Gold nanoparticles delivery in mammalian live cells: a critical review

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Abstract

Functional nanomaterials have recently attracted strong interest from the biology community, not only as potential drug delivery vehicles or diagnostic tools, but also as optical nanomaterials. This is illustrated by the explosion of publications in the field with more than 2,000 publications in the last 2 years (4,000 papers since 2000; from ISI Web of Knowledge, ‘nanoparticle and cell’ hit). Such a publication boom in this novel interdisciplinary field has resulted in papers of unequal standard, partly because it is challenging to assemble the required expertise in chemistry, physics, and biology in a single team. As an extreme example, several papers published in physical chemistry journals claim intracellular delivery of nanoparticles, but show pictures of cells that are, to the expert biologist, evidently dead (and therefore permeable). To attain proper cellular applications using nanomaterials, it is critical not only to achieve efficient delivery in healthy cells, but also to control the intracellular availability and the fate of the nanomaterial. This is still an open challenge that will only be met by innovative delivery methods combined with rigorous and quantitative characterization of the uptake and the fate of the nanoparticles. This review mainly focuses on gold nanoparticles and discusses the various approaches to nanoparticle delivery, including surface chemical modifications and several methods used to facilitate cellular uptake and endosomal escape. We will also review the main detection methods and how their optimum use can inform about intracellular localization, efficiency of delivery, and integrity of the surface capping.

Keywords: gold nanoparticles; cell delivery; bionanotechnology; nanomaterials; photothermal microscopy; cell imaging; intracellular fate

Gold nanoparticles have emerged as attractive nanomaterials for biological and biomedical applications because of their physical and chemical properties (1–13) (Fig. 1). The gold core is inert

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and essentially non-toxic to cells (14). The particles absorb and resonantly scatter visible and near-infrared light upon excitation of their surface plasmon oscillation. The plasmon resonance band can be tuned over a wide spectral range by changing intrinsic parameters such as the material (bi-metallic or hybrid particles), the size, or the shape (sphere, rod, cube, triangle, cage, etc.) (15). The light-scattering signal is intense and much brighter than chemical fluorophores and does not photobleach or blink (16). This constitutes an advantage for their applications in single molecule imaging, where the use of dyes, fluorophores, or quantum dots is limited by low signal intensities, complex blinking phenomena, and photobleaching. For particles below 30 nm, the absorption becomes dominant over scattering and can be used for detection by photothermal microscopy (17). Gold nanoparticles with varying core size are prepared by the reduction of gold salts in the presence of stabilizing agents to prevent nanoparticle agglomeration and control growth (18-22). Particle suspensions are also commercially available. Furthermore, gold nanoparticles can be easily functionalized by anchoring thiol linkers in their monolayers. A wide variety of functional bionanomaterials has been obtained, including nanoparticles modified with peptides, proteins, antibodies, oligosaccharides, and nucleic acids (23-29) (for review, see References 6, 12). This allows the nanomaterials to act as multifunctional platforms for both therapeutic and diagnostic purposes (5, 30, 31).

The use of functionalized gold nanoparticles for biological and biomedical applications includes bio-imaging, single molecule tracking, biosensing, drug delivery, transfection, and diagnostic. For example, through proper functionalization, the particles can be engineered to accumulate preferentially in tumor cells using targeting ligands, providing a tool for cancer diagnosis and gene therapy (32). Sensor arrays have been developed to differentiate normal, cancerous, and metastatic cells using the fluorescence quenching properties of gold nanoparticles (30). The interactions and fate of a broad range of functionalized nanoparticles is currently under investigation in a wide diversity of biological models, ranging from whole organisms to tissues to cells in culture, and also to yeast (33) and prokaryote bacteria (34). The needs of intracellular delivery depend on the applications. For cancer cell targeting and killing, the requirement is proper cell recognition and uptake independently of the ultimate localization (e.g. vesicular localization is not a problem in this context). However, for intracellular imaging and sensing, cell recognition is less important, whereas the ultimate intracellular localization of the nanomaterial is crucial and needs to be fully addressed.

Here, we will focus on the delivery and detection of nanoparticles (with a particular emphasis on gold) into mammalian cells. Many biomedical applications will ultimately necessitate a targeted intracellular delivery and availability of the nanomaterial, not only to specific cells, but also to specific subcellular compartments. Currently, the main challenge is to avoid endosomal localization and to control the stability of the functional capping after cellular uptake. This review discusses how the physico-chemical properties of the particles affect cellular uptake and the current approaches under study for intracellular delivery and control of subcellular localization. We will review the different techniques used for gold nanoparticles imaging and quantification.

