶ Even though metallic silver is not soluble in water, ¹⁵⁷ when in the nanometer size range, it can be oxidized releasing silver ions from its surface. Such Ag⁺ ion release (leaching) in aquatic environments is associated with the tox-

icity of sub-10 nm nanosilver, 158 which has been threatened with labelling as a pesticide by the US Environmental Protection Agency. 159 Combined scanning tunneling microscopy and DFT calculations 160 revealed that oxidation of (110) Ag surface (Fig. 2a) takes place by dissociation of O2 and formation of O-Ag-O complex (Fig. 2b) that weakens the Ag-Ag bond and produces a vacancy (Fig. 2c) and a Ag adatom (Fig. 2d). The O-Ag-O complex diffuses to the closest hollow site producing a concerted motion (Fig. 2e), which induces a new vacancy (Fig. 2f). Oxidation of Ag NPs occurs by the formation of a core-shell structure, where a metallic Ag core is formed covered by a silver oxide layer. 161 The surface oxide layer predicted by reactive MD simulations thickens with decreasing nanoparticle size, consistent with ion-selective electrode experiments, 43 while sufficiently small (<2 nm) Ag NPs suffer internal oxidation.¹⁶¹ An Arrhenius-type relationship describes the oxidation rate constant, showing faster oxidation at higher temperatures and lower activation energies with decreasing nanosilver size (Fig. 2g). 161 Even though the NP size plays a key role in oxidation and thus antibacterial activity of nanosilver, NP shape also affects particle oxidation. For Ag NPs with similar surface area, NP geometries with higher surface fractions of (100) and (111) facets oxidize faster compared to nanostructures with large fraction of (110) facets that are structurally less stable and lose their crystallinity more easily. 162 The influence of facets reactivity is also observed by gas-solid interfacial energy calculations for Cu¹⁶³ showing that Cu₂O monolayers can form and grow on (110) Cu facets more easily than on (100) ones, in agreement with environmental transmission electron microscopy imaging.

Likewise, platinum (Pt) is subject to rigorous scrutiny due to their potential in biomedical applications. It is used as a catalyst in the production of silicone breast implants, raising toxicological concerns associated with the residual Pt that might enter the body, either through the intact shell or in the event of implant rupture.²⁵ Platinum NPs have shown varying levels of toxicity, with some studies indicating cytotoxic effects through mechanisms like oxidative stress, 164 DNA damage, 165 and cell cycle arrest, 166 while others demonstrate limited harm, depending on factors such as size,167 coating,168 and concentration.⁷⁷ Although FDA recognizes Pt as a safe material when in its zero-oxidation state, small Pt NPs (around 1 nm) exhibit greater toxicity upon oxidation, likely due to increased Pt2+ ion release that disrupts DNA structures. 169 The exact mechanisms behind this varied Pt toxicity remain unclear. MD simulations using the COMB3 force field have shown that higher oxygen coverage led to destabilization of the Pt NP surface atoms. 170 Larger Pt nanoparticles, however, were more stable due to their lower surface-to-volume ratio and lower oxygen adsorption energy, making them more resistant to oxidation.

The metallic surface oxide layer that forms upon oxidation is linked not only to nanoparticle toxicity, but also to reduced performance. For example, a 0.4 nm silver oxide layer reduces surface-enhanced Raman scattering (SERS) enhancement factors by three orders of magnitude compared to unoxidized nanosilver. 171 Similarly, oxidation of Co NPs results in the formation of a Co/CoO core-shell structure, which deteriorates the overall NP magnetization due to the antiferromagnetic CoO shell dominating over the ferromagnetic Co core. 172 These experimental findings have been corroborated by DFT calculations of Co oxidation revealing the formation of a nonmagnetic Co₃O₄ layer¹⁷³ that drastically reduces NP magnetization, rendering them unsuitable for cancer theranostics.

While pure metallic NPs, such as Ag, Fe, and Cu, are susceptible to oxidation, they can achieve improved biocompatibility through surface treatment, such as coating with inert shells or functional molecules, or by combining with other materials. Such multicomponent NPs not only exhibit significantly reduced oxidation and ion release, leading to lower tox-

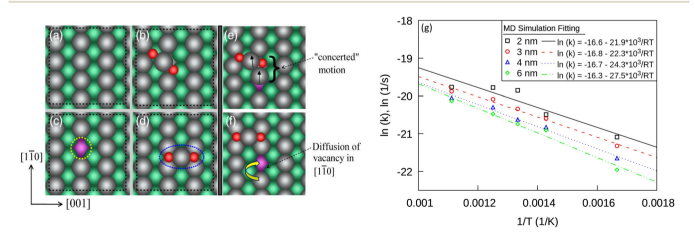


Fig. 2 Ag vacancy formation on a (110) facet: (a) pristine surface, (b) formation of O-Ag-O complex by oxygen dissociation, (c) formation of Ag vacancy, (d) Ag adatom stabilized by oxygen, (e) diffusion of O-Ag-O and Ag atom and (f) diffusion of vacancy. Ag atoms in the first, second and third layer are shown in gray, green, and purple, respectively; O atoms are shown in red (adapted from ref. 160 with permission from the American Physical Society, copyright 2017). (q) Arrhenius plots of the Ag NP oxidation determined by the shrinking core model at 600-900 K (adapted from ref. 161 with permission from the American Chemical Society, copyright 2023).

icity, but also demonstrate enhanced functional performance. For example, iron exhibits remarkable interest for magnetically directed drug delivery and hyperthermia applications, due to its higher magnetic strength compared to the more commonly used iron oxides. 174 However, iron oxidizes easily, is toxic, and tends to aggregate leading to thrombosis. 175 In contrast to pure Fe₁₃ clusters that form stable oxides (Fig. 3a) which reduce their magnetic moment, Au coatings prevent Fe oxidation (Fig. 3b), while maintaining the magnetic properties of the core Fe. 174 Similarly, substitutional introduction of Au into the corners and edges of Pt NPs inhibits oxide formation and Pt dissolution in aqueous solution, enhancing its stability. 176 In addition, Au-Co bimetallic NPs with L10 ordered structure exhibit higher magnetic anisotropy energy compared to core-shell or disordered (alloyed) ones, highlighting potential for magnetic NP hyperthermia applications.¹⁷⁷ Recently, ReaxFF MD simulations 178 revealed that even though

the crystal phase and metallic core are preserved during early stages of sintering of Ni/NiO and Cu/Cu2O metal/metal oxide core-shell NPs, the iron core in Fe/Fe₂O₃ NPs is fully oxidized due to higher oxygen mobility (Fig. 3c).

The effect of the local surface structure on NP physicochemical stability has also been demonstrated by DFT and electronic structure calculations of metal oxides, such as CeO₂ ¹⁷⁹ and TiO₂. ¹⁸⁰ Oxygen diffusion leads to disordered but partially stable CeO2 oxide surfaces, which are further stabilized by water adsorption. 179 Furthermore, water molecules preferentially bond to low coordinated atoms of the TiO2 surface, enhancing the overall NP stability. 180 Coordination analysis of surface Ti atoms indicated that smaller NPs exhibit more undercoordinated Ti (coordination number <6) and less fully coordinated Ti atoms, enhancing the overall NP affinity to water adsorbates (Fig. 4).

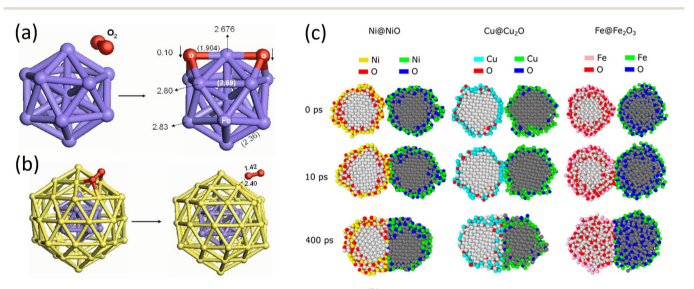


Fig. 3 Initial and optimized geometries for (a) $Fe_{13}-O_2$ and (b) $Fe_{13}@Au_{42}-O_2$. The bond lengths (Å) (in parentheses) and magnetic moments are given for $Fe_{13}-O_2$, while the bond length of O_2 and the distance between O_2-Au are given for $Fe_{13}@Au_{42}-O_2$. Purple: Fe, red: O, and yellow: Au. (c) Cross section snapshots of two coalescing core-shell NPs. Atoms are colored based on their initial position (core, surface, starting NP), with O atoms shown in red (left NP) and blue (right NP, adapted from ref. 178 with permission from the American Chemical Society, copyright 2021).

