Surface Functionalisation of Upconversion Nanoparticles with Different Moieties for Biomedical Applications

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Abstract: Lanthane ion-doped upconversion nanoparticles (UCNPs) that can convert low-energy infrared photons into high-energy visible and ultraviolet photons, are becoming highly sought-after for advanced biomedical and biophotonics applications. Their unique luminescent properties enable UCNPs to be applied for diagnosis, including biolabeling, biosensing, bioimaging, and multiple imaging modality, as well as therapeutic treatments including photothermal and photodynamic therapy, bio-reductive chemotherapy and drug delivery. For the employment of the inorganic nanomaterials into biological environments, it is critical to bridge the gap in between nanoparticles and biomolecules via surface modifications and subsequent functionalisation. This work reviews the various ways to surface modify and functionalise UCNPs so as to impart different functional molecular groups to the UCNPs surfaces for a broad range of applications in biomedical areas. We discussed commonly used base functionalities, including carboxyl, amino and thiol moieties that are typically imparted to UCNPs surfaces so as to provide further functional capacity.

Keywords: upconversion; nanoparticles; lanthanide; surface modification; functionalisation; ligand engineering; silanisation

1. Introduction

Extensive research over the past decade has led to an expanded understanding and development of nanomaterials for their considerably broad use across both the scientific and technological spectrums, with disciplines such as nanomedicine, biotechnology, and forensics exhibiting excellent potential for application. Recent advancements have seen the evolution of nanoparticles for their uses in theranostics, including bioimaging, selective therapeutic delivery and biosensing applications where theranostics is a concept that was derived to describe the development of specific, individualised therapies for diseases, whilst merging therapeutic and analytical approaches through multi-modal application [1,2].

The intrinsic photo-physical properties of upconversion nanoparticles make them particularly attractive candidates for diagnostic operations whilst offering the capability to phototherapy. A comprehensive and wide-ranging selection of surface modification methods allows for the specific design of physico-chemical, toxicological, and pharmacological properties, demonstrating their promising versatility across a myriad of practical applications [3-5]. Therefore, the surface modification approaches for UCNPs are selected basically depending on improvements to nanoparticle biocompatibility whilst allowing them to be protected from surrounding aqueous media, thus
enhancing both the dispersion and denaturation stability of the nanoparticles, preventing their
dissolution into ionic species [4,6,7].

Popular choices for the surface modification of UCNPs include molecular ligands or silanisation
as they naturally encapsulate and shield the nanoparticles from external interactions, thus substantially
minimising nanoparticle dissolution and unwanted exchanges [5,8]. Another important role of the
surface modified polymer molecules or silica shell is to provide further functionality to the UCNPs for
desired operations. Design-enhanced colloidal stability [5,9–17] and bio-conjugation [2,15,16,18–26] are just
a few examples from the extensive of significant benefits that can be gained from the functionalisation
of UCNPs with functional moieties. There is a small range of common base functional moieties,
including carboxyl (–COOH), amino (–NH₂) and thiol (–SH) groups, but their utilisation across a broad
spectrum of applications differs. Environmental influences such as pH, magnetic fields, light, salts,
and temperature are manipulated in various ways so as to achieve success through interactions with
these functional moieties [27]. Although the moieties can be the same, the desired effect of the external
influences may be completely opposing based on the given application.

2. Surface Modifications

Commonly synthesised UCNPs are typically capped by surfactant ligands (such as oleate) that
aid in the stabilisation of the particles from aggregation. The ligands help to control the growth
of the nanoparticles via surface co-ordination while also acting as a solvent during synthesis [5,28].
However, these ligands tend to exhibit strong hydrophobic qualities or have minimal functional
moieties for further application [3–5,7,29]. Additionally, hydrophilic ligands used in hydrothermal
synthesis that co-ordinate via amine or carboxylic groups can be used in the synthesis to impart
hydrophilicity but may not supply or allow the modification of necessary functional moieties for
desired use [3,5]. Therefore, subsequent modification of UCNPs surfaces allows for specifically
tailored or additional physical, chemical or biological attributes, which differ from those originally
present on the nanoparticles, to be augmented onto the material surface. This confers changes to
hydrophilicity, surface charge and surface energy amongst other properties, for their use in aqueous
mediums (specifically in biological systems) [5–7]. Biocompatibility is a significant priority in the
design of surface modifications for particles that are intended for use in nanomedical applications,
however, a lack of ubiquitous and consistent methodology in the field presents a substantial obstacle
along the way to further developments [30]. Moreover, nanoparticles intended for use in biomedical
applications require uniformity in morphology and size which presents a greater difficulty in obtaining
ideal samples [29].

Various modification strategies exist that have tried to develop universal methods and overcome
these limitations but stability in aqueous environments is still a considerable challenge [10]. The four
main strategies for surface modification include: (a) ligand engineering [2,3,5,7,9,10,12,31]; (b) layer-by-
layer assembly [2,3,5,7,32]; (c) ligand attraction [2,3,5,7,33]; and (d) surface polymerisation [2,3,5,6,
11,13,15–17,34–39]. Each modification method as well as the specific molecules selected, has its own
benefits and hindrances. While there are many ways to modify the surfaces of UCNPs, this review
aims at covering the two most popular methods, that being ligand engineering and silanisation.

2.1. Ligand Engineering

Ligand engineering comprises two methods of surface modification. The first being the oxidation
of ligands present on UCNPs surfaces (e.g., oleate) via an oxidiser such as the Lemieux-von
Rudloff reagent or ozone. Figure 1 shows a typical ligand oxidation process, which could result
in terminal carboxylates, aldehydes or epoxides [5,17,40]. The oxidative alteration of ligands is not a
commonly used approach to UCNPs surface modification due to the poor dispersal in aqueous
media of resultant UCNPs and the limited variability in functional moieties [4,5,29]. Although
ligand oxidation exhibits no influence on morphology or size of UCNPs [41,42], work conducted
by Naccache, R. et al. (2009) [43] determined that prolonged oxidation (>2 h) of oleate ligands on
NaGdF₄: Yb³⁺, Ho³⁺ UCNPs resulted in an MnO₂ precipitate that was challenging to remove, reduced
the hydrophilicity of the UCNPs in water and significantly weakened the luminescent emission of the particles. Interestingly, Deng, R. et al. (2011) [42] found that the addition of glutathione (GSH) in low amounts to MnO$_2$-modified NaYF$_4$: Yb$^{3+}$, Tm$^{3+}$ UCNPs led to a substantial recovery of emission from the particles. It is asserted that the enhancement of the optical properties was a result of GSH-mediated reduction of MnO$_2$ to Mn$^{2+}$. Therefore, in the case of prolonged oxidation, it may be possible to remove the MnO$_2$ and recover emission of the particles if necessary.

![Figure 1](image1.png)

**Figure 1.** Scheme of the oxidation of oleate with the Lemieux–von Rudloff reagent (MnO$_2^-$/IO$_4^-$). Note that the dimensions of the NPs and ligands are not to scale. Reprinted with permission from [41]. Copyright 2008 American Chemical Society.

The second method defined under ligand engineering is the ligand exchange process, which is an effective technique to replace hydrophobic ligands on the UCNPs surface with the ligands that exhibit both greater affinity and hydrophilicity. The process is easy to operate, and exhibits a negligible effect on the morphology of the yielded UCNPs. The driving force behind this reaction is that hydrophilic ligand has a stronger coordination ability to lanthanide ions than the original hydrophobic ligand [44]. Figure 2 is the scheme of the ligand exchange process for surface modification of UCNPs. A ligand exchange reaction does not necessarily involve full replacement of all the hydrophobic ligands present on the surface, however, the complete exchange of ligands is ensured if an excess of the new ligand is added to a suitable solvent at elevated temperatures, with the choice of solvent being reliant on the dynamic solvability of both the hydrophobic and new hydrophilic ligands [5].