**Fig. 1.** Properties and potential applications of gold nanoparticles in biology and medicine.

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inside fixed and living cells and the possible methods to probe the integrity of the functional capping inside the cellular compartment.

**Non-specific cellular uptake**

The plasma membrane defines the separation between the interior of a cell and the outside environment. It is semi-permeable and allows free diffusion of small and non-polar molecules. However, bigger objects such as nanomaterials are incapable of crossing the plasma membrane and require uptake mechanisms such as endocytosis (for review, see References (35) and (36)). Most gold nanoparticle bioconjugates are easily taken up by the cells through endocytotic mechanisms, but they remain trapped in endosomal vesicles and are incapable of reaching the cytosol. Although endocytotic uptake is the norm for a broad range of nanomaterials, its efficiency is dependent on the nanoparticle surface chemistry and the physical properties (size and shape) of the material.

**Surface chemistry**

Incubation of nanomaterials with live cells implicates their exposure to cell culture medium often containing serum. There are evidences suggesting that non-specific interaction of serum proteins has a major effect on uptake. Increase in uptake after 12 h incubation in fetal calf serum (FCS) was observed for superparamagnetic iron oxide nanoparticles (SPIONs) covered with a silane layer (37). Significant difference of uptake was observed for different surface chemistry, such as layer terminated with COOH, NH2, and PEG, but this difference essentially vanished when the particles had been exposed to FCS for several hours (37). In another study, an excellent correlation was found between adsorption of albumin and uptake of a series of functionalized polystyrene particles (Fig. 2) (38). In addition, after removing selected components from the serum, the authors concluded that the uptake does not depend on the identity of proteins adsorbed to particle surfaces, but rather on the capacity of the surfaces to bind protein and they propose that an assessment of adsorptive capacity could be used to predict nanoparticle–cell interactions.

Using mixed monolayers containing oligonucleotides on 13 nm gold nanoparticles, Giljohan et al. observed a correlation between oligonucleotide loading and cell uptake. They also observed an increase in size after exposure to serum correlated with the amount of oligonucleotide and concluded that serum proteins play a key role in mediating nanoparticle–cell interaction and uptake of oligonucleotide-capped nanoparticles (39).

One way to prevent non-specific interactions of serum proteins with nanoparticles is surface functionalization with polyethylene glycol (PEG) (40). There is a consensus on the role of PEG to strongly reduce nanomaterial interaction with cells, thereby impairing their intracellular uptake. PEGylation of nanorods was used to reduce non-specific binding of heceptin-coated nanorods, leading to specific recognition of the cancer cell lines presenting the antigen (41). No significant uptake could be obtained for 10 nm gold nanoparticles decorated with PEG 5000, however an increase of more than a factor 10 was obtained by including a small proportion of a signaling peptide in the layer through ligand exchange (42). Nativo et al. found that uptake in HeLa cells of 16 nm nanoparticles prepared by the Turkevich-Frens method was completely stopped if these particles were protected by a ligand shell terminated with tetraethylene glycol (43). Even after prolonged incubation for 24 h or after a 10-fold increase in concentration, no particles were found inside the cells by transmission electron microscopy (TEM), and no gold was detected by atomic emission spectroscopy (AES) (43). Similarly, the uptake of large core-shell polymer beads was reduced by PEGylation (44).

However, two studies seem to contradict the apparent consensus on the effect of PEG. In the first study, Liu et al. reported high uptake of 4.7 nm PEGylated gold nanoparticles in CT26 cells and subsequent X-ray irradiation cell damage (42). One possible explanation resides in the fact that the PEG molecule used is not
thiolated and the nanoparticles may therefore not be functionalized with PEG. In the second study, Gu et al. observed a very slow (plateau after ~30 h) but significant uptake of 3–4 nm gold nanoparticles that had been functionalized with PEG in a two-step functionalization process (45). Even more surprisingly, the paper presents convincing evidence of localization of the nanoparticles in the nucleosomes. We suggest that the end-group of the particular PEG used (amino-propyl) may play a role, but a definite interpretation of this observation will require further investigation.