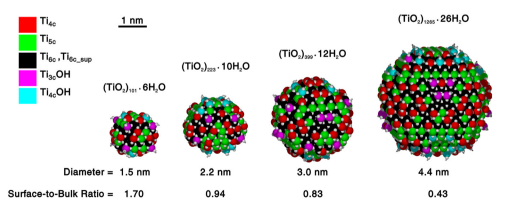


Fig. 4 DFT-optimized geometries of anatase TiO₂ NPs, corresponding to the most stable configurations after annealing at 300 (diameters of 1.5 and 2.2 nm) and 500 K (3 and 4.4 nm). Color code shows different coordination types for surface Ti atoms (adapted from ref. 180 with permission from the American Institute of Physics, copyright 2017).

2.2 Nanoparticle interactions in solutions: surface reactivity, dissolution, and thermal properties

Evaluation of nanoparticle properties in solutions, such as dissolution and surface reactivity, is crucial for biomedical applications due physiological constraints. For example, surface reactivity affects the properties of nanocarriers in drug delivery, while nanoparticle solubility (or dispersibility) affects their compatibility with biological systems. Poor solubility promotes aggregation which, in turn, limits NP interactions with target cells or tissues. However, controlled NP dissolution is often desirable for antibacterial activity. 181

Reactive MD using COMB3¹⁸² has shown that the degree of Pt NP oxidation affects its dissolution. Pt NPs with low fraction of oxygen adsorbates (<40%) on surface atoms release multiple highly charged Pt ions (Fig. 5a: red ions with q = +0.85e), while evenly sized Pt NPs with high oxygen coverage (>60%) do not dissolve. Water absorbs on Pt(111), (100), (110), (211), and (321) surfaces via the oxygen atom (Fig. 5b), ¹⁸³ and dissociates more easily on the (110) Pt surface than on (111), underscoring the significance of both the surface facet and coordination in Pt reactivity. The Pt(110) surface is more active than the (321)

facet, similar to Ag surfaces. 184 In anatase TiO2, water dissociation occurs at bridging oxygen sites, forming hydroxyl groups. 185 These sites on the edges and corners on the TiO2 NP facilitate faster water adsorption and desorption cycles compared to flat surfaces. Spontaneous dissociative adsorption of water on both anatase and rutile TiO2 nanosurfaces has also been observed, leading to formation of hydroxyl and hydrogen ions. 186 These computational insights are crucial for understanding water interactions with TiO2 surfaces in physiological media and highlight the potential for surface functionalization to enhance biocompatibility in drug delivery applications. CeO₂ NPs, studied by MD simulations and DFT calculations, exhibit increased reactivity at higher Ce3+/Ce4+ ratios and oxygen vacancies in CeO2, as highlighted by electrostatic potential maps (Fig. 5c-f). 187 This makes CeO₂ NPs susceptible to oxygen release in the presence of water, a property that can be exploited to enhance their effectiveness as nanocarriers in drug delivery.

The physicochemical properties of nanomaterials can also influence the thermal performance of nanofluids in medical applications. Enhanced thermal conductivity can optimize drug release, 188 aid cancer treatment with radiofrequency abla-

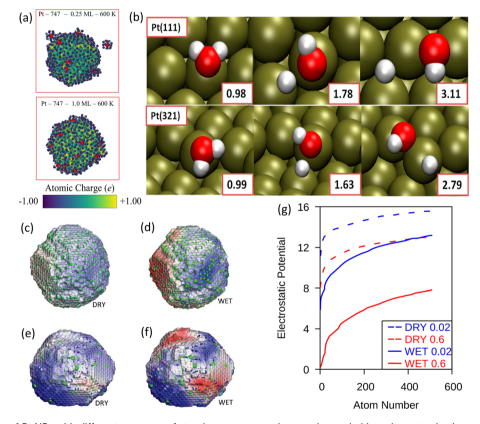


Fig. 5 (a) Dissolution of Pt NPs with different coverage of atomic oxygen monolayer, color-coded based on atomic charge (adapted from ref. 182 with permission from the Royal Society of Chemistry, copyright 2023). Atoms with a charge higher than 0.85e⁻ (red) dissociate from the monolayer at low oxygen coverage but not at higher levels. (b) Optimized geometries for the initial (left panel), transition (middle panel), and final states (right panel) of water dissociation on (111) (top) and (321) Pt facets (bottom), along with the cleaved O-H bond distances in Å (white: H, red: O, green: Pt) (Adapted from ref. 183 with permission from the American Chemical Society, copyright 2014). (c-f) Electrostatic potential maps of (c and e) dry and (d and f) wet CeO₂ NPs with Ce³⁺/Ce⁴⁺ ratio of (c and d) 0.6 and (e and f) 0.02. 187 Green spheres represent the positions of Ce³⁺ ions. (g) Calculated electrostatic potentials of the CeO₂ NPs of Fig. 5 (c-f) (adapted from ref. 187 with permission from the Royal Society of Chemistry, copyright 2013).

tion, 189 and improve flexibility of biological tissues during peristalsis facilitating endoscopy procedures. 190,191 Efficient heat transfer from nanomaterials to surrounding liquids is crucial in diagnostic tools, photothermal therapies, and drug delivery devices. 192,193 The thermophysical properties of nanofluids depend on NP characteristics such as size194 and morphology. 195 MD simulations have shown an increase in the specific heat capacity of Cu NP-water nanofluids for NP sizes of 1-10 nm, 194 approaching the bulk heat capacity for NPs larger than 4 nm. Addition of Cu and CuO NPs in water-based nanofluids increases the thermal conductivity by forming a heat-conductive channel around the immersed nanoparticles. 196 This is consistent with MD simulations 197 of Pt zig-zag nanochannels containing water and Cu NPs, where NPs demonstrated a preference to accumulate near the nanochannel walls, leading to increased density and thermal conductivity of the nanofluid, accompanied by a decrease in the temperature and molecular velocity of the nanofluid. Similar effects have been observed for CuO198 and Fe₃O₄ NPs. 199,200

The chemical environment within aqueous suspensions is also key in dictating the behavior of these engineered nanomaterials by altering their morphology and surface reactivity. The relationship between NP size and its impact on reactivity and toxicity is illustrated by MD simulations of 1-20 nm Ag NPs in water, ²⁰¹ revealing that smaller particles exhibit higher surface energy, making them less stable. Interestingly, within the 3-10 nm range, hybrid structures with high surface energies were identified, indicating that the reactivity and toxicity of small Ag NPs can change significantly even for slight size variations in current analytical methods. The roughness of such Ag NPs increases with increasing number of atoms, peaking at 8000 atoms before decreasing as the number of Ag atoms grow further, indicating an optimal particle size for NP roughness which affects NP interactions with ligands and biomolecules. Rough surfaces provide more stabilizing contacts compared to smooth surfaces, 202 enhancing binding efficiency and surface reactivity.

2.3 Antibacterial activity

Molecular dynamics and molecular docking simulations have been crucial in understanding the antibacterial activity of Ag NPs and their interactions with bacterial components. Such calculations have identified likely regions that nanosilver interacts most effectively with different bacteria, 203 and have elucidated the mechanism of Ag NPs cytotoxicity stemming from their interactions with proteins involved in apoptosis and oxidative stress generation.²⁰⁴ Furthermore, coarse-grained MD simulations demonstrated that when Ag nanoclusters (<1 nm) are coated with amphiphilic ligands, they exhibit superior antibacterial effects due to overcoming energy barriers more easily than Ag NPs larger than 2 nm. 205 IR spectra measurements supported by MD simulations²⁰⁶ indicated enhanced stability of Ag NP-nisin conjugates, due to stronger interactions between the N-terminal residues of nisin peptide and the NP core. In contrast, C-terminal residues are dominated by noncovalent interactions with the anionic membrane that destabilize the NP when it interacts with the cell membrane of bacteria.