![Figure 2](image2.png)

**Figure 2.** Schematic illustration of the ligand exchange process where the original oleate capping ligands on the surface of the upconversion nanoparticles are replaced by the PEG-Phosphate ligands. Note that the dimensions of the NPs and ligands are not to scale. Reprinted with permission from [45]. Copyright 2010 American Chemical Society.
Various hydrophilic ligands have been introduced onto the surfaces of UCNPs via ligand exchange, including ligands such as citrate [3,5,29], hexanedioic acid [3,5,29], polyethylene glycol (PEG) derivatives [5,10,29,31], 6-aminohexanoic acid [3,5,46], polyacrylic acid (PAA) derivatives [10,43,47], and phosphate derivatives [10,31,48]. Newly affixed ligands tend to require a certain length in order to be able to provide functionality that can be accessible for subsequent modification but this can be advantageously used in the design of nanoparticle stability. For instance, 1,10-decanedicarboxylic acid (DDA) has been demonstrated to closely crosslink NaYF$_4$: Yb$^{3+}$, Er$^{3+}$ UCNPs and Fe$_3$O$_4$ NPs via terminal carboxylic acid moieties. Due to the short length of the DDA ligand compared to the oleate ligand, oleate concentration could be optimized so as to functionalise the UCNPs with both oleate and DDA in an advantageous manner via the balance of the ligand exchange progress dynamically. The length of the oleate ligands prevent undesired interactions with anchored carboxylic acid groups present on the shorter DDA ligands by means of steric hindrance, thereby preventing substitution of the crosslinker [49]. Preferably, the newly affixed ligands should be multidentate as the type and number of co-ordination sites on the hydrophobic and hydrophilic ligands strongly influence the efficacy of the exchange. For example, the higher co-ordination displayed by carboxylates makes them a favoured choice over amine-containing ligands, particularly when coating positively-charged UCNPs (i.e., NaYF$_4$: Yb$^{3+}$, Er$^{3+}$) [50].

2.1.1. Carboxyl Moiety Modified UCNPs via Ligand Engineering

The exchange of ligands present on as-synthesised UCNPs with ligands exhibiting a carboxylic acid functional moiety would be the most effective method of modification under ligand engineering to impart such functionality. Due to the hydrophilic properties exhibited by carboxyls as well as the ease of coupling between carboxylic moieties and amine residues commonly found on biomolecules, it is a prevalent functionality for bio-sensing applications [41,51–53]. Furthermore, an additional benefit gained by carboxyl functionalisation is the capability to store particles for lengthier periods at an alkaline pH due to deprotonation of the terminal hydroxyl present on the carboxylic acids. The presence of a formal negative charge gives rise to a larger negative zeta potential and, thus, greater colloidal stability of the sample, substantially reducing agglomeration of the nanoparticles over time [17]. Table 1 summarised the surface modified UCNPs via ligand engineering with carboxyl moieties and their corresponding applications.

<table>
<thead>
<tr>
<th>UCNPs</th>
<th>Ligand Coordinating Moiety</th>
<th>Functional Moiety</th>
<th>Method of Functionalisation</th>
<th>Application</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>NaYF$_4$:Yb$^{3+}$, Tm$^{3+}$</td>
<td>DTPA</td>
<td>Carboxylate</td>
<td>EDC Chemistry</td>
<td>Hg$^{2+}$ DNA-based biosensor</td>
<td>51</td>
</tr>
<tr>
<td>NaYF$_4$:Yb$^{3+}$, Tm$^{3+}$</td>
<td>DPTA</td>
<td>Carboxylate</td>
<td>EDC/NHS Chemistry</td>
<td>DNA-based biosensor</td>
<td>52</td>
</tr>
<tr>
<td>NaYF$_4$:Yb$^{3+}$, Tm$^{3+}$</td>
<td>PAA</td>
<td>Carboxylate</td>
<td>EDC/NHS Chemistry</td>
<td>Ligase assisted DNA-based biosensor</td>
<td>53</td>
</tr>
<tr>
<td>NaYF$_4$: Yb$^{3+}$, $X^{3+}$ (Ho$^{3+}$, Er$^{3+}$ or Tm$^{3+}$)</td>
<td>AA</td>
<td>Carboxylate</td>
<td>Ligand Exchange</td>
<td>ZnPc-based photodynamic Therapy</td>
<td>41</td>
</tr>
<tr>
<td>NaYF$_4$: Yb$^{3+}$, Er$^{3+}$</td>
<td>SOC-Chitosan</td>
<td>Carboxylate</td>
<td>Ligand Exchange</td>
<td>UCNP-based platform for bio-application</td>
<td>54</td>
</tr>
<tr>
<td>NaYF$_4$: Yb$^{3+}$, Er$^{3+}$</td>
<td>MA-PEG</td>
<td>Carboxylate</td>
<td>Electrostatic Interactions</td>
<td>pH-manipulated drug delivery</td>
<td>55</td>
</tr>
<tr>
<td>NaYF$_4$: Yb$^{3+}$, Er$^{3+}$</td>
<td>PAA</td>
<td>Carboxylate</td>
<td>Electrostatic Interactions</td>
<td>Multifunctional liposomal nanocarriers</td>
<td>56</td>
</tr>
<tr>
<td>NaGdF$_4$: Yb$^{3+}$, Er$^{3+}$</td>
<td>Citrate</td>
<td>Carboxylate</td>
<td>Electrostatic Interactions</td>
<td>Multifunctional liposomal nanocarriers</td>
<td>26</td>
</tr>
</tbody>
</table>
Two representative works published by Kumar and Zhang (2009–2010) [51,52] established the realisation of DNA-based bio-sensors via the modification of NaYF$_4$: Yb$^{3+}$, Tm$^{3+}$ UCNP surfaces with diethylenetriaminepentaacetic acid (DTPA). A ligand exchange with oleate allowed for modification of DTPA onto the UCNP surface whereby exposed carboxyl groups were then coupled to terminus –NH$_2$ groups on ss-DNA via carbodiimide chemistry without the addition of NHS derivatives. Whilst the bio-sensors are similar in concept, they are slightly different in practice. The first sensor [51] relies on ss-DNA that is present on the surface of the UCNP to convert to a hairpin conformation following interactions with free Hg$^{2+}$ ions in solution. The change in conformation not only entraps the free Hg$^{2+}$ ions but an intercalating dye (SYBR Green I) that is also present in solution. When the nanoparticles are excited, their emission is absorbed by SYBRG (~477 nm) followed by SYBRG emission between 490–520 nm thus indicating entrapment of Hg$^{2+}$ and the realisation of a DNA-based mercuric ion bio-sensor. The secondary sensor [52] works via the conjugation of complimentary ss-DNA in solution to the ss-DNA present on the UCNP surface. If complimentary ss-DNA successfully links to the UCNP, the intercalating dye (SYBRG) present in the solution gets trapped between the cross-linked DNA, so when the UCNPs are excited their emission is absorbed by the dye and a subsequent emission at 490–520 nm is seen. This signals the presence of complimentary DNA, which aims to be correlated to identify diseases, mutagens or pathogens by association.

Using the same carboxy group exposed on the UCNP surface, further work conducted by Wang and Zhang (2014) [53] determines a ligase-assisted DNA-based bio-sensor by addition of surface carboxyls through ligand exchange of oleate-capped NaYF$_4$: Yb$^{3+}$, Tm$^{3+}$ with poly(acrylic) acid. Amine-terminated ss-DNA was conjugated to the UCNP via carbodiimide and NHS chemistry, followed by introduction of an intercalating dye (SYBRG), complementary ss-DNA target ss-DNA into solution. The ss-DNA present on the UCNP as well as the complementary ss-DNA in solution are matching segments to the target ss-DNA, where a successful match leads to DNA hybridisation followed by ligation, dehybridisation of target DNA and a subsequent hairpin formation between the segments (Figure 3). Ligation of the DNA and subsequent hairpinning would not occur if the segments of ss-DNA were mismatched to the target ss-DNA in solution. As a consequence, this mechanism of action allows for both accurate selectivity and sensitivity regarding determination of target ss-DNA presence. Furthermore, system temperatures during the experiment were manipulated to achieve desired results; this is due to hybridisation occurring at low temperatures and dehybridisation occurring at high temperature, so alteration of temperatures allowed for the acceleration of these processes. Advantageously, the addition of PAA to the surface of the UCNP allowed for polymeric encapsulation of the nanoparticle cores, which is ideal for the protection from the surrounding aqueous environment, thus prolonging the storage life of the particles.