Surface structure

In 2004, Jackson et al. reported the synthesis of stripy nanoparticles based on scanning tunneling microscopy observation (46). In subsequent papers, a number of unusual properties were claimed for these new materials, including, most recently, their ability to penetrate the plasma membrane without disruption (47). Both the structure and the properties of these particles are, however, a matter of controversy (48).

Size and a few small complications

Size

The synthesis of gold nanoparticles can be controlled to obtain particles in a large range of sizes. This has allowed Chithrani et al. to evaluate nanoparticle entry into HeLa cells for particles in the 14–100 nm size range (49). The authors found that maximum entry is observed for a 50-nm diameter sphere (Fig. 3A). More recently, building on progress in the size-controlled synthesis of silica, Lu et al. have measured in the same cell line how entry of spherical mesoporous silica beads vary in the 30–280 nm size range (50) (Fig. 3B). The latter paper concluded that their results are in agreement with Chithrani et al., but careful observation of the results indicates a more complicated picture. While in the first paper, uptake is measured in number of particles per cell, in the second, it is measured in picogram of materials per cell. If expressed in pg/cell, the results by Chithrani et al. indicate a plateau around 100 nm rather than a peak at 50 nm (Fig. 3C).

The debate on a maximum size for internalization is still open. Some authors report a maximum size for

Fig. 3. Comparison of nanoparticle uptake as a function of size reported by (a) Chithrani et al. (49) and (b) Lu et al. (50). (c) The results of Chithrani et al. are re-plotted with the particle uptake expressed in pg/cell (as in Reference 50) instead of number of particles per cell. (A and C) Adapted with permission from Chithrani et al. (49). Copyright 2006 American Chemical Society. (B) Lu F, Wu SH, Hung Y, Mou CY. Size effect on cell uptake in well-suspended, uniform mesoporous silica nanoparticles. Small 2009; 5: 1408–13. Copyright Wiley-VCH Verlag GmbH & Co. KGaA. Reproduced with permission.
internalization by non-phagocyte cells, e.g. 1 μm latex beads are not taken by mouse melanoma B16 cells (51). However, gold nanowires that have a 200-nm diameter and a length of several micrometers have been observed to be taken up by fibroblast and HeLa cells (52).

Kinetics of entry

One difficulty in comparing various studies is that most papers present a still picture for a given time of incubation. There have been very few comprehensive quantitative studies of the kinetics of interaction and uptake of nanomaterials in cells. According to some reports, equilibrium (or saturation) takes several hours and is size dependent. Rejman et al. present significant changes in the dynamics of uptake for latex beads between 50 and 500 nm in murine melanoma cell line B16-F10. The saturation time increased with size from ~30 min (50–100 nm beads) to several hours (200 nm beads) (51). Hartig et al. present a detailed kinetic analysis of 140 nm positively charged fluorescent polymer beads interactions with endothelial cells (53). They observe extensive cooperativity and do not approach saturation uptake in the range of concentration accessible. At the complete opposite, a report on carbon nanotubes indicates an almost instantaneous equilibrium with the cell having a similar endo- and exocytosis rate that keeps the intracellular concentration constant (54).

Clustering

Another difficulty in comparing various published studies is to know exactly which species are taken up by the cell. Is it single nanoparticles or aggregates formed prior to, or during interaction with the membrane? Salmaso et al. used particles capped with a thermoresponsive polymer to control aggregation. The results showed an increase in uptake upon aggregation (55). Surface presentation has been proposed as an original approach aiming at better controlled delivery conditions (56). Chithrani et al. have reported that single 50 nm transferrin-coated gold nanoparticles are able to enter cells, while 14 nm transferrin-coated nanoparticles were only taken up when clustered in groups of at least six particles (57). However, in our own experiments on the entry of peptide-capped nanoparticles in HeLa cells, we have observed endosomes containing single 10 nm nanoparticles (Fig. 4). A possible explanation for these apparently conflicting results may be the functionalization of the nanoparticles (transferrin versus peptide/PEG) and its effect on uptake mechanism. Real-time single particle imaging will be necessary to better understand this phenomenon and its impact on uptake.