Modification of Ag surface with polymers, such as poly(Nvinyl-2-pyrrolidone) (PVP) and chitosan, enhances the stability and antibacterial activity of Ag NPs by preventing aggregation. Tang et al.²⁰⁷ explored the adsorption of Ag NPs on activated carbon fibers using chitosan as binding agent, demonstrating excellent antibacterial activity. Fig. 6a-f shows coarse-grained MD simulations of the adsorption of Ag NPs on chitosancoated carbon fibers, forming small aggregates comprised of 1 to 5 NPs, leading to a decrease in their surface area. These clusters attach to chitosan chains, which transition from stretched to wrapped configurations around the cluster, leading to Ag NP deposition on the carbon, in accord with scanning electronic microscopy images (Fig. 6g and h). 207

2.4 Nanoparticle-based drug delivery

One of the game-changing applications of nanomaterials lies in targeted drug delivery. The discovery of new drug candidates is a slow and expensive process, taking years to identify a viable candidate.²⁰⁸ When using NP formulations, multiple variables must be taken into account, such as NP solubility (as discussed in section 2.2), drug loading capacity, release kinetics, functionalization, and interaction with biological targets. Computational approaches can accelerate the discovery of new drug candidates and improve the design of appropriate nanomaterials by predicting their properties and behavior in biological media. For example, McLean and Yarovsky²⁰⁹ have outlined the physicochemical²¹⁰ and biofouling^{211,212} properties of functionalized silica, illustrating its potential for biomedical applications. Computational screening of ligand-functionalized surfaces allowed for evaluation and selection of amorphous silica surfaces that resist protein adsorption, critical for developing efficient biomedical devices. 212 Both MD and experiments have revealed that silica NPs can translocate through giant unilamellar vesicles when exposed to high-frequency electromagnetic fields. This was also observed by MD and experiments, highlighting the impact of electromagnetic fields on enhancing membrane permeability, opening new possibilities for precise targeted drug delivery. 210

CGMD simulations have revealed that nanoparticle targeting of lipid membranes is influenced by various factors. Globular proteins adsorb on NPs in orientations that maximize contact between aminoacids and the NP surface.213 The permeation of fcc NPs in intestinal membranes varies with NP shape and surface charge. Nonpolar rod-shaped NPs cross the lipid membrane model more easily compared to spherical and disc-shaped ones, while polar NPs were unable to permeate the membrane regardless of their shape. 214 Additionally, CGMD simulations with the MARTINI model²¹⁵ have shown that the length and density of ligands in functionalized NPs with hydrophobic cores can tune their location in membranes. Hydrophobic NPs tend to occupy the raft domain of membranes while less hydrophobic NPs occupy the non-raft domain. Larger ligands can also affect the localization by hindering the exposure of NP's hydrophobic core.

Review

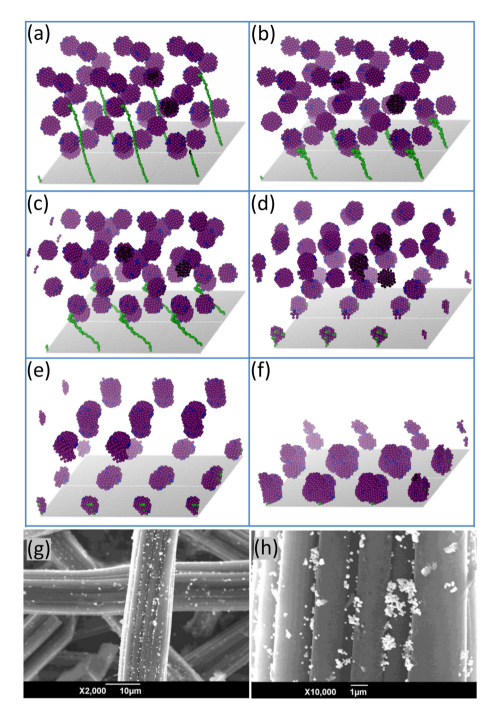


Fig. 6 (a-f) Adsorption of Ag NPs on chitosan-active carbon fibers (purple: non-charged Ag NPs, blue: negatively charged Ag NPs, green: chitosan monomers, silver: carbon surface) and (q and h) scanning microscopy images of Aq NPs supported on carbon (adapted from ref. 207 with permission from Elsevier, copyright 2017).

Biocompatibility and biodistribution are essential in nanocarrier applications. Natural rubber latex has been shown to improve Ag NP biocompatibility, increasing the release of antiinflammatory molecules, such as metronidazole (MET), 216 with potential in the treatment of bone or tissue infections. Simulations and experiments²¹⁷ revealed that MET-Ag NP interactions are favored over rubber-Ag NP ones, as indicated by the local softness in rubber latex (Fig. 7a) and MET mole-

cules (Fig. 7b), corroborated by in vitro experiments of MET release.

Surface coatings, such as poly-(ethylene glycol) (PEG), are commonly employed to increase NP stability, improve cellular uptake, reduce the toxicity of nanocarriers, and prevent unwanted interactions with biomolecules. MD, DFT, and simulated annealing Monte Carlo (MC) simulations have illustrated the adsorption of PEG molecules on Ag NPs, identifying

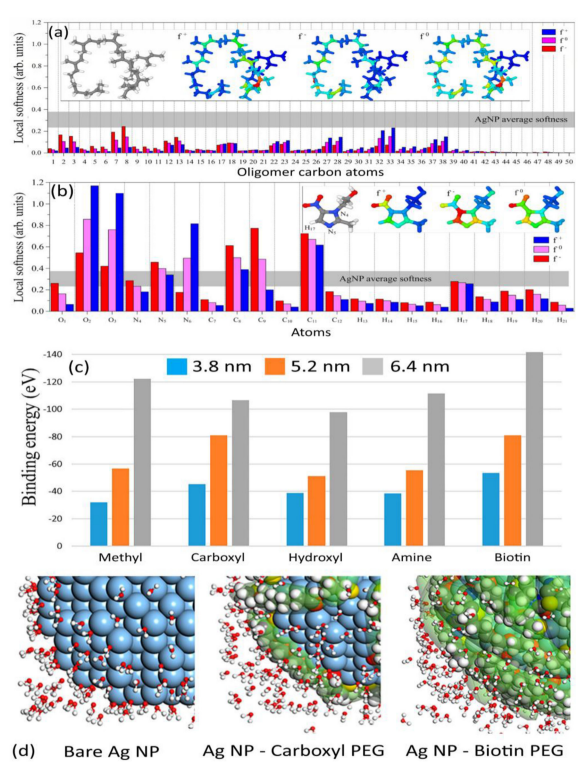


Fig. 7 Local softness as a function of atom type for (a) natural rubber latex and (b) MET molecule for the Ag NP functionalization (adapted from ref. 217 with permission from Springer Nature, copyright 2022). (c) Binding energy as a function of the number of functionalized PEG ligands on Ag NPs. 218 (d) Direction of oxygen in water molecules during interactions with a 6.4 nm Ag NP (blue: Ag, red: O, white: H, and green: carboxyl or biotin functionalized PEG ligands) (adapted from ref. 218 with permission from Taylor and Francis, copyright 2023).

methyl, hydroxyl, carboxyl, amine, and biotin functional groups as reactive centers. 218 Larger Ag NPs exhibited lower binding energies for all functional groups (Fig. 7c), but the carboxyl and biotin groups affected the directionality of water molecules (Fig. 7d), resulting in the most stable Ag NP-PEG structures. This potentially leads to water deprotonation and

local pH changes, impacting the cellular media. Similarly, PEG-coated ${\rm Fe_3O_4}$ NPs loaded with 5-fluorouracil demonstrated that NP coating and size can tune drug delivery. ²¹⁹

Multiscale modeling²²⁰ of PEGylated Au and Ag NPs revealed that increasing the density of PEG coating weakened NP-protein binding, while proteins exhibited preferential coating towards pristine Ag NPs rather than pristine Au. Simulations of immobilized proteins on Au NPs at various rotation angles (Fig. 8a-c) revealed that side chain interactions can guide the design of nanocarriers and nanosensors. Such multiscale approaches have been employed to optimize nanocarrier design targeting tumor cells, by estimating the binding energies between blood plasma or various dietary proteins and Ag NP surfaces. 221 The simulations have shown that biomolecule adsorption is more favorable on the (110) and (111) Ag NP facets rather than (100) ones, with aromatic residues showing strong binding on the (111) facet due to its electron density distribution, whereas linear molecules bind more strongly on the (110) facets (Fig. 9a). Such efficient screening of biomolecule interactions with different NP surfaces can aid in the rational design of nanocarriers. Similar computational approaches²²² have shown that TiO₂ NPs favored hydrophilic interactions over affinity to protein residues, while moderate interactions of Au NPs with gelatin²²³ and pectin, ²²⁴ primarily through physisorption, showcased their potential as drug carriers for curcumin, an antioxidant compound.