Although not a typical approach, the oxidation of ligands present on the surface of UCNP as-synthesised can also be a means of achieving bio-sensing capabilities. Research conducted by Chen et al. (2008) [41] demonstrated the biosensing application of various lanthanide-doped UCNP (NaYF$_4$: Yb$^{3+}$ with activator variations of Er$^{3+}$, Ho$^{3+}$ or Tm$^{3+}$) coated with azelaic acid (AA) ligands. The oleate-capped UCNP were directly oxidised via addition of the Lemieux-von Rudloff reagent, yielding an exposed terminal carboxylic acid group on the azelaic acid ligands. Addition of the carboxyl moiety imparted both hydrophilicity and bio-conjugation functionality, thus allowing the subsequent cross-linking of streptavidin to the terminal carboxyls via carbodiimide chemistry involving NHS. The streptavidin was further conjugated to capture-DNA that was added to solution with complimentary reporter-DNA attached to a TAMRA quencher. Given that the complimentary reporter-DNA successfully conjugates to the capture-DNA present on the UCNP, the quencher would then be located in close proximity to the UCNP. This allows for luminescent resonance energy transfer (LRET) to occur as phosphorescence emitted by the UCNP is absorbed by the quencher due to the absorption spectrum of the quencher overlapping the green emission (~544 nm) of Er$^{3+}$. 
The use of carboxyl moieties is not limited to biosensor applications but can also be used for multi-modal theranostic purposes as well, with one possible example being the use of chitosan coated UCNPs for their use in photo-dynamic therapy as demonstrated by Cui et al. [54]. Amphiphilic chitosan (specifically $N$-succinyl-$N'$-octyl chitosan) was coated onto NaYF$_4$: Yb$^{3+}$, Er$^{3+}$ UCNPs whereby the hydrophobic photosensitiser ZnPc was loaded into the chitosan. However, due to the amphiphilic nature of the chitosan (SOC), it can be either coated via layer-by-layer attraction (hydrophobic interactions with surface residing oleate alkyl chains) due to exposed octyl groups on SOC or through a ligand exchange via terminal carboxyl groups. Either process would not change the overall mechanism of the application but the latter would be an example of carboxyls being used exclusively for their co-ordination abilities (in the same way oleate coordinates to UCNPs). ZnPc was loaded internally into the SOC via hydrophobic interactions whereby excitation of the UCNPs leads to 660 nm visible light emission that activates the release of reactive oxygen species (ROS) from activated ZnPc molecules. The researchers determined a loading capacity of 10.8%, a marked improvement from a comparison where ZnPc loading in mesoporous silica showed only ~0.1% loading capacity [57], most likely due to the hydrophilic nature of the silica. Furthermore, induced cell apoptosis was determined in MCF-7 cells as a result of successful ROS release, thereby indicating the successful realisation of an UCNP-based photodynamic therapy system.

Similarly, Liebherr et al. (2012) [55] developed colloidal-stable NaYF$_4$: Yb$^{3+}$, Er$^{3+}$ UCNPs via ligand exchange of oleate with maleimide-polyethylene glycol (MA-PEG) that exhibits terminal carboxyl groups. The carboxyl groups were used to coordinate the MA-PEG to the UCNPs, thus resulting in exposed thiol-sensitive maleimide groups that can be used for bioconjugation. Coupling of FITC-labelled $\gamma$-globulin to the UCNPs was accomplished through addition of tris(2-carboxyethyl)phosphine (TCEP), a reduction agent that reduces existing cysteine disulphides present in the $\gamma$-globulin, allowing for thioether bond formation with the exposed maleimide groups. Confirmation of successful coupling was proven via the measuring of fluorescence of the labeled $\gamma$-globulins on both activate and inactive maleimide-functionalised UCNPs whereby active UCNPs demonstrated greater relative red emission while dispersed in water than inactive UCNPs. The reported UCNPs can be utilised as a platform in various bioconjugation applications involving thiols, such as biosensing, bioimaging, and therapeutic delivery.
The aforementioned examples represent not only the various ways to modify the surface of UCNPs with terminal carboxyls but for the most part, typical applications involving the carboxyl moiety being used for the addition of hydrophilic properties, biomolecule cross-linking capabilities and/or co-ordination capabilities. However, surface moieties can exhibit complex multi-functional properties that can give rise to effective utilisation in various applications. Jia et al. (2013) [56] developed surface modified NaYF₄: Yb³⁺, Er³⁺ UCNPs with poly(acrylic) acid for their use in a pH-triggered controlled drug delivery system that demonstrates various functionalities of terminal carboxyl groups in practice. At a neutral pH, the anti-cancer drug doxorubicin (DOX) was bound to terminal carboxyl groups residing on the poly(acrylic) acid surface via electrostatic interactions following deprotonation of the carboxyls. As DOX exhibits a positive charge, it becomes tightly bound to the negatively charged carboxyls, leading to the conjugation of DOX molecules surrounding the surface of the UCNPs. Moreover, alteration of pH conditions can be used to manipulate desired operations; the loading rate of DOX was determined to be high in weakly alkaline environments (approx. 1% at pH 2, and 32% at pH 7) whilst the rate of release was determined to be high in acidic environments. Further assessment by the group testified to successful drug delivery within in vitro HeLa cells by cytotoxicity assays, where cell viability decreased by 50%. In this case, the carboxyl moieties demonstrate multi-functionality, that being: (1) the capacity to conjugate the anti-cancer drug DOX; (2) the control of the rates of both loading and release via manipulation of environmental pH; (3) imparting hydrophilic properties to the UCNPs so as to increase biocompatibility through surface residing carboxyls; and (4) coordinating poly(acrylic) acid to the UCNPs through internal carboxylate interactions.

Finally, NaGdF₄: Yb³⁺, Er³⁺ UCNPs were modified to achieve multifunctional upconverting liposomal nanocarriers for their use in anti-cancer drug delivery treatments by Huang et al. (2016) [26]. The surface of the UCNPs was modified with citrate ligands which exhibit tri-carboxyl moieties that demonstrate multiple functionalities in practice (as shown in Figure 4). Firstly, deprotonation of the carboxyls leads to subsequent coordination of the citrate ligands to the surface of the UCNPs via a terminal carboxylate moiety. The two remaining surface carboxyl groups then are used to (1) impart hydrophilic properties to the UCNPs; (2) provide a hydrophilic environment for the loading of the anticancer drug DOX; and (3) conjugate as well as stabilise DOX molecules through ionic interactions (carboxylates exhibit a net negative charge whilst DOX exhibits a net positive charge). As DOX has an absorption range overlapping with part of the UCNP emission range (515 nm–570 nm), UV–VIS can be used to quantify the loading capacity of the UCNPs. Following encapsulation of the UCNPs in a liposome made of 1,2-dioleoyl-sn-glycero-3-phosphocholine (DOPC) and cholesterol in a ratio of 2:1, DOX was loaded into Lipo@DOX and UCNP@DOX@Lipo samples. It was determined that the loading efficiency of Lipo@DOX was 98% whilst UCNP@DOX@Lipo demonstrated a loading efficiency of 72%, the group asserted that this is most likely due to the decreased internal volume of the liposome from UCNPs occupying internal space. Passive drug release of the UCNP@DOX@Lipo particles was tested in both solutions of phosphate buffered saline (PBS) (imitates in vitro conditions) and 50% foetal bovine serum (FBS) in PBS (imitates in vivo conditions), with passive leaking only accounting for 6% loss of DOX in PBS over a 12 h period at 37 °C. As the overall passive release was minimal, PBS can be considered a viable storage solution whilst passive release of 39% in 50% FBS + PBS solution indicates the influence of external factors, such as serum protein and lipid bilayer destabilisation interactions that further the complexity of application in vivo.
Figure 4. Schematic representation showing the structures of the blank liposome, Lipo-UCNPs, Lipo-DOX, and Lipo-UCNPs-DOX. When UCNPs and DOX were incorporated into the liposome, UCNPs and DOX were located in the inner core of the liposome. Upon laser irradiation (980 nm), energy transfer occurred from UCNPs to DOX, where the upconverted green emission is partially absorbed by the DOX. Reprinted with permission from [26]. Copyright 2016 American Chemical Society.

2.1.2. Amino Moiety Modified UCNPs via Ligand Engineering

Much like carboxyl moieties, amino groups are favoured due to the hydrophilic properties they impart onto UCNPs and their ability to also be cross-linked via the well-established carbodiimide coupling protocol. Conversely, they exhibit positive charges at physiological pH which may be desirable for certain applications. The summary of the surface modified UCNPs via ligand engineering process with amino moieties is exhibited in Table 2.

Table 2. Summary of the surface modified UCNPs via ligand engineering with amino moieties.