Shape and rigidity

It has been suggested that uptake was higher for gold nanoparticles than for gold nanorods (49). This was based on the observation of enrichment of the number of spheres in endosomes when HeLa cells were exposed to a suspension of particles containing a mixture of rods and a small proportion of spheres. Similar trends were reported in a study of the influence of shape and rigidity on uptake by macrophages using polystyrene particles (58). Polystyrene particles of identical volumes but different shapes were obtained by stretching the particles embedded in a polymer film. The extended flexible (‘worm-like’) particles exhibited negligible phagocytosis when compared to spherical particles (58). However, the same question was also addressed using a series of size-controlled nanohydrogel particles and in these conditions the high aspect ratio particles (d = 150 nm, h = 450 nm) were internalized by HeLa cells approximately four times faster than the more symmetric low aspect ratio particles (d = 200 nm, h = 200 nm). In the latter study, the particles were being dosed at constant particle mass (15 μg/ml) (59).

It is remarkable that after several thousand publications on the interaction of particles with cells, simple questions such as the effect of size or particle chemistry on nanoparticle uptake in cells are not settled. The most likely explanation is that while the question is simple, the answer is not: the effect of size is not independent of other parameters such as surface chemistry, cell line, colloidal stability, non-specific interactions, etc. It is therefore possible that high-throughput approaches with systematic variations of material parameters will be required to reach a better understanding of nanomaterials-cell interactions. Passive uptake of nanomaterials always results in endosomal localization. This is a stringent limitation for biological and biomedical applications and therefore considerable efforts are being made to escape or bypass the endocytic pathway. In the next section, we examine...
facilitated and active delivery routes for nanoparticle entry into cells.

Facilitated delivery
A number of strategies for biomolecule delivery, especially anti-sense oligonucleotides and DNA have been developed (for review, see Reference (60)). They include the use of penetrating peptides, polycationic molecules, liposomes, and methods involving physical transient disruption of the plasma membrane. Most of these methods have been applied to nanomaterials and gold nanoparticles and their efficiency for intracellular delivery and cytoplasmic availability of the cargo will be discussed.

Cell-penetrating peptides
Over the past 15 years, carrier delivery methods have been developed to facilitate the entry of short oligonucleotides, plasmids, peptides, and proteins. Since the discovery of the Antennapedia homeodomain (61), short cationic peptides that are able to translocate the cell membrane without disruption have caused much excitement. These peptides were called cell-penetrating peptides (CPPs) (62, 63). Since then, different CPPs have been described, including Tat, Sweet Arrow peptide, transportan, and polyarginines (for review, see Reference (64)), and their method of internalization has been extensively studied, discussed, and has raised some controversy (65–69). To summarize, CPPs utilize more than one mechanism of endocytosis to translocate through the plasma membrane. Imaging of Tat peptide-conjugated quantum dots in living cells has shown that the peptide-conjugated nanoparticles are entering cells via macropinocytosis. Tracking data demonstrated that the quantum dots-loaded vesicles were actively transported by molecular machines to an asymmetric perinuclear region (70). Although this study was performed with quantum dots, it is likely that the same mechanisms occur with gold nanoparticles. Direct translocation of CPPs through the membrane has also been reported, especially when they are present in higher concentration and in the presence of endocytotic inhibitors (67, 68). CPPs are attractive as a transfection method, because only incubation is needed and no physical disruption and manipulation of the membrane is necessary.

Gold nanoparticles functionalized with CPPs have been prepared and their efficiency for delivery has been evaluated. There is a clear consensus that intracellular uptake is increased by CPPs. However, the subcellular localization and escape from endosomes is inconsistent. Gold nanoparticles (3 nm) functionalized with Tat peptide were shown by electron microscopy to be localized in the nucleus and in the cytoplasm (71). However, other studies using gold nanoparticles functionalized with amphipathic proline-rich peptide (12 nm gold nanoparticles-C-SAP) (72), Tat, or penetratin (10 nm gold nanoparticles) (43) showed the nanoparticles trapped in endosomes. Similar vesicular localization was also observed with Tat-quantum dots (70). The large discrepancy obtained between different studies can be due to the cell lines used, the size of the cargoes (73), the chemical composition of the capping layers (74), and could also originate from artifacts coming from the fixation required for electron microscopy (75). Careful consideration of cell shape and viability must also be taken into account. A number of studies claim intracellular uptake and even nuclear localization, but show pictures of dead/dying cells; such studies are not significant as membrane permeability is a hallmark of dying cells.