Super-paramagnetic Fe₃O₄ NPs have emerged as promising drug delivery agents due to their non-toxic nature, magnetic resonant activity, biocompatibility, and cost-effectiveness. 225 Their ability to maintain magnetic properties when complexed with polymers like chitosan enhances their potential in theranostic applications.²²⁶ Chitosan's positive charge density enables strong interaction with negatively charged drug molecules, making it an ideal carrier. MD simulations²²⁷ have revealed strong affinity of chitosan with the Fe₃O₄(111) surface, compared to other facets, with hydrogen and nitrogen from chitosan's amino group showing higher interaction probability with oxygen atoms on the (111) Fe₃O₄ facet. Furthermore, experiments and MD simulations of the adsorption of bovine serum albumin (BSA) protein on Fe₃O₄ NPs revealed the formation of multiple protein monolayers on the NP surface, with BSA protein spreading over the Fe₃O₄ NP surface and eventually relaxing into a compact configuration (Fig. 9b). 228 This strong protein adsorption, confirmed by transmission electron microscopy and UV-vis measurements, suggests BSA-coated Fe₃O₄ NPs as highly resistant drug delivery vehicles.

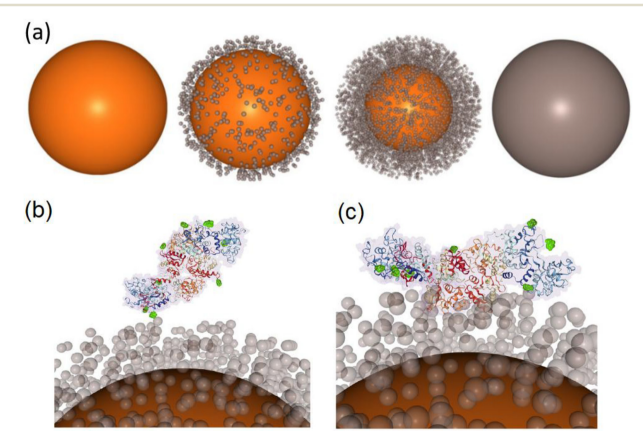


Fig. 8 (a) CG structures of pristine and PEGylated Au NPs. CG structures of docked protein apo-human serum transferrin on PEGylated Au NPs with rotational coordinates (b) $\varphi = 180^{\circ}$, $\theta = 90^{\circ}$ and (c) $\varphi = 200^{\circ}$, $\theta = 25^{\circ}$ (adapted from ref. 220 with permission from the Royal Society of Chemistry, copyright 2023).

Nanoscale

(a) spreading (t=3ns) relaxation (t=9ns)

Trp134
Trp213
Tr

Fig. 9 (a) Adsorption energies (in k_BT) for biomolecules on (100) (red), (110) (blue), and (111) (green) Ag surfaces (adapted from ref. 221 with permission from the American Chemical Society, copyright 2022). (b) Conformational changes of BSA adsorption on a Fe₃O₄ NP (yellow, adapted from ref. 228 with permission from the Royal Society of Chemistry, copyright 2016). (c) Binding energy of TMZ on Fe₃O₄ NPs as a function of NP size.²²⁹ (d) TMZ attains perpendicular orientation on 1.2 (left), 4.2 (center), and 6.5 nm (right) Fe₃O₄ NPs (adapted from ref. 229 with permission from Elsevier, copyright 2019).

Nanoparticles offer targeted delivery solutions for various chemotherapeutic agents, enhancing efficacy while minimizing side effects. For example, functionalized NPs with optimized size and charge can improve brain delivery of Temozolomide (TMZ), which exhibits poor permeability in the brain. DFT/MD studies²²⁹ have shown that Fe₃O₄ NPs smaller than 4 nm enhance TMZ permeability through chemisorption (Fig. 9c) and potential modification of TMZ's charge distribution, while for larger NPs chemisorption, physisorption, or hydrogen binding may occur. Chemisorption can result in modifications in the molecular structure of the drug molecule, whereas physisorption and hydrogen bonding leave the TMZ molecule intact, preserving its physical properties upon its release from the nanocarrier. The Fe₃O₄ NP loading capacity also varies with NP size, favoring a perpendicular TMZ configuration on the NP surface (Fig. 9d). Similarly, Doxorubicin (DOX), another widely used chemotherapeutic agent, which poses challenges due to its cardiotoxic side effects benefits from the encapsulation of TiO₂ nanotubes, that facilitate DOX aggregation within the nanotube as confirmed by DFT/MD simulations, 230 thereby reducing interaction with the cardiac

Nanoparticle Diameter (nm)

tissue. Furthermore, Phenindione, an anticoagulant with severe side effects, can be effectively incorporated into galactose-loaded Ag NPs, promoting slow release and slowing down blood coagulation, as demonstrated by DFT calculations of galactose-phenindione interactions. ⁴⁶ Similar *in silico* studies have been reviewed by Ashwini *et al.*²³¹

In antiviral and antibacterial therapies, NP-based approaches can mitigate side effects while enhancing therapeutic efficiency. Morad *et al.*²³² explored the adsorption of COVID-19 therapeutic agents, hydroxychloroquine and chloroquine, on various metallic NPs (Ag, Au, Ag/Au, and Pt NPs) by QM and MD. The hydroxyl group in hydroxychloroquine exhibited the strongest binding on Pt. Increasing Ag NP size from 1.6 to 4.6 nm led to weaker interactions with both drugs. Among metal clusters of 147 atoms and different compositions, affinity followed the trend: Ag NP < AuAg NP < Au NP < Pt NP.

2.5 NP-based computer simulations for neurodegenerative disease treatment

As these advancements in understanding NP behavior in biological systems continue, there is increasing attention towards

specialized medical challenges, such as treatment of neurodegenerative diseases. Alzheimer's (AD) and Parkinson's disease (PD) share features including accumulation of proteins in cells (amyloids), abnormal metal ions concentration in cerebral media, and high oxidative stress levels in the brain.²³³ Although a handful of drugs approved by the U.S. Food and Drug Administration are available for clinical use, their therapeutic benefits are short-lasting and unable to halt disease progression.²³⁴ The urgency to develop more effective treatments against neurological disorders has led researchers to explore multitarget strategies including nanomaterials. Computational simulations have proven essential in understanding complex interactions between biomolecules and nanocarriers, helping optimize drug design. However, most computational research has centered on molecular drug design, 235-237 with less emphasis placed on inorganic NPs, despite their potential in drug distribution.²³⁸ A major challenge in AD and PD drug design is ensuring that drug candidates cross the blood-brain barrier (BBB) while designing nanocarriers that avoid undesirable interactions during this BBB crossing.

One of the main hallmarks of AD is the aggregation of unfolded amyloid-beta (Aβ) monomers into fibrils that interact with NPs. Tavanti et al. 239 explored by MD the interaction between Aβ 40 monomers and citrate-capped Au NPs in water. Contact probability analysis between the functionalized Au NP and 1-3 Aβ monomers (Fig. 10a, top), revealed two binding Aβ sites, 13HHQK16 and 24VGSNKGAI31, with more binding sites

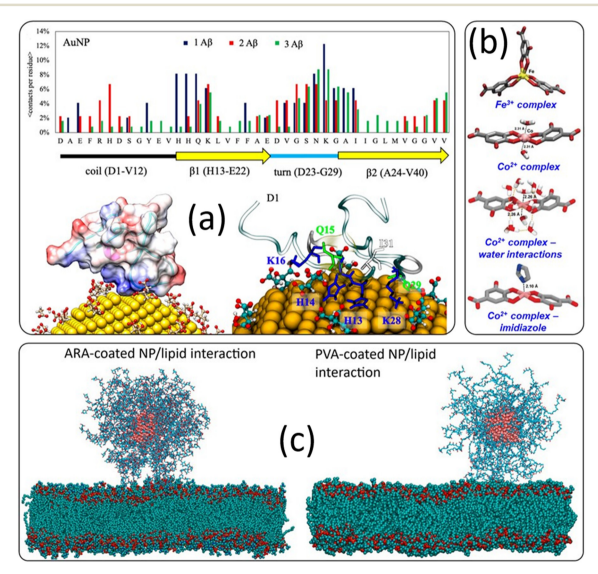


Fig. 10 (a) Contact probability between AuNP and A β (1–40) residues for three different amyloid concentrations (top); Au NP-A β complex interacting with A β , showing the surface potential map of A β (bottom-left) and amino acid residues involved during A β attachment to the Au NP surface (bottom-right). Negatively charged residues are shown in blue (adapted from ref. 239 with permission from MDPI, copyright 2021). (b) Structures of Fe (yellow) and Co (pink) complexes with tannic acid in the presence of water molecules and imidazole ring of histidine (adapted from ref. 241 with permission from the Royal Society of Chemistry, copyright 2019). (c) Interaction between lipid layer model and Fe₃O₄ NP coating with ARA and PVA (adapted from ref. 243 with permission from Elsevier, copyright 2020).

observed as the number of AB monomers increased, including a C-terminal contact site. Positively charged Aß regions (K28, H13, H14, and K16) interact with negatively charged citrate ligands (Fig. 10a, bottom left), while other residues (Q15, Q29, and I31) bind to Au atoms via van der Waals forces (Fig. 10a, bottom right). These interactions disrupted AB aggregation, highlighting the potential role of Au NPs in altering AD-related protein aggregation, with similar results seen in MD studies relevant to PD.²⁴⁰ Zhang et al.²⁴¹ synthetized Au NPs coated with metal-phenolic networks, with inhibitory activity towards amyloid formation. Polyphenolic networks between tannic acid and different metal ions (Al3+, Fe3+, Co2+, Ni2+, Cu2+ and Zn²⁺) were explored, with cobalt-tannic acid complexes exhibiting the highest inhibitory activity. MD simulations revealed Co complexes with uncoordinated axial positions allowing them to interact with water molecules and amyloid amino acids (Fig. 10b, top). In contrast, iron complexes formed stable octahedral structures, reducing solvent exposure. QM calculations showed that while water interactions with Co centers were unfavorable, histidine and methionine interact strongly with Co (Fig. 10b, bottom), potentially reducing oxidative stress and preventing Aβ aggregation commonly associated with AD.