<table>
<thead>
<tr>
<th>UCNP</th>
<th>Ligand</th>
<th>Coordinating Moiety</th>
<th>Functional Moiety</th>
<th>Method of Functionalisation</th>
<th>Application</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>NaYF₄: Yb³⁺, Tm³⁺</td>
<td>PEA</td>
<td>Phosphate</td>
<td>Amino</td>
<td>EDC/NHS Chemistry</td>
<td>Remotely triggered anti-cancer system</td>
<td>[25]</td>
</tr>
<tr>
<td>NaYF₄: Yb³⁺, Tm³⁺</td>
<td>PEA</td>
<td>Phosphate</td>
<td>Amino</td>
<td>Reductive Amination Nucleophilic Thiourea Formation</td>
<td>Solid-phase biosensing</td>
<td>[58]</td>
</tr>
<tr>
<td>NaGdF₄: Yb³⁺, Er³⁺</td>
<td>PMAM</td>
<td>Amino</td>
<td>Amino</td>
<td></td>
<td>Targeted lectin recognition</td>
<td>[59]</td>
</tr>
</tbody>
</table>

An exchange between oleate and o-phosphorylethanolamine (PEA), which exhibits both terminal amino and phosphate groups, was carried out on NaYF₄: Yb³⁺, Tm³⁺ UCNPs in research conducted by Fedoryshin et al. (2014) [25]. The exchange led to the affixing of primary amines to the surface of the particles for their use in a remotely triggered anti-cancer drug delivery system (as displayed in Figure 5). As phosphate exhibits both greater coordination sites and binding affinity than carboxylates, the exchange favours the replacement of oleate carboxylate groups with terminal phosphates [10]. Conjugation of the photocleavable prodrug ONB-5-FU to the UCNPs via carbodiimide chemistry involved the addition of NHS, resulting in the formation of an amide bond. Due to ONB-5-FU exhibiting an overlapping UV absorption range with that of UV emission range of the UCNPs,
excitation of UCNPs resulted in 364 nm UV light emission that efficiently photocleaved the prodrug. This allowed for detachment and consequent release of 5-fluorouracil, where cleavage is proposed to occur at the C-N bond that couples ONB and 5-FU. It was determined that the rate of release could be tuned with laser power output (10, 30 and 80 mW) and that as high as 77% of the prodrug was effectively released via remote triggering through NIR excitation in comparison to traditional direct UV excitation. Laser excitation at 80 mW resulted in an observed initial rate constant (k_0) of 130 μM min⁻¹ while, conversely, at 10 mW, a k_0 of 18 μM min⁻¹ was observed.

Similarly, work presented by Doughan et al. (2014) [58] utilises PEA to realise a biosensing application through solid-phase covalent immobilisation of UCNPs (Figure 6). A significant problem presented to the applications of UCNPs in aqueous mediums is the agglomeration of particles in solution, which can be overcome through the use of UCNPs in solid-phase at interfaces. Standard as-synthesised NaYF₄: Yb³⁺, Tm³⁺ UCNPs were transitioned to a hydrophilic phase via a ligand exchange with oleate carboxylate groups with terminal phosphates

Glass slides, which were used as the interface medium, were functionalised with surface silanol groups by washing with NH₄OH and HCl in the presence of H₂O₂. Subsequently they were amine functionalised via (3-aminopropyl)trimethoxysilane (APTMS) followed by reductive amination with sodium cyanoborohydride (NaBH₃CN). The resulting aldehyde-functionalised glass slides provided an interface for which the PEA-modified UCNPs could now be coupled and immobilised to through covalent bonding. Incubation of the glass slides over 2 h followed drop spotting of UCNPs with NaBH₃CN on desired locations so as to allow reductive amination to occur and the formation of a subsequent alkyl amine bond between the underside of the UCNPs and slide (as shown in Figure 6).

The surface residing amino groups on the UCNPs were functionalised with thrombin after residual aldehyde groups on the slide were inactivated with ethanolamine in the presence of NaBH₃CN. Amine-modified thrombin specific aptamer-1 was drop cast onto active UCNP spots after the surface amines were reacted with glutaraldehyde. As quantum dots (QDs) exhibit a spectral overlap with UCNPs, displaying an absorbance over their emission range (~330–490 nm), they can be used as effective energy acceptors in an LRET-based energy donor-acceptor system of UCNPs and QDs. As a result, QDs were functionalised with thrombin specific aptamer-2 and subsequently incubated for 6 h with UCNPs so as to allow the aptamers to attach. If the aptamers have successfully attached, the UCNPs will undergo a LRET process with the QDs when excited at 980 nm. Confirmation and quantification can be achieved via the consequent QD emission at 614 nm and peak ratio, whereby it was determined that the assay showed strong selectivity for thrombin through control testing. In contrast to 0.1 μM thrombin, a 1 μM solution of BSA displayed a signal that was 80% lower while
also demonstrating a signal that was 90% lower for the non-specific adsorption of QDs onto the surface of the interface.

Figure 6. Schematic showing immobilization of PEA-UCNP on aldehyde functionalised coverslip and use in the detection of thrombin (grey) using thrombin-specific aptamer-1 (red) and aptamer-2 (blue). Reprinted with permission from [58]. Copyright 2014 American Chemical Society.

An endeavour to produce carbohydrate-coated UCNPs for lectin recognition was undertaken by Bogdan et al. (2010) [59]. After NaGdF\(_4\): Yb\(^{3+}\), Er\(^{3+}\) UCNPS were synthesised with oleate ligands present on the surface, they subsequently underwent a ligand-exchange with poly(amido)amine (PMAM). As PMAM exhibits four terminal amino moieties, one moiety can exchange and co-ordinate to the UCNP surface while the remaining three groups provide both hydrophilicity and further bioconjugation capabilities. Synthetic multivalent carbohydrates demonstrate efficient inhibition of protein-carbohydrate interactions due to strong binding affinities. Therefore, by cross-linking synthetic carbohydrates to the UCNPs, the UCNPs gain the ability to interact and detect bacteria or virus infections. The synthetic carbohydrate p-isothiocyanatophenyl a-D-mannopyranoside was coupled to the UCNP's via thiourea bond formation between the terminal amino groups residing on the UCNP surface and the terminal thiocyanate on the synthetic carbohydrate. Further testing through an LRET assay led to the determination that the mannose moiety successfully conjugated to protein receptors by use of a RITC-labelled plant lectin Con A (RITC-Con A). As the RITC-Con A absorption overlaps with the emission range of the UCNPs (550 nm), it gives off a characteristic emission at 585 nm that can indicate successful attachment of the carbohydrate to protein receptors after UCNP excitation. The group demonstrated that there was a matching boost in acceptor fluorescence to UCNP green emission decline after excitation of the UCNP's at 980 nm. Furthermore, the addition of Gd\(^{3+}\) to the nanocore allows for great contrast when imaging via magnetic resonance imaging (MRI) as it exhibits a considerable magnetic moment.

2.1.3. Thiol Moiety Modified UCNPs via Ligand Engineering

Equivalently to carboxyl and amino moieties, thiol moieties can be used to co-ordinate to UCNPs, impart hydrophilicity and grant the capability to further conjugate biomolecules for
extended application. Although they cannot undergo carbodiimide reactions, thiols can present an alternative option for operations where carboxyl and amino cannot be applied effectively. Table 3 summarised the surface modified UCNPs via ligand engineering with thiol moieties and their corresponding applications.

**Table 3.** Summary of the surface modified UCNPs via ligand engineering with thiol moieties.

<table>
<thead>
<tr>
<th>UCNP</th>
<th>Ligand</th>
<th>Coordinating Moiety</th>
<th>Functional Moiety</th>
<th>Method of Functionalisation</th>
<th>Application</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe₃O₄/NaYF₄: Yb³⁺, Er³⁺</td>
<td>MUA</td>
<td>Carboxylate</td>
<td>Thiol</td>
<td>Affinity to Fe₃O₄</td>
<td>Fe₃O₄ enhanced superparamagnetic UCNPs</td>
<td>[49]</td>
</tr>
<tr>
<td>NaYF₄: Yb³⁺, Er³⁺</td>
<td>TGA</td>
<td>Thiol</td>
<td>Carboxylic Acid</td>
<td>Electrostatic Interactions</td>
<td>Ag⁺-assisted photothermal therapy Dual-mode luminescence</td>
<td>[60]</td>
</tr>
<tr>
<td>Yb³⁺, Tm³⁺: BaF₂/Ln³⁺:SrF₂</td>
<td>TGA</td>
<td>Thiol</td>
<td>Carboxylic Acid</td>
<td>Ligand Exchange</td>
<td>Active-core/Active-shell synthesis of UCNPs</td>
<td>[61]</td>
</tr>
<tr>
<td>NaYF₄: Gd³⁺, Eu³⁺</td>
<td>MPA</td>
<td>Thiol</td>
<td>Carboxylic Acid</td>
<td></td>
<td>Combined optical and MR imaging</td>
<td>[62]</td>
</tr>
</tbody>
</table>

Successful development of superparamagnetic Fe₃O₄/NaYF₄: Yb³⁺, Er³⁺ hybrid nanoparticles was demonstrated by Shen et al. (2010) [49] whereby 11-mercaptoundecanoic acid (MUA) was surface modified onto NaYF₄: Yb³⁺, Er³⁺ UCNPs via a ligand exchange with oleate ligands present on the as-synthesised particles. As MUA exhibits both terminal thiol and carboxylic moieties, it displays the ability to co-ordinate to the UCNPs via the carboxyl moiety whilst being capable of further cross-linking through residual thiol moieties. These thiol moieties play a crucial role in the cross-linking process due to the specific affinity they boast toward Fe₃O₄ NPs. Compared to UCNPs coated with 1,10-decanedicarboxylic acid (DDA) which exhibits terminal carboxyl moieties, the UCNPs@MUA were not only smaller in size than the UCNPs@DDA but also absorbed Fe₃O₄ NPs more densely. It is inferred that the MUA perhaps inhibits the growth of Fe₃O₄ on the surface of the UCNPs due to a robust interaction between Fe and S. Furthermore, the group present evidence to support effective magnetic separation ability whereby they explain that the ligand spacing between Fe₃O₄ and the UCNPs allows for the superparamagnetic properties of Fe₃O₄ to be kept intact.