Nuclear localization has been observed using nuclear localization sequences (NLS) in combination with CPPs and PEG (43). However, only a few percent of nuclear localization was achieved and most of the gold nanoparticles remained trapped in endosomes. Mandal et al. also observed endosomal localization using gold nanoparticles capped with a monolayer composed of CPP and NLS (76). Successful nuclear localization using microinjection of gold nanoparticles (~20 nm) capped with nucleoplasmin containing a NLS was observed by Feldherr et al. However, no nuclear import was observed with the NLS sequence only (77). Tkachenko et al. have also observed endosomal trapping and no nuclear localization using the same NLS sequence from the SV40 large T antigen with 20 nm gold nanoparticles incubated on Hep2G cells (78, 79). The same team eventually managed to visualize, by video-enhanced color differential interference contrast (VEC-DIC) microscopy, nuclear delivery of gold nanoparticles using an adenoviral fiber protein containing receptor-mediated endocytosis (RME) and NLS sequences. However, the unique cell shown as proof of the successful delivery doesn’t look very healthy and is likely to be engaged in a death program. This re-emphasizes the requirement to assess the cell shape and viability for these types of experiments to avoid artifacts. Taken together, and excluding a number of studies showing nuclear localization in dead cells, no conditions where the majority of particles had a nuclear localization has been achieved using NLS, indicating that endosomal escape constitutes a prerequisite for successful intracellular targeting.

Protein-mediated delivery: use of transferrin
Proteins have also been used as vectors to facilitate delivery of gold nanoparticles. The interaction between transferrin and its receptor has been used as a potential pathway for cellular uptake of drugs and genes (80). Transferrin plays an important role in iron transport for hemoglobin synthesis. It has been successfully applied for enhanced internalization of 20 nm gold nanoparticles.
capped with transferrin (96 nm hydrodynamic radius) as monitored by atomic force microscopy (AFM) (81). The uptake of 25 nm gold particles capped with transferrin was also studied by laser scanning confocal microscopy (82). The subcellular localization is not detailed in these studies, yet the receptor-mediated endocytosis observed by AFM (81) and the spotty fluorescent signal detected by confocal microscopy (82) suggests endosomal localization. In a third study with particles of various sizes and shapes, TEM and fluorescence imaging indicate endosomal localization as well (57).

**Transfection reagents**

Polyethylenimine (PEI) is a polycationic molecule that has been used as a DNA transfection agent due to the charge-based interaction with negatively charged DNA. It has been used to cap gold nanoparticles and enhance siRNA and DNA delivery (83, 84). However, these studies were not focused on the delivery of the gold nanoparticles. Other cationic molecules, such as cationic lipids, have been developed and are commercially available for the purpose of oligonucleotide transfection. These compounds include Fugene-6, Effectene, GeneSHUTTLE, lipofectamine, etc. Again, gold nanoparticles have been used in combination with these compounds to enhance DNA transfection efficiency (85). We have used Fugene-6 for gold nanoparticle delivery, and found that it increased the aggregation of 10 nm CALNN-capped gold nanoparticles (86). Although a high-level cellular uptake was observed, all gold nanoparticles were localized inside endosomes (Fig. 5). Similar results were obtained with the protein delivery reagent BioPorter (86).

**Liposomes**

A liposome is a vesicle made out of a bilayer of phospholipids that envelops a core aqueous compartment. They can be composed of naturally derived phospholipids or surfactant components. They have been developed for drug delivery as they can carry either hydrophilic (in the center) or hydrophobic (in the membrane) molecules. Initially, the suggested mechanism for delivery was their fusion with the plasma membrane to deliver their contents into the cytosol, but it is now accepted that the main delivery mechanism is endocytosis (for review about cationic liposomes-based delivery, see Reference (87)). One of the approaches to achieving gold nanoparticles delivery is to incorporate the nanoparticles inside or on the surface of the liposome. Liposome intracellular uptake occurs via endocytosis leading to the gold cargo delivery inside endosomes (43, 88). Small gold nanoparticles (1.4 nm) have been successfully delivered by liposomes into cancer cells (89). In this case, the delivery efficiency was increased by 1,000 fold for the small gold nanoparticles. Yet, again, the investigation of intracellular distribution clearly showed lysosomal localization (final degrading organelles) within 40 min of incubation (89).