To overcome BBB permeability restrictions, non-invasive techniques using magnetic fields have been investigated. Magnetic Fe₃O₄ NPs have emerged as effective contrast agents for magnetic resonance imaging (MRI) for AD or epilepsy diagnosis due to aggregation of these NPs in the brain by the high magnetic activity of the neural network in this zone. 242,243 Lazaratos et al. 243 performed MD simulations and experiments to examine the interactions between lipid bilayers and magnetic Fe₃O₄ NPs coated with polyvinyl alcohol (PVA) and polyarabic acid (ARA) (Fig. 10c). Stronger interactions and shorter Fe₃O₄ NP-bilayer distances were observed for the PVA-coated NP compared to ARA-coated ones. The potential of Au-coated Fe₃O₄ NPs to cross the BBB has also been explored by MD under magnetic fields.²⁴⁴ The Au/Fe₃O₄ nanocomposites could temporarily open gaps in the membrane, allowing passage and reversible deformation. By adjusting the magnetic field strength and pattern, the BBB crossing time could be precisely controlled, offering significant potential for NP-based treatments targeting the brain.

Parkinson's disease is characterized by dopamine deficiency, leading to loss of coordination and impaired movement. Simulations of Fe₃O₄ NPs for dopamine replacement therapy²⁴⁵ revealed that as the number of dopamine molecules on uncoated Fe₃O₄ NPs increases, their binding energy increases, though the rate of energy change slows down as more molecules are added. Larger PEG-coated Fe₃O₄ NPs exhibited even higher binding energies with dopamine compared to uncoated NPs, due to the hydrodynamic size increase in the presence of PEG, without sacrificing stability. Furthermore, albumin-coated Fe₃O₄ NPs enhance BBB permeability. Increasing the dopamine-to-albumin ratio at constant PEG led to higher binding energies, indicating stronger ligand-Fe₃O₄ NP interactions. However, increasing PEG for a given amount of dopamine reduced NP stability. These findings suggest that adjusting the ratios of NP coating, dopamine, and albumin offers a potential strategy to optimize NP size and stability for PD treatment. CeO_2 NPs also show potential to inhibit α -syn amyloid aggregate formation, critical in neurodegenerative diseases. Molecular docking and MD simulations revealed strong interactions between CeO_2 NPs and α -syn fibrils. These interactions induced conformational changes in amyloid structure when in proximity to CeO_2 NPs, such as large spacing among neighboring protein chains, leading to partial disaggregation of α -syn, consistent with spectroscopy measurements. 115

2.6 Other biomedical applications

The physicochemical properties of NPs change upon interaction with biofluids, as proteins adsorb to form a protein the NP's biological corona that affects response. Computational simulations are instrumental in predicting how these interactions affect NP stability, bioactivity, and the formation of the protein corona. For example, multiscale simulations have been employed to examine the formation of a protein corona on 3.2 and 10 nm Ag NPs using the antimicrobial peptide ovispirin-1.²⁴⁶ Hydrophilic interactions between the NP surface and certain peptide residues were key to the initial adsorption, with water solvation also contributing to the protein anchoring. An adsorption-desorption-readsorption cycle was observed for a 3.2 nm Ag NP, indicating an unstable attachment, while the larger 10 nm Ag NP showed more stable peptide anchoring due to its larger surface area, allowing peptide diffusion and stronger adsorption (Fig. 11a).

Power et al.²⁴⁷ developed a coarse-grain multiscale model to rank the adsorption affinity of proteins on Au NPs accounting for preferred orientations and the influence of NP size and shape. Rouse and Lobaskin²⁴⁸ developed a hard-sphere model to mimic protein corona formation on spherical and cylindrical NPs, finding that NP size and geometry affect protein packing and corona composition. Kinetic MC simulations confirmed these results, predicting different protein coverages depending on NP size and shape. Specific surface interactions with some adsorbates can give rise to more complex phenomena, such as enhanced Raman effect. For example, the interaction of PVA with Ag NPs, investigated by DFT,249 revealed that the hydroxyl oxygen atoms of PVA attach to a 55-atom Ag cluster, without significantly altering the NP surface. Negative binding energies indicated a thermodynamically favorable interaction between the Ag NP surface and PVA, consistent with surface-enhanced Raman scattering experiments.

Even though the biocompatibility of Au NPs makes them suitable for theranostics, their aggregation in physiological media must be prevented to preserve cellular uptake and low toxicity. Canepa *et al.* Explored this issue by CGMD simulations of ligand-functionalized 2–4 nm Au NPs interacting with a bilayer that mimics neuronal plasma membrane in physiological salt solution. Fig. 11b shows the spontaneous adsorption of a single functionalized Au NP onto the lipid, altering lipid–liquid phase separation. Larger Au NPs (4 nm) promote the formation of stable aggregates (Fig. 11b) by indu-

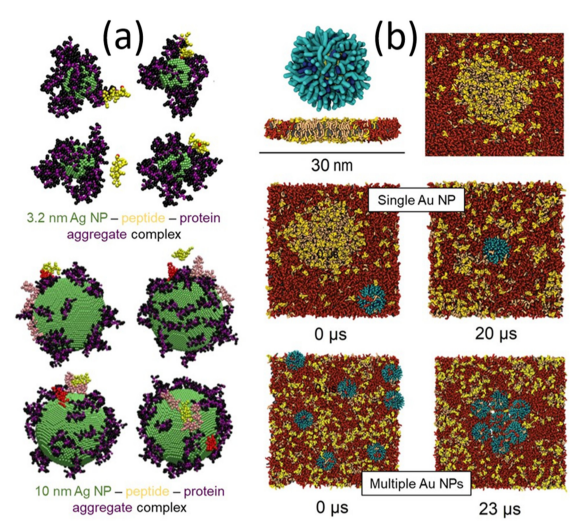


Fig. 11 (a) The adsorption—desorption—readsorption cycle for a 3.2 (top) and a 10 nm (bottom) Ag NP (adapted from ref. 246 with permission from the American Chemical Society, copyright 2022). (b) Coarse-grained representations of NP and lipid bilayer model (top and side views), where hydrophilic ligands are shown in cyan, hydrophobic ligands in blue, glycosphingolipids (gangliosides) in yellow, 1,2-dilinoleoyl-sn-glycero-3-phosphocholine in brown, sphingomyelin in pink and cholesterol in gray. The lipid—liquid phase separation when a single 4 nm Au NP is embedded in the bilayer is compared oncean aggregate is formed by clustering of multiple Au NPs (adapted from ref. 252 with permission from the Royal Society of Chemistry, copyright 2020).

cing perturbations in the lipid bilayer, while smaller ones (2 nm) form transient dimers. These results are consistent with CGMD simulations of fullerene aggregation in lipid membranes, showing that increasing fullerene size promotes aggregation. CGMD simulations fullerene size promotes aggregation. CGMD simulations fullerene size promotes aggregation. Showed that short-range charge-charge interactions (ion bridging), lipid depletion, and membrane curvature drive aggregation of adsorbed NPs on membranes, confirmed by cryo-electron microscopy experiments. Recently, Ahmed et al. Studied the assembly of 5–50 nm Au NPs with bacteriophages by the united atom approach, revealing that these composite nanostructures can minimize the formation of biomolecular corona, maintaining the targeting ability in the biological fluids, which is desirable in sensing applications.