Inversely, research by Dong, B. et al. (2011) [60] establishes the use of the thiol moiety as a co-ordinating moiety to the surface of NaYF₄: Yb³⁺, Er³⁺ UCNPs for therapeutic photothermal applications (as displayed in Figure 7). Notably, it has been reported that ligands which have been exchanged to UCNP surfaces via thiol coordination require additional functionalisation so as to undergo cell internalisation effectively [5]. As-synthesised UCNPs underwent a ligand exchange with thioglycolic acid (TGA). Absence of the characteristic -SH peak at 2600 cm⁻¹ as determined via FTIR analysis led the group to assert that the -SH group is not only responsible for the connection to the UCNPs but the impartation of hydrophilic properties as well. The deprotonation and subsequent negative charge exhibited by the now surface residing carboxyl moieties was used to conjugate Ag⁺ ions to the surface of the UCNP, forming an Ag⁺ ionic shell. Addition of the Ag⁺ shell to the nanoparticles granted an increase in temperature when the UCNPs were excited at 980 nm, compared to UCNPs without the Ag⁺ shell present on the surface. Ag⁺-coated UCNPs in solution reached an increased temperature of 303 K compared to 293 K from uncoated and irradiated UCNPs at the same power density range. Further tests led to the determination of significantly reduced cell viability of both HepG2 and BCap-37 cells, whereby after 20 min of irradiation viability reached as low as 4.62% and 5.43%, respectively.
Similarly, a novel method for the synthesis of sub-10 nm Ln\textsuperscript{3+}-La\textsubscript{2}F\textsubscript{2}/Ln\textsuperscript{3+}-SrF\textsubscript{2} UCNPs was developed by Chen et al. (2011) \cite{61}, where Ln\textsuperscript{3+} = La – Lu. Synthesis of Tm\textsuperscript{3+}, Yb\textsuperscript{3+}-BaF\textsubscript{2}/Ln\textsuperscript{3+}-SrF\textsubscript{2} active-core/active-shell UCNPs was followed by exchange of the terminal oleate ligands with TGA for the sole purpose of aqueous dispersal. Again, the thiol moiety co-ordinates to the surface of the UCNPs whilst the terminal carboxyl moieties reside on the surface of the UCNPs can grant further bioconjugation abilities. Design of an active-core/active-shell provides NIR-NIR dual-mode luminescence due to the upconverting emission of Tm\textsuperscript{3+} at 802 nm upon excitation of Yb\textsuperscript{3+}, as well as the NIR downconverting emissions both of Yb\textsuperscript{3+} at 975 nm and Nd\textsuperscript{3+} at 1054 nm. Furthermore, shell encapsulation of the core allows for protection of core ions from external interaction such as solvents and ligands that may cause vibrational deactivation.

Lastly, related work published by Kumar et al. (2009) \cite{62} also establishes the use of the thiol moiety as a co-ordinating moiety to the UCNP surface for a combined optical and magnetic resonance imaging application. After the synthesis of Na\textsubscript{3}YF\textsubscript{4}:Gd\textsuperscript{3+}, Eu\textsuperscript{3+}, Er\textsuperscript{3+} UCNPs, as-present oleate ligands were exchanged with 3-mercaptopropionic acid (MPA), a ligand that has both a thiol and carboxyl moiety incorporated into its structure. The thiol moiety of MPA co-ordinates to the surface of the UCNPs whilst the residual carboxyl moieties provide further functionality by means of bioconjugation. Various tumour-specific antibodies were conjugated to the nanoparticles, such as anti-mesothelin, anti-claudin 4, transferrin via carbodiimide chemistry between terminal amino groups on the antibodies and the surface carboxyl groups of the UCNPs. Confocal images of Panc 1 cells confirmed substantial enhancement of cell uptake for antibody modified UCNPs compared to UCNPs without antibody modification, a result of a receptor-mediated process. Furthermore, it has been reported that corresponding receptors to the antibodies are overexpressed on the surface of these particular cells \cite{63} and as a result, a high amount of UCNPs can be seen targeting and binding specifically to the receptors on the surface of the cells.

2.2. Silanisation of UCNPs

Silanisation, a popular method for UCNP surface modification due to well-established surface chemistry \cite{3}, allows for the controlled growth and simultaneous functionalisation of a silica shell on UCNPs. Dense (dSiO\textsubscript{2}) \cite{18,64,65} and mesoporous (mSiO\textsubscript{2}) \cite{11,15,25,37,39,66,67} silica shells are common surface modifications to UCNPs due to enhanced bio-compatibility and stability within hydrophilic mediums, such as biological fluids, and present minimal influence on luminescence \cite{5}. Encapsulation of nanoparticles in a silica shell coating allows them to be protected from surrounding aqueous media, thus enhancing both the dissolution and denaturation stability of the nanoparticles and preventing their dissolution into individual ionic species \cite{4,6,7}. This aids in avoiding the release of toxic ions comprising the nanoparticles and a substantially decreased likelihood of an anaphylactic response. Furthermore, the chemically inert nature of silica (SiO\textsubscript{2}) ensures that no degradation of the shell will occur whilst terminal hydroxyl (–OH) groups on the surface of the shell allow for subsequent
functionalisation (–COOH, –NH₂ or –SH) and/or conjugation of biomolecules that can be used for bio-sensing as well as drug and gene delivery applications [4–6,35,36,40,65].

The option to simultaneously functionalise the silica shell during synthesis or post-synthesis can be done easily with silanes that typically undergo a further condensation reaction at the terminal hydroxyls present on the silica shell surface. Examples of such silanes include (3-aminopropyl)trimethoxysilane [4,5,36,37,65] (APTES) which imparts amine (–NH₂) functionalisation, carboxyethylsilanetriol [4,5,17,68] (CEST) that provides carboxylic (–COOH) functionalisation and (3-mercaptopropyl)trimethoxysilane [37,69] (MPTMS) which grants thiol (–SH) functionalisation. Amine-terminated functional groups are the most common silica surface functionalisation, followed behind by carboxyl functionalisation [3–5,29,36]. This is due to the hydrophilic nature of these moieties and, as previously stated, the established coupling process between these functional groups and many biomolecules via carbodiimide chemistry that links amines and carboxylics to form an amide bond [3–5,7,34]. One of the most notable advantages of silanisation is ability to tune the thickness of the silica shell, a variable that is dependent on the concentration of the selected silyl ether. This is particularly useful in both the realisation of drug delivery systems and bioassays where the thickness of the shell can be optimised for drug loading as well as luminescent resonance energy transfer (LRET) mechanisms where by luminescent energy emitted by the UCNPs is commonly transferred to a bio-conjugated compound or biomolecule [3,19,24,25,29,30].

2.2.1. Carboxyl Moiety Modified UCNPs via Silanisation

Functionalisation of silica with carboxyl moieties is a common approach for the preparation of UCNPs for various biomedical operations. The negatively-charged electrostatic interactions they exhibit may be attractive in use. Furthermore, their use in forming amide bonds with biomolecules through well-established carbodiimide chemistry makes them desirable candidates. A summary of the surface modified UCNPs via silanisation with carboxyl moieties can be seen in Table 4.