The nonmagnetic nature of graphene has drawn interest in biomedical imaging, as the microstrain effects may induced by NPs may confer magnetic properties on graphene-based nanostructures. Idisi et al. 256 explored this effect on the magnetic properties of graphene oxide nanosheets doped with Fe-O from Fe₃O₄ NPs via spin polarizable DFT calculations. The negative formation energies obtained for different Fe-O dopant configurations incorporated in the graphene nanosheet proved the stability of these structures, with significant magnetization enhancement when multiple Fe-O sites are present. Spin density distribution and density of states analysis suggested charge transfer leading to interaction among C, Fe, and O atoms. This charge transfer, along with the NP size, likely explains the origin of the enhanced magnetization observed in experiments. Thomas et al.257 explored the structure of resilin-like peptides on graphene by experiments and MD simulations. Such materials exhibit enhanced elastomeric properties and can create artificial muscles or can be used in

Table 2 Summary of simulation studies of NP-macromolecule interactions relevant for biomedical applications. The computational models are described in terms of the force fields for MD and CGMD, and the basis sets and pseudopotentials in DFT

NP	NP characteristics	Computational model	Biomedical relevance	Ref.
Ag	Ag NPs (<3 nm), coated with adamantanethiolates	CGMD: MARTINI FF and Morse potential for water	Interaction between Ag NPs and multidrug-resistant bacteria membranes	205
	10 nm Ag NP	MD: GROMOS96 FF	Efficiency of the antimicrobial peptide nisin upon assembly at Ag NP surface	206
	Ag NPs, chitosan chains coating	CGMD: LJ-Coulombic potentials, implicit solvent model	Mechanism of Ag NP loading on chitosan chains for fabrication of antibacterial materials	207
	PVP 4.5 nm Ag NP	MD: G53a6 FF with DFT and LJ potential	Polymer-capped NPs with tunable properties	263
	2 nm Ag NP functionalized with MET	MD: AMBER FF, DFT: PBE/6-31G (d,p)	MET and Ag NPs supported on biocompatible rubber latex for drug delivery applications	217
	PEG-coated 1–3.2 nm Ag NPs	MD: consistent-valence FF, DFT: PW91, MC	NP size and PEG functionalization effect in Ag NP stability to supress NP aggregation	218
	Ag surfaces, 29.2 and 43 nm Ag NPs	CGMD: INTERFACE and CHARMM36 FFs, UA potential	Evaluation of hydrophobicity and lipophilicity of Ag NPs to predict their toxicity as nanocarriers	221
	3.2 and 10 nm Ag NPs	CGMD: Martini and CHARMM36 FFs	Size and hydrophilicity effects of Ag NPs on peptide corona formation	246
	PVA coated Ag nanocluster (<1 nm) 2–10 nm thiolate-coated Ag NPs	DFT: PW91 MD: ReaxFF	PVA binding on Ag NP surfaces for nanofiber fabrication Analysis of thiolates adsorption on Ag NPs (111) surfaces for tuned antibacterial properties	249 264
	$25 \times 25 \text{ Å}^2$ (111) Ag slabs surrounded by water and ligands	MD: AgP-CHARMM and CHARMM36m FFs, TIP3P model	Thermodynamics of the adsorption of different capping agents used for surface stabilization of Ag NPs	265
Cu	DNA and small Cu NPs (<1 nm) composites	DFT: B3LYP-PM6, MD: parmbsc1 FF	Size effect of Cu NPs in DNA binding and structure for their use as anti-fungal agents	266
Au	Bare 4 nm Au NPs	CGMD: MARTINI FF	Adhesion of Au NPs on mixed zwitterionic/anionic lipid membranes relevant in biomedical applications	124
	2 nm Au NPs functionalized with thiol ligands	CGMD: MARTINI FF	Adsorption of Au NPs on mixed lipid bilayers to understand their fate in biological fluids	267
	2 nm alkylamines functionalized Au NPs	MC/MD: CHARMM36m FF, CGMD: POL/BMW-MARTINI FF	Interaction between Au NPs and lipid bilayer membranes	268
	2 nm Au NPs functionalized with PEGylated ligands	MD: OPLS-AA FF, LJ and RB potentials	Solvation of PEGylated Au NPs in water which could inhibit corona protein formation	269
	10 atoms cluster to mimic Au NP surfaces	DFT: B3LYP/LanL2DZ pseudopotential	Determination of optimal NP size for high curcumin drug delivery yield	223
	10 atom cluster bonded to pectin chains	DFT: B3LYP/LanL2DZ pseudopotential	Electronic structure and adsorption study between Au NPs and pectin biopolymer to unveil drug delivery activity of coated Au NPs	224
	4 nm citrate-capped Au NPs with Aβ amyloids attached	MD: GROMOS 54a7 FF	Folding mechanism of Aβ amyloids on Au NPs surfaces to understand formation of toxic plaques in AD	239
	12 nm citrate-capped Au NPs bonded with α-syn protein	MD: CHARMM FF	Binding of α-synuclein with coated Au NPs associated with Parkinson's disease treatments	240
	Coated Au NPs (<50 nm) in water	DFT: B3LYP/6-311G, MD: COMPASS FF	Mechanistic insights of the inhibition of A β (1–40) amyloids with cobalt–tannic coated Au NPs	241
	2–4 nm coated Au NPs in multidomain bilayer model	CGMD: MARTINI FF	Effect of amphiphilic and negatively charged Au NPs on neuronal plasma membrane for sensing applications	252
	31.5 × 31.5 nm ² surface for Au nanorod with placed DNA receptors	CGMD: oxDNA2 model	Effect of DNA receptor density on DNA-modified Au nanorods for bioimaging and drug delivery purposes	270
	Au(100) nanosurfaces	DFT: B3LYP/6-31G, MD: CHARMM FF, TIP3P model	Binding affinities of capping molecules for Au nanorods growth with multiple biomedical applications	271
	Au(111) nanosurfaces	MD: CHARMM22 FF, TIP3P model	Transition of peptide structure on Au NP by variation of aminoacid fragment affects optical properties	272
TiO_2	DOX solution confined inside 2.5 nm TiO_2 nanotubes walls	DFT: B3LYP/6-311+G, MD: OPLS-AA and SPc FFs	TiO ₂ nanotubes as controllable drug-delivery systems of anticancer DOX compound	230
	2.2 nm TiO ₂ and 50 methoxy-PEG polymer chains	DFT: B3LYP, MD: MA and CHARMM36 FFS, CGMD: MARTINI 2 FF	Cell permeation efficiency estimation of TiO ₂ coated with PEG-based chains	261
	Polymorphs TiO ₂ slabs	MD: CHARMM36, LJ potential, TIP3P model	Adhesion of Arg-Gly-Asp on ${\rm TiO_2}$ surfaces for osseointegration applications	273
	2.2 nm anatase TiO ₂ NP with DOX and hydrophobic ligands	MD: GAFF2 and modified MA FFs, TIP3P model	pH-triggered release of DOX from functionalized TiO ₂ NPs to assess drug loading capacity	274
	PEGylated 2.2 nm anatase TiO ₂ NP with folic acid molecules	MD: CHARMM36 and CGenFF	Protonation state and folic acid density effect on the interaction of ${\rm TiO_2}$ NP-based nanodevices with folate	275
	TiO ₂ nanosurfaces, 4 nm Anatase NP	CGMD: Slipids FF, TIP3P model	receptors from tumor cells Adsorption of lipids near TiO ₂ NPs and nanosurfaces relevant to assess their toxicity	276

Review Nanoscale

Table 2 (Contd.)

NP	NP characteristics	Computational model	Biomedical relevance	Ref.
Fe_3O_4	PEGylated and bare 1–7 nm Fe ₃ O ₄ NPs	DFT: PW91, MC/MD: pcff and COMPASS FFs	Coating (PEGylation) and size effect on the drug delivery efficiency of Fe_3O_4 NPs	219
	Slab models of Fe ₃ O ₄	MD: COMPASS FF	Adsorption of chitosan chains on Fe ₃ O ₄ crystallographic planes	227
	6 nm Fe ₃ O ₄ NP with BSA proteins attached	MD: CHARMM27 FF, implicit model GBIS	BSA protein adsorption mechanism for super- paramagnetic nanocarrier design	228
	1–7 nm Fe ₃ O ₄ NPs with TMZ	DFT: PW91, MC/MD: pcff and COMPASS FFs	Effect of water and NPs size in the bonding between Fe ₃ O ₄ surface and anti-cancer TMZ compound	229
	2 nm Fe ₃ O ₄ NP coated with 2 Å thick Au shell	MD: CHARMM27 FF, TIP3P model	Application of a magnetic field for the crossing of core/ shell Fe ₃ O ₄ /Au NP through BBB	244
	3 nm Fe ₃ O ₄ NPs with ARA and PVA in water	MD: CHARMM36 FF, TIP3P water model	Assessment of interactions Fe ₃ O ₄ NPs with model cell membranes for theranostic purposes	243
	1–7 nm Fe ₃ O ₄ NPs functionalized with dopamine compounds	MC/MD: pcff and COMPASS FFs	Bonding of PEGylated dopamine with albumin molecules on Fe ₃ O ₄ NPs to enhance BBB permeation	245
	Graphene oxide nanosheets doped with Fe ₃ O ₄ NPs	DFT: PBE/Vanderbilt pseudopotentials	Fe ₃ O ₄ NPs doping in graphene oxide nanosheets to enhance magnetic properties for theranostic applications	256
CeO ₂	2–3 nm CeO ₂ NPs and α-syn monomers/fibrils	MD: universal FF	Aggregation study between CeO_2 NPs and α -syn amyloids associated with neurodegenerative diseases	115
Other	35.7 and 52.7 nm PEG-coated Au/ Ag NPs	CGMD: UA potential	Protein binding on the surface of PEG-coated NPs for nanocarriers and biosensors design	220
	5–200 nm Au/TiO ₂ NPs	CGMD: UA potential	Development of a model for fast estimation of protein affinities on NPs surfaces	222
	Clusters, (111) slabs for Ag/Au/Pt, 1.6–4.6 nm Ag NPs	DFT: PBE-D3/TZP, MD: Gromos53a6, SPC FFs	Adsorption of hydroxychloroquine on Ag, Au and Pt NPs for antiviral properties against COVID-19	232

wound healing. The binding affinities of various peptides indicated the formation of a dimer in solution that inhibits their attachment to graphene. Considering an additional binding domain, however, all peptides significantly increase their binding, which could guide the design of elastomeric nanomaterials

The superior photocatalytic activity of TiO2 NPs can be exploited in cancer therapy and diagnostics⁹⁵ but their broader application is hindered by potential toxicological manifestations observed in mice rats or other animal models.^{258,259} TiO₂ functionalization with PEG chains improves their biocompatibility avoiding premature elimination from the blood circulation by adsorption with serum proteins or by phagocytic uptake, enhancing NP efficacy in reaching tumor sites.²⁶⁰ Atomistic and coarse-grained MD simulations showed that PEGylated TiO2 NPs are less thermodynamically stable in lipid bilayers compared to polyethylene (PE)- or PE-PEG-coated NPs, which spontaneously migrate to the membrane, regardless of the chemical nature of the terminal group in PE-PEG chains.261

Peroxidases play a pivotal role in detoxification by catalyzing the reduction of hydrogen peroxide. While widely used in biosensors, these enzymes are expensive and sensitive to environmental conditions. As an alternative, metallic oxide nanomaterials, particularly iron oxide, offer similar catalytic activity, higher stability and large surface area at a lower cost. Shen et al. 262 evaluated the peroxide-like activity for a set of 15 iron-oxide slabs with different facet orientations and Fe vacancies by DFT, revealing dissociation of the chemisorbed H₂O₂ into hydroxyl groups. This led to a predictive chemical

descriptor for peroxidase-mimicking nanomaterials, in accordance with experimental data, proving the ability of computer simulation to aid the rational design of nanomaterials. Table 2 provides a summary of simulation studies of NP-macromolecule interactions relevant for biomedical applications, classified by nanomaterial type.

Multiphysics mesoscale modeling

The atomistic and coarse-grained simulations covered in section 2, usually model small systems of spherical NPs. However, fractal-like aggregates, typically in the range of tens to hundreds of nanometers, are often encountered in complex biomedical environments, where they interact with each other and with external fields, or are dispersed in solutions. In such scenarios, multiphysics mesoscale models can bridge experimental parameters and NP characteristics, helping to uncover structure-property relationships and optimize NP design. Various modeling approaches are available to address different aspects of NP behavior and performance. Discrete element method, Brownian dynamics, and continuum particle dynamics are commonly used to model NP dynamics during their synthesis, ²⁷⁷ focusing on particle movement, interparticle forces, and their collective behavior over time. These simulations offer insights into NP agglomeration 278,279 and transport behavior of irregularly shaped aggregates. 280 Applications of these techniques span from nanotoxicology^{281,282} and ecotoxicology²⁸³ to engineered nanofluid stability.²⁸⁴

Brownian dynamics, for example, have been used to predict the settling velocity of fractal-like SiO2 aggregates (Fig. 12a and b) suggesting a linear relationship with the ratio of the agglomerate mass to hydrodynamic diameter (Fig. 12c). 280 However, the assumption of monodisperse primary particles often made in settling rate calculations^{285,286} underestimates their settling velocity, as particle polydispersity and agglomerate morphology can significantly influence NP transport. For instance, in agglomerates with a polydispersity of 1.5 and a mass mobility exponent of 2.2 (i.e., open fractal-like aggregates), assuming monodisperse particles can lead to a 33% underestimation of settling velocity. This can potentially result in underestimating NP dose assessment in nanotoxicology studies or overestimating nanofluid stability. Ma et al. 287 explored the size-dependent dissolution of Ag NPs, by mass transport rate equations assuming first-order kinetics of Ag ion release. A faster volumetric reduction percentage was observed for octahedral Ag NPs compared to rods and cubes (Fig. 12d-f), corresponding to higher ion release rate, owing to the varying fraction of reactive edges and corners on NPs surface which are more prone to dissolution, as revealed by ab initio²⁸⁷ and reactive MD. Higher NaCl concentrations accelerate Ag+ dissolution (Fig. 12d) by the increase of the

mass transfer rate of Cl-. Similar first-order kinetics analysis was used to evaluate leaching from alloyed Ag-Au NPs. 288 Considering a shrinking-particle model, the continuous decrease in NP diameter was predicted along with enrichment in Au during the initial stages of dissolution. These leaching rate results combined with Mie theory simulations of scattering spectra were able to reproduce experimental leaching profiles.

Furthermore, multiphysics models are essential for understanding NP performance in external fields, such as electric 289 or magnetic fields. 290,291 Monte Carlo (MC), 3D full-wave computational models, or discrete dipole approximation techniques are commonly employed to evaluate the optical performance of plasmonic NPs, sought after for non-invasive photothermal therapies and thermally modulated drug delivery, while recent advancements in the simulation of light scattering²⁹² have increasingly focused on accelerating calculations to larger particles or more complex NP systems, such as NPs near planar surfaces.²⁹³ For example, finite-difference timedomain and finite element method (FEM) have been used to predict the absorption spectra and charge distribution of Fe@Au core-shell nanostars under near-infrared (NIR) irradiation (Fig. 13a). 294 NIR irradiation along the longitudinal

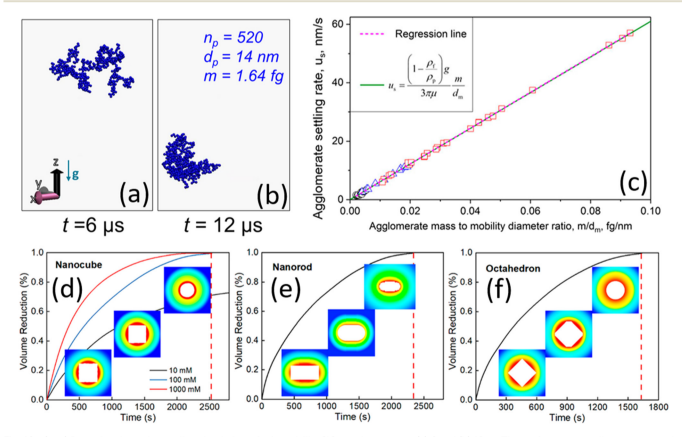


Fig. 12 (a-c) Brownian dynamics settling simulation of a fractal-like SiO₂ agglomerate at (a) 6 and (b) 12 µs. The agglomerate settling rate, µs, scales linearly with the agglomerate mass to mobility diameter ratio, $m/d_{\rm m}$ (c) (adapted from ref. 280 with permission from the American Institute of Physics, copyright 2018). (d-f) COMSOL multiphysics simulations of the volumetric reduction of Ag (d) nanocube, (e) nanorod, and (f) octahedron during dissolution (adapted from ref. 287 with permission from the American Chemical Society, copyright 2023).