<table>
<thead>
<tr>
<th>UCNP Type</th>
<th>Silica Type</th>
<th>Functional Moiety</th>
<th>Method</th>
<th>Application</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>NaYF₄: Yb³⁺, Er³⁺</td>
<td>dSiO₂</td>
<td>Carboxylic Acid</td>
<td>Electrostatic Interactions</td>
<td>RCA-120 functionalised UCNPs for biolabeling</td>
<td>[17]</td>
</tr>
<tr>
<td>NaYF₄: Yb³⁺, Er³⁺ and NaGdF₄: Yb³⁺, Er³⁺</td>
<td>dSiO₂</td>
<td>Carboxylic Acid/Amino Acid</td>
<td>EDC/NHS Chemistry</td>
<td>Electrophoretic separation and purification of UCNPs@dSiO₂</td>
<td>[70]</td>
</tr>
<tr>
<td>NaYF₄: Yb³⁺, Er³⁺</td>
<td>dSiO₂</td>
<td>Carboxylic Acid</td>
<td>EDC/NHS Chemistry</td>
<td>Highly sensitive DNA-based biosensor</td>
<td>[20]</td>
</tr>
<tr>
<td>NaYF₄: Yb³⁺, Er³⁺</td>
<td>dSiO₂</td>
<td>Carboxylic Acid</td>
<td>EDC/NHS Chemistry</td>
<td>Oligonucleotide-based biosensor</td>
<td>[22]</td>
</tr>
</tbody>
</table>

A versatile system for biolabeling applications was developed by Liu et al. (2012) [17] following the dense silanisation of NaYF₄: Yb³⁺, Er³⁺ UCNPs. As-synthesised UCNPs with residing oleate capping agents underwent a reverse microemulsion reaction with TEOS to provide shell thicknesses between 1.5 nm and 6 nm. Silica shells not only provide hydrophilicity to the UCNP but also allow for protection of the UCNP nanocore via encapsulation, as well as granting further functionalisation capabilities by interaction with residual surface silanol groups. Following the coating of silica, addition of CEST to the UCNP solution led to terminal triol groups undergoing condensation with the surface silanols, a result of ammonia-assisted hydrolysation. Ultimately, a robust ether bond is formed between the UCNPs and CEST, therefore, affixing terminal carboxyl groups to the surface of the UCNPs. Furthermore, the effects of pH on the carboxyl moieties were investigated (displayed in Figure 8), demonstrating that while the particles displayed excellent colloidal stability at alkaline pH, an increase in particle aggregation (correlating with particle size) is seen at neutral pH. Although this data demonstrates the ability to store the nanoparticles long-term in alkaline solutions, it also
highlights the greatest obstacle in the field, which is the difficulty in obtaining monodispersed and un-agglomerated nanoparticles for use at a biological pH. This behaviour can be explained by the protonation of the terminal carboxylate groups (and an increase in the concentration of protons) at lower pH levels. The addition of a proton to the carboxylate neutralises the formal negative charge held by the hydroxyl group and, thus, reduces the previous amount of repulsion between the negatively charged carboxylate and silica shell, leading to an increase in agglomeration (and hence greater particle size). To demonstrate the biolabeling and LRET application of the UCNPs, the group conjugated Ricinis Communis Agglutinin (RCA 120) to the UCNPs so as to provide interaction with HeLa cells. Upconversion luminescence microscopic imaging illustrated that no upconversion was observed when UCNPs@dSiO$_2$-COOH were incubated with HeLa cells whilst UCNPs coupled with RCA-120 demonstrated significant upconversion, suggesting the RCA-120 successfully targeted and interacted with the HeLa cells.

![Figure 8. Zeta-potential and DLS size distribution as a function of the solution pH value. Reprinted with permission from [17]. Copyright 2013 Royal Society of Chemistry.](image_url)

The use of agarose gel electrophoresis for the detection, separation and purification of various UCNS based on their surface charge was reviewed by Hlaváček et al. (2014) [70]. Various UCNPs (NaGdF$_4$ and NaYF$_4$: Yb$^{3+}$, Er$^{3+}$) were coated with dSiO$_2$ and subsequently functionalised with either APTES or CEST (as displayed in Figure 9). Following this, bare UCNPs@dSiO$_2$, UCNPs@dSiO$_2$-COOH and UCNPs@dSiO$_2$-NH$_2$ were separated onto vertical agarose gel whereby electrophoresis was carried out for 30 mins at 100 V. Separation of particles can be done size-dependently due to both the terminal silanol groups (pK$_a$ = 7.0) and carboxyl groups (pK$_a$ = 4.5) providing a pervasive negative charge across the surface of the UCNPs. UCNPs exhibiting bare silica shells demonstrated a longer diffuse zone than carboxylated UCNPs, with bare silica particle being observed to form several distinct bands. Furthermore, a fraction of the bare particles did not entirely the agarose gel with the formation of several bands indicates varying clumps of agglomerated nanoparticles. Conversely, UCNPs functionalised with terminal carboxyl groups exhibited only a single band when separated under electrophoresis, indicating monodispersed and unagglomerated particles. Moreover, it was determined that fluorescently doping the UCNPs for greater characterisation did not affect the overall surface charge, size or agglomeration of particles. Following electrophoretic separation, the terminal carboxyl groups were inked to terminal amino groups present on the bovine serum albumin (BSA) via carbodiimide chemistry. To further test the separation abilities of the particles via electrophoresis, the newly bioconjugated particles were added to agarose gel under electrophoresis. Although the presence of the UCNPs was well-defined in a single band, the electrophoretic mobility of the particles...
was substantially reduced following bioconjugation. Separation via electrophoresis not only led to single band separation of conjugated UCNPs but of excess BSA as well, thus successfully allowing the separation of conjugated UCNPs from left over BSA reagent.

Figure 9. (a) Schematic drawing of silica-coated UCNPs. UCNPs were coated with either a (b) bare or (c) carboxylated (black dots) silica shell. In parallel approaches, fluorescein (green) was added to the (d) bare or (e) carboxylated silica shell. Reprinted with permission from [70]. Copyright 2014 American Chemical Society.

Alonso-Cristobal et al. (2015) [20] developed carboxyl-functionalised UCNPs@dSiO$_2$ as a platform for a highly sensitive DNA-based sensor (displayed in Figure 10). Following the synthesis of NaYF$_4$: Yb$^{3+}$, Er$^{3+}$ UCNPs, the particles were subsequently coated with a dSiO$_2$ shell via reverse microemulsion. Additionally, the surface silanols groups were condensed with APTES to provide terminal amine functionality and hydrophilicity. However, the terminal amine groups were subsequently subjected to a reaction with succinic anhydride, leading to a ring open reaction that results in formation of an amide bond and affixation of terminal carboxyl groups to the silica shell as shown in Figure 10. Not only does the conversion to carboxyl moieties provide greater hydrophilicity to the UCNPs but it provides the capability to bioconjugate ss-DNA covalently through its terminal amino group. By using carbodiimide chemistry, the terminal carboxyl groups were used to consequently form an amido bond between the UCNP and ss-DNA. As ss-DNA can undergo hybridisation with complementary strands of DNA, complementary DNA attached to graphene oxide (GO) was added to a solution of UCNP@dSiO$_2$@ss-DNA. Hybridisation of the DNA leads to double stranded DNA (ds-DNA) that does not actively interact with the attached GO and therefore allows emission from the UCNPs to be observed once excited at 980 nm. Conversely, non-complementary DNA attached to GO will not undergo hybridisation with ss-DNA present on the UCNP and therefore will interact with the GO. As the absorption spectrum of GO completely overlaps with that of the emission spectrum of the UCNPs, the emission of the UCNPs is quenched via an LRET process. The system described here permits the realisation of a highly selective and sensitive DNA-based biosensor that can qualitatively detect DNA through a turn “on”/turn “off” mechanism.
In a similar manner, Mendez-Gonzalez et al. (2017) [22] functionalised NaYF₄: Yb³⁺, Er³⁺ UCNPs for their use in an oligonucleotide-based sensor (shown in Figure 11). The particles were coated with a dSiO₂ shell where they were then subsequently functionalised with APTES to provide hydrophilicity to the UCNPs. Following this, the residual amino groups were reacted with succinic anhydride to afford terminal carboxyl terminals to the surface of the silica shell via a ring opening reaction as previously mentioned. As a result, the formation of a newly formed amide bond between the terminal amines on the UCNP and the anhydride give rise to carboxyl moieties on the surface of the UCNPs. Although the carboxyl moieties provide greater hydrophilicity than the amine groups, their main role is to provide cross-linking capabilities. They granted the opportunity to cross-link azide-modified ss-DNA to the surface carboxyl groups via carbodiimide chemistry interaction with terminal amino groups present on the DNA. As a consequence, the now terminal azide group present on the surface of the UCNPs affords the ability to undergo highly selective and sensitive click chemistry reactions. The double bonds present on the terminal azide groups can undergo 1,3-dipolar cycloaddition with the alkyne bonds present on the dibenzocyclooctyne (DBCO), which has been modified onto an end of biotin-modified complementary ss-DNA. After addition of the DBCO-ssDNA-biotin into a solution containing the carboxyl-modified UCNPs, the UCNPs were selectively trapped on a streptavidin-coated surface via click chemistry interaction. Consequently, upconversion emission intensity, which was proportional to the target concentration present in the sample, was emitted on UCNP excitation at 980 nm. Not only can this system be used as a screening device to detect both RNA and DNA oligonucleotides but the group also determined a substantially low detection limit of around 1 × 10⁻¹⁷ moles (100 fM).

2.2.2. Amino Moiety Modified UCNPs via Silanisation

Surface functionalisation of silane modified UCNPs with amino moieties can be considered to be the most prevalent of the base functionalisations for silica modifications. As aforementioned, this is due to their ease of involvement with familiar carboxidiimide chemistry, the additional hydrophilicity that they impart and the positive charge that they can exhibit. A summary of surface modified UCNPs via silanisation with amino moieties can be seen in Table 5.
Figure 11. An experimental illustration for the synthesis and functionalisation of UCNPs by Mendez-Gonzalez et al. for a LRET-based oligonucleotide sensor. Reprinted with permission from [22]. Copyright 2017 American Chemical Society.

Table 5. Summary of the surface modified UCNPs via silanisation with amino moieties.

<table>
<thead>
<tr>
<th>UCNP</th>
<th>Silica Type</th>
<th>Functional Moiety</th>
<th>Method</th>
<th>Application</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>NaYF₄: Yb³⁺, Er³⁺</td>
<td>dSiO₂</td>
<td>Amino/Carboxylic Acid</td>
<td>Electrostatic Interactions</td>
<td>Investigation of the effect of surface charge on cellular internalisation</td>
<td>[71]</td>
</tr>
<tr>
<td>NaYF₄: Yb³⁺, Tm³⁺</td>
<td>dSiO₂</td>
<td>Amino</td>
<td>EDC/NHS Chemistry</td>
<td>Photocaged and controlled chemotherapy System</td>
<td>[72]</td>
</tr>
<tr>
<td>NaYF₄: Yb³⁺, Tm³⁺@NaYF₄</td>
<td>mSiO₂</td>
<td>Amino</td>
<td>EDC/NHS Chemistry</td>
<td>Azobenzene-gated remotely triggered drug delivery system</td>
<td>[24]</td>
</tr>
<tr>
<td>NaYF₄: Yb³⁺, Er³⁺</td>
<td>mSiO₂</td>
<td>Amino</td>
<td>Electrostatic Interactions</td>
<td>Targeted delivery of siRNA</td>
<td>[73]</td>
</tr>
</tbody>
</table>

A unique research angle taken by Zhang et al. (2014) [71] investigated the surface charge of dSiO₂ coated NaYF₄: Yb³⁺, Er³⁺ UCNPs and their effect on both cellular interactions as well as cytotoxicity. The dSiO₂ was functionalised with terminal surface carboxyls via the use of carboxyethylsilanetriol (CEST), which allowed for conjugation of NH₂-PEG-NH₂ ligands by the use of carbodiimide chemistry involving sulfo-NHS. The formation of an amide bond resulted in UCNPs@dSiO₂@PEG-NH₂ particles that exhibit terminal amino groups, undergoing protonation at physiological pH and resulting in positively charged surface moieties. The cellular internalisation and uptake of the UCNPs was measured via luminescence and inductively-coupled plasma mass-spectrometry (ICP-MS). Upconversion luminescent intensity was found to be highest for UCNPs@dSiO₂@PEG-NH₂ followed
by UCNPs@dSiO$_2$-COOH when imaged after incubation with HepG-2 and HeLa cells, with measurements conducted via ICP-MS confirming the uptake efficiency of the particles. Further testing through the use of a conventional MTT assay determined the amine terminated nanoparticles to have substantially greater cytotoxicity than the carboxyl terminated particles. Moreover, ICP-MS measurements determined UCNPs@dSiO$_2$@PEG-NH$_2$ to release greater intracellular amounts of Y ions than UCNPs@dSiO$_2$-COOH, as a response to intracellular ROS. Overall, it was determined that a net positive surface charge account for greater cell internalisation of particles and, thus, a greater cytotoxicity than those with a net negative surface charge, for HepG-2 and HeLa cells.

Chien et al. (2013) [72] showcases the development of a targeted and photocontrolled caged chemotherapy UCNP-based system (illustrated in Figure 12) where NaYF$_4$: Yb$^{3+}$, Tm$^{3+}$ UCNPs are coated by TEOS to provide a dSiO$_2$ shell. They were subsequently functionalised with APTES, thereby imparting terminal amino groups to the surface of the UCNPs and greater hydrophilicity. Successful functionalisation of the amino groups to the surface of the silica was confirmed through an additional coupling of N-succinimidyl 3(2-pyridyldithio)-propionate (SPDP) to the amine groups, which can then be cleaved through reduction of its disulphide bond by dithiothreitol (DTT). The cleavage results in the release of pyridine-2-thione (P2T) and demonstrates absorbance at 343 nm that can be used to quantify the release of the P2T and subsequently the successful functionalisation as well as occupation of the amino groups. DOX molecules were thiolated followed by consequent conjugation to the surface residual SPDP molecules where release of DOX molecules can be monitored and quantified via consequent P2T release absorbance at 343 nm as well as fluorescent emission of free-DOX in the supernatant, at 555 nm. It was determined that ~2/3 of the amino groups were occupied by SPDP, allowing for further conjugation of remaining residual surface groups with PEG molecules via carbodiimide chemistry involving NHS. As PEG exhibits terminal carboxyl groups, they were used to further cross-link folic acid (FA) via the terminal -NH$_2$ FA groups by a subsequent carbodiimide reaction. The targeting capabilities of FA coupled with lysosomal enzymes and their ability to cleave the SPDP disulphide bond allows for the realisation of a targeted chemotherapy delivery method. Moreover, the group demonstrated further functionalisation between the terminal FA-COOH groups and 2-nitrobenzylamine (NBA), a photocaging molecule via another carbodiimide reaction. The photocaging functionality displayed by NBA can allow for the remote triggering of NBA removal and subsequent uptake by desired cells in a localised region. Although this design is impressive in approach, both inefficient endocytosis and folate receptor expression levels inhibit the overall release of doxorubicin when used in application.

Liu et al. (2013) [24] examined the controlled release of doxorubicin from NaYF$_4$: Yb$^{3+}$, Tm$^{3+}@$ NaYF$_4$ core shell UCNPs that were functionalised with azobenzene-gated mesoporous silica (as demonstrated in Figure 13). After the coating of mSiO$_2$, it is possible to separately functionalise the surface of the silica shell as well as the pores of the silica. Consequently, APTES was modified with 4-(phenyl)azobenzoyl chloride via the terminal APTES amino group followed by condensation of the resultant N-(3-triethoxysilyl)propyl-4-phenylazobenzamide with terminal hydroxyl groups present inside the mSiO$_2$ pores. The surface of the silica shell was functionalised with APTES, imparting terminal amino groups that were then used to cross-link TAT peptides via carbodiimide chemistry, for enhanced cellular uptake. Following the loading of DOX, it was determined that the positive zeta potential (+2.2 mV) of doxorubicin allowed for strong attachment to the inside of mesopores due to the negative zeta potential (−40.6 mV) exhibited by the silica, leading to strong charge interactions and hydrogen bond formations with residual terminal silanols on the silica surface. Advantageously, this charge-charge interaction allows for doxorubicin to be transported throughout aqueous mediums with minimal to no loss of the drug from the pores unless remotely triggered via conformations in the azobenzene molecule (Figure 13). These are a result of LRET mechanisms arising from both UV and visible light emissions of the UCNP after excitation at 980 nm, where the absorption range of the azobenzene molecule (330 nm–440 nm) partially overlaps with the emission range of the UCNPs (350 nm–450 nm). Furthermore, it is demonstrated that visible light emission resulted in a
trans conformation of the molecule whilst conversely UV light emission result in a cis conformation, thus resulting in a wagging motion that pushes out the loaded DOX molecules in a localised region.

![Image of photocaged UCNPs following NIR laser activation to remove cage molecules and subsequent targeting of cancer cells.](image)

**Figure 12.** Illustration of photocaged UCNPs following NIR laser activation to remove cage molecules and subsequent targeting of cancer cells. Reprinted with permission from [72]. Copyright 2013 American Chemical Society.

Although the following research does not directly involve the mesoporous encapsulation of UCNPs, the same principles can easily be applied to mSiO₂ coated UCNPs for drug delivery applications. Manzano et al. (2008) [74] investigated the effect of amine functionalised mesoporous silica on the controlled drug delivery of ibuprofen. They determined the amine functionalisation to
have increased drug loading capacity by ~10% (due to conjugation of drug to surface of particles as well as pores) as well as resulting in a controlled, slow release of ibuprofen in comparison to non-functionalised, irregular shaped mesopores. This is due to the interactions between the positively protonated $-\text{NH}_3^+$ groups within the pores and negatively-charged $-\text{COOH}$ moiety present on the ibuprofen. Further research conducted by Basaldella and Legnoverde (2010) [75] determined the effects on sulphonic and amine functionalised mesoporous silica in the delivery of cephalixin, which exhibited a two-step prolife: (i) fast delivery (<5 h) and (ii) slow release (>30 h). Similarly, they demonstrated that the slower rate of delivery (>30 h) was attributed to the presence of terminal amines interacting with the $-\text{COOH}$ moiety present on cephalixin. Contrastingly, mesoporous silica spheres functionalised with terminal sulphonic acid led to a fast drug release rate (<5 h) due to a lack of interaction between cephalixin and the sulphonic acid, regardless of the amino moiety present on cephalixin. The extensive research in mesoporous aided drug-delivery displays the tremendous potential in individualised delivery methods for both specific drug molecules and treatments, with additional diagnostic capabilities if coupled with UCNPs.