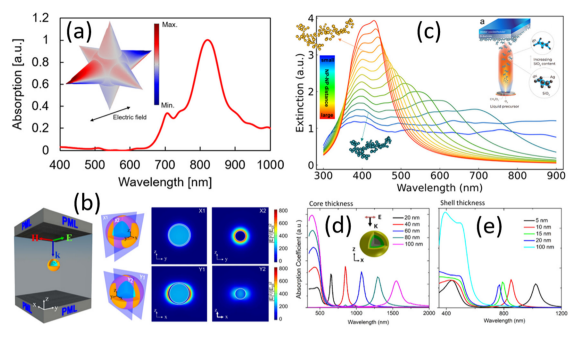


Fig. 13 (a) Simulated absorption spectrum of magnetic core-Au shell nanostar under NIR irradiation and charge distribution at the resonance wavelength (inset, adapted from ref. 294 with permission from Nature Portfolio, copyright 2020). (b) Local field enhancement of the SiO₂@Au coreshell structure at the localized surface plasmon resonance wavelength of 800 nm (adapted from ref. 295 with permission from Nature Portfolio, copyright 2016). (c) Simulated extinction spectra of fractal-like Ag agglomerates with different interparticle NP-NP distance.²⁹⁶ The inset shows nanoaggregates with large (yellow, top-left inset) and short (green, bottom-left inset) interparticle distances and a schematic of flame aerosol deposition experiments of as-prepared Ag/SiO₂ nanoaggregates (top-right inset) (adapted from ref. 296 with permission from the American Chemical Society, copyright 2021). (d) Absorption coefficient as a function of the wavelength at different Ag core sizes for a constant SiO₂ and Au shell thickness of 10 nm. The inset shows a multilayered Ag@SiO2@Au NP with a Ag core size of 40 nm surrounded by a SiO2 and Au shells. (e) Absorption coefficient as a function of the wavelength at different Au shell thicknesses for a 10 nm thick SiO₂ layer and Ag core size of 40 nm (adapted from ref. 298 with permission from Nature Portfolio, copyright 2019).

axis of the nanostars produced two absorption peaks but also induced a strong dipole moment in the direction of the incident light, leading to charge separation along the branches and tips of the nanostar (Fig. 13a: inset), confirming their potential for NIR-triggered drug release. In addition, 3D fullwave computational modeling showed that SiO₂@Au coreshell structures (Fig. 13b) and Au nano-cages exhibit higher local field enhancement compared to Au nanorods due to higher degree of rotational symmetry, making them more efficient photothermal transducers. 295 Merkl et al. 296 demonstrated, through a combination of UV-Vis spectra measurements and extinction spectra calculations, 297 that primarily interparticle spacing affects plasmonic coupling in flamemade Ag/SiO₂ nanoaggregates, rather than the constituent particle size and overall aggregate morphology. A reduction in interparticle spacing was achieved by controlling the thickness of the SiO₂ dielectric spacer, leading to a progressive increase in NIR extinction (Fig. 13c), highlighting the potential of these nanoaggregates as functional coatings for medical devices.

The development of materials for biomolecular detection with single-molecule sensitivity in the tissue transparent NIR region holds great promise in biomedicine. Optimizing nearfield enhancement and tunable resonance in multilayered Ag@SiO2@Au NPs is crucial for Raman scattering and sensing. FEM simulations²⁹⁸ show that increasing the Ag core diameter from 40 to 80 nm, shifts the absorption peak into the IR region, when maintaining the SiO₂ dielectric spacer and Au shell thickness at 10 nm (Fig. 13d). In contrast, decreasing the Au shell thickness from 100 to 5 nm shifts the absorption peak from red to NIR for a 40 nm Ag core (Fig. 13e), while reducing the interparticle spacing for a Ag@SiO₂@Au dimer maximizes the near-field enhancement.

The resonance between electron oscillations of metallic NPs (e.g., Au and Pt NPs), with electromagnetic waves generates significant heat, enhancing the photothermal and thermoplasmonic properties of these materials, making them suitable for theranostics and cancer therapy applications. While Au NPs have been studied extensively, 299,300 Pt-based NPs have received little attention. Samadi et al. 301 investigated the thermoplasmonic properties, photothermal effects on cancer cells, and the toxicity of Pt NPs, using FEM simulations to support their experimental findings. The simulations showed that the absorption of light and irradiated temperature of Pt NPs matched the experimental data. Larger Pt NPs exhibited significantly higher irradiated temperatures and maintained structural integrity at high temperatures. Notably, the formation of a vapor layer around irradiated Pt NPs suggested potential structural changes. Simulated temperature profiles at a water/glass interface indicated that the Pt NP could become embedded in the

glass surface if vapor layers formed, with temperatures reaching around 900 $^{\circ}\mathrm{C}.$

Additionally, a FEM-based model was developed to predict the plasmonic resonance of Au NPs embedded in breast cancer cells. By selecting appropriate factors, such as NP assembly, surrounding medium (refractive index), and interparticle distance, the model explained experimental results, such as the higher scattering contributions of larger NPs in the optical response after light irradiation.

4 Conclusions and outlook

Nanomaterials emerge as versatile tools for biomedical applications, offering transformative possibilities from combating neurodegenerative disorders and cancers to enhancing medical equipment with antibacterial coatings and advancing biosensors. A thorough understanding of nanoparticle (NP) behavior in biological fluids and their interactions with biomolecules is essential to fully integrate them into biomedical applications and achieve clinical translation. Computational approaches facilitate the development of innovative nanomaterials by enabling exploration of NP stability, antibacterial activity, potential toxicological manifestations, and drug release rates in NP-drug systems. This review highlights the significance of computational simulations in understanding NP physicochemical characteristics, including chemical stability, oxidation, and reactivity in biofluids. Density functional theory (DFT) and molecular dynamics (MD) simulations can unravel the intricacies of nanoscale interactions that cannot be observed directly experimentally, by elucidating NP reactivity, and long-term behavior of NPs in biological media. Case studies of NP-protein interactions and drug delivery systems underscore the utility of these techniques in designing biocompatible, functional NPs with enhanced targeting capabilities. The potential of NPs in treating neurodegenerative diseases, such as Alzheimer's and Parkinson's, is particularly compelling. Computational simulations can aid the discovery of nanomaterials tailored for these applications, the rational drug design, and the understanding of nano-biomolecular interactions.

However, force field development is pivotal for the next generation of computational nano-biomedical research. Surface NP characteristics, including crystal facets and surface chemistry, are critical to NP interactions with biological molecules. Current forcefields frequently face limitations in transconditions. ferability across different systems and Furthermore, advancements in nanoparticle science and the creation of functional multicomponent nanoparticles necessitate the development of accurate forcefields. For example, doped oxides such as Co-doped iron oxide, 303 superparamagnetic iron oxide nanoparticles (Zn_{0.5}Fe_{2.5}O₄ Mn_{0.5}Fe_{2.5}O₄) interacting with drug molecules, ³⁰⁴ hybrid bioglass - metal oxide nanoparticles (CeO₂ or Mn₃O₄), 305 and other nanocomposites such as SiO₂-coated TiN, 306 recently synthesized in academic laboratories show promise for applications in oral drug delivery, wound healing, and radiation therapy, highlighting this need. The reactivity and/or electromagnetic properties of these nanoparticles are also critical for their efficacy in these applications, requiring sophisticated force fields to precisely model such properties.

A second major challenge in computational biomedicine is the direct comparison between simulations and experiments, due to the different length and time scales each can accurately track. Simulations are typically limited to small nanoparticles, while experiments cannot routinely monitor the dynamics of nanoparticle-biomolecule interactions at the nanoscale. The implementation of mesoscale and multiscale simulations for modeling NP transport and NP response to external fields, can bridge the gap between experimental parameters and nanostructural characteristics, supporting the design of NPs for nontoxic drug delivery, photothermal therapy, and MRI contrast agents for biological sensing and disease diagnosis.

Although still in their early stages of development, advanced computational approaches and technologies, such as physics-informed machine learning (ML) and quantum computing, show promise for effectively bridging length and time scales in simulations. These ML models harness potential to significantly enhance the efficiency and accuracy of predictions in molecular modeling, while quantum computing can revolutionize molecular simulations by drastically accelerating computations, enabling more realistic, extensive, and sophisticated simulations. However, realizing these advancements requires fostering multidisciplinary collaborations and overcoming communication barriers across chemistry, physics, engineering, computer science, and mathematics disciplines.

Author contributions

D. C.: conceptualization, writing – original draft. E. G.: conceptualization, writing – review and editing, funding acquisition, supervision.

Data availability

No primary research results, software or code have been included and no new data were generated or analysed as part of this review.

Conflicts of interest

The authors declare no competing financial interest.

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Review

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