Lastly, Jiang, S. et al. (2009) [73] demonstrate coated NaYF$_4$: Yb$^{3+}$, Er$^{3+}$ UCNPs with mSiO$_2$ for their use in the targeted delivery of small interfering RNA (siRNA) to cancer cells. The surface silanols groups exhibited by mSiO$_2$ were used in a condensation reaction to couple N-[3-(trimethoxysilyl)propyl]ethylenediamine (AEAPTMS) to the UCNPs. Following this, the newly affixed terminal amino groups were used to further functionalise the UCNPs with an Anti-Her2 antibody by reaction of its terminal amino group through carbodiimide chemistry. Addition of the antibody allowed for the targeting of surface Her2 receptors that are overexpressed on SK-BR-3 cells and was coupled to the UCNPs via carbodiimide chemistry. Subsequently, absorption of siRNA into the mesopores of the silica was achieved via incubation with the UCNPs in solution, after which the gene silencing effect of the siRNA was studied via an exogenous luciferase gene expression assay. Down-regulation of the luciferase gene expression was measured to be 45.5% whilst control samples did not illustrate any down-regulation of the expression, allowing for the successful realisation of a targeted siRNA delivery system. Furthermore, an MTT assay was used to assess the cytotoxicity and subsequent cell viability of SK-BR-3 cells, with cell viability being determined to be above 98.6% with a UCNP concentration less than 50 µg mL$^{-1}$, while a decrease to 92.5% when the concentration was increased to 80 µg mL$^{-1}$ was observed.

### 2.2.3. Thiol Moiety Modified UCNPs via Silanisation

Thiol moieties tend to be the least common of the three base functionalisations for silica, perhaps mainly due to the large variety of biomolecules that exhibit terminal amino or carboxyl moieties, therefore allowing easy coupling to the same moieties due to the great extent of which the methodology surrounding carbodiimide chemistry has been covered. Thiols are particularly ideal for applications which involve a thiol-sensitive maleimide moiety that can be interacted with by thiol moieties present on the surface of UCNPs [72,76]. Table 6 summarised the surface modified UCNPs via silanisation with thiol moieties.

<table>
<thead>
<tr>
<th>UCNP</th>
<th>Silica Type</th>
<th>Functional Moiety</th>
<th>Method</th>
<th>Application</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>NaYF$_4$: Yb$^{3+}$, Tm$^{3+}$</td>
<td>dSiO$_2$</td>
<td>Thiol</td>
<td>Thiol-Maleimide “Click” Chemistry</td>
<td>Photocaged UCNPs for remotely triggered cellular transduction</td>
<td>[76]</td>
</tr>
<tr>
<td>NaYF$_4$: Yb$^{3+}$, Tm$^{3+}$/Au-NP</td>
<td>dSiO$_2$</td>
<td>Thiol</td>
<td>Sulfur-Gold Bond Interaction</td>
<td>UCNPs with attenuated X-Ray interaction as a theranostic platform</td>
<td>[77]</td>
</tr>
</tbody>
</table>

Work conducted by Gao et al. (2015) [76] saw the development of a photocaged upconversion based system that could allow for the remote triggering of intracellular signal transduction via the excitation of UCNPs (displayed in Figure 14). Specifically, NaYF$_4$: Yb$^{3+}$, Tm$^{3+}$ were surface modified
with a dSiO$_2$ shell after which they were subsequently functionalised with terminal thiol groups via the addition of MPTMS. The thiol groups provide additional hydrophilicity to the nanoparticles (on top of the hydrophilicity imparted by the silica shell), thus affording them to be compatible with physiological medium as well as providing further functionality pathways. Following this, PEG3000 was grafted onto the dSiO$_2$ surface through interaction of terminal thiol-sensitive maleimido groups exhibited by the PEG molecules with the newly affixed terminal thiol groups.

![Figure 14. Caged PKA/UCNP complex design and the process of upconversion-assisted PKA uncaging. Reprinted with permission from [76]. Copyright 2015 American Chemical Society.](image)

Further conjugation of previously prepared protein kinase A (PKA) and photocaged PKA to the UCNPs was undertaken, whereby the protein was immobilised via electrostatic interactions [78]. The activity of the PKA was determined by measuring the formation of NADH as a result of phosphorylation of the peptide after coupling with lactate dehydrogenase and pyruvate kinase, leading to oxidation. As PKA can unbundle and disintegrate stress fibres in cells, the group demonstrated that the caged PKA UCNPs caused no disintegration of stress fibres in REF52 cells, resolving that the UCNPs were enzymatically inert if they were not intentionally irradiated with UV or NIR light. Furthermore, disintegration of stress fibres was confirmed quantitatively through the staining intensities of nanoparticle regions relative to background stained stress fibres in the cells. This is a particularly interesting application of UCNPs as there have been minimal attempts at designing remotely-activated enzymatic systems in regards to upconversion.

The development of hybrid nanoparticles consisting of both lanthanide-doped UCNPs and gold (Au) nanoparticles was exhibited by Wang et al. (2011) [77], which displayed attenuated X-ray interaction. After synthesis of NaYF$_4$: Yb$^{3+}$, Tm$^{3+}$ UCNPs, the group coated the particles with a dSiO$_2$ shell followed by subsequent functionalisation via MPTMS to provide surface thiol moieties. Due to the particularly sturdy interaction exhibit between Au and thiolsates, the citrate present on the as-synthesised Au-NPs can be replaced by the terminal thiol moieties present on the UCNP silica shell surface. Thus, mixing of the two nanoparticle solutions can lead to effective hybridization between the UCNPs@dSiO$_2$ and Au-NPs, leading to UCNPs@dSiO$_2$@Au-NPs. Furthermore, it is asserted that the thiolate ligands on the particles can aid in the prevention of particle agglomeration through steric driven repulsion, thus imparting greater colloidal stability when the particles are dispersed in aqueous solutions. As Au-NPs exhibit luminescent absorption near ~541 nm, they can be effectively used as an energy acceptor in a system whereby UCNPs act as energy donors via FRET mechanisms. This is due to a close overlap with upconversion emission exhibited by UCNPs at ~541 nm and leads to the quenching of upconversion luminescence. However, it was determined that the UCNPs@dSiO$_2$@Au-NPs still exhibited significant upconversion luminescence in comparison to unhybridised particles. Furthermore, it was demonstrated that the UCNPs@dSiO$_2$@Au-NPs had greater X-ray attenuation than unhybridised Au-NPs at the same concentration. This system can serve as a platform that can allow for the realisation of various diagnostic, as well as therapeutic applications due to the great multi-modal functionality it exhibits.
3. Conclusions

There is a multitude of ways in which it is possible to approach the surface modification of UCNPs to provide various functional moieties for further use in applications such as biosensing, bioimaging, and drug delivery. As discussed, the primary base functionalities (–COOH, NH₂ and -SH) can imparted onto the surfaces of UCNPs through techniques such as ligand exchange and silanisation. These functionalities can be used to coordinate ligands to the surfaces of UCNPs, as well as grant hydrophilic properties to the UCNPs for their use in aqueous mediums, which is a key objective for bio-applications. Moreover, the affixation of these functional moieties provides further capabilities such as bio-conjugation. Amino and carboxyl groups are evidently favoured modifications due to the extensive methodology surrounding carbodiimide chemistry, the popularity of which can be explained by the abundance of terminal carboxyl and amino groups on biomolecules. DNA and protein are common examples of such biomolecules that are actively conjugated to UCNPs for their use in various bio-applications and can be coupled to UCNPs via carbodiimide chemistry. Surface modifications themselves offer functionality, with silica for example providing encapsulation of the nanocore thereby protecting it from unwanted external stimuli. As a result, dissolution and desaturating of the particles are substantially decreased. Mesoporous silica and polymeric ligands can grant the ability to load therapeutic drug cargo or photosensitisers, an advantage over ligands which require external conjugation of drugs or biomolecules that may display unwanted interactions due to their placement on the surface of particles. Furthermore, it allows for complex design around the working mechanisms of the desired application, for example, separate functionalisation of mSiO₂ pores to the surface can award greater control over mechanisms such as drug release times whilst maintaining the ability to specifically target and localise treatment. Although there only a few base functionalities, the way they are utilised in various applications can be extremely diverse given the necessary conditions for the operation. Manipulation of properties via pH changes, temperature and electrostatic characteristics, for example, can allow for these same functional moieties to be used differently and more effectively to achieve either similar or drastically different goals.

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