**Bacterial toxins**

The use of toxins of microbial origin constitutes a different approach for intracellular delivery. Fifteen years ago, it was used for oligonucleotides delivery, especially anti-sense oligonucleotides. They allow reversible permeabilization by transiently forming pores in the cell membrane (90-92). Streptolysin-O (SLO) increased by more than 100 times the intracellular delivery of oligodeoxynucleotides in the KY01 myelogenous leukemia cells compared to conditions without SLO (93, 94). More recently, SLO has been successfully used for protein (95) and peptide nucleic acid (PNA) delivery (96). We have therefore tested this method on 10% CALNN-PEG – 90% CALNN functionalized gold nanoparticles (10 and 5 nm), and obtained improvement of cellular uptake (Fig. 6), suggesting that this route has to be further explored.

**Fig. 5.** TEM images of 10 nm CALNN-capped gold nanoparticles internalized with Fugene-6 (15 µL Fugene-6, 15 µM CALNN-capped gold nanoparticles). TEM images clearly show endosomal localization as well as some large aggregates taken up by the cells by macropinocytosis (right panel).
endosome content into the cytosol (98). This toxin loses its activity in the neutral cytosol and has therefore minimal toxicity (99). The use of these toxins is currently not described for nanoparticles delivery, yet they might offer attractive properties for cytoplasmic delivery.

**Active delivery using physical methods**

Several methods for active delivery involving disruption of the cell membrane, physically or chemically, have been tested with nanoparticles.

**Sonoporation**

Sonoporation uses ultrasound to generate transient non-selective pores on the cell membrane and has been exploited as an intracellular drug and gene delivery strategy. The technique is based on ultrasound in conjunction with microbubbles to drive the cargo into the cell by causing violent bubble cavitation and jet streaming, resulting in pore formation that may be followed by rapid annealing if the membrane damage is not too severe. The size of the pores has been estimated to be \( \sim 110 \pm 40 \) nm in a Xenopus laevis oocytes model (102). A combination of ultrasound and microbubbles has been used for chemotherapeutic drug (103), siRNA (104), DNA (105), and peptide (106) delivery. Ultrasound has been used for delivery of 100 nm nanogold-dipalmitoyl phosphatidyl ethanolamine (DPPE) that are not spontaneously internalized by endocytosis (101). It clearly increased intracellular uptake, without causing significant cell death. However, TEM images showed once again a vesicular localization.

**Genegun**

Genegun is a physical means of molecule delivery. It uses a high pressure helium gas burst to accelerate the cargo and deliver it into the cells and was first described by Klein et al. (107). It has primarily been used to transfect cells with DNA or plasmids (in particular in plant cells). Nanoparticles are frequently used as an adjuvant in these studies, but their fate and localization is not reported because the focus is on DNA transfection efficiency. It has even been used *in vivo* for gene delivery into superficial hepatocytes (108). It has recently been used to transfer nanosensors into several adherent cell lines. The cells displayed a good nuclear uptake of the nanosensor according to the fluorescence levels and distribution measured by confocal microscopy, but the viability remained questionable according to the phase contrast images (109).

**Microinjection**

Another mechanical delivery method is microinjection to single cells. It allows control of the delivery dosage and precise timing of delivery. It has been widely used in many research areas, including the transfection of cells refractory to common transfection reagents such as primary cells (for review, see Reference (110)). However, inappropriate manipulation (injection pressure, needle positioning) can lead to cytotoxicity or cellular stress. Surprisingly, only a few papers show micro-injected gold nanoparticles and their intracellular localization. It has been reported with nucleoplasmmin-capped gold nanoparticles in BALB/c 3T3 cells (77). Targeted optical injection of gold nanoparticles has recently been described by Dholakia et al. (111). They have used a combination of optical tweezing and opto-injection to deliver single 100 nm nanoparticles into the nucleus of single mammalian cells. Although the approach is very neat and well controlled, it seems to be technically extremely challenging, requires sophisticated lasers, and is limited to large nanoparticles. Laser irradiation has also been used with 15–30 nm gold nanoparticles to increase membrane permeabilization and nanoparticle delivery without causing cell death (112), but the intracellular distribution was not evaluated.
**Assisted endosomal escape**

The main limitation with all the techniques described above is the ultimate vesicular localization and absence of availability in the cytosol or in the nucleus in a healthy living cell (for a summary, see Table 1). The same limitations have already been discussed for quantum dots (for review, see Reference (113)). In the field of DNA and plasmid delivery, it has previously been shown that lysosomotropic agents, such as chloroquine or sucrose (114), can improve DNA delivery and subsequent exogenous gene expression. Chloroquine is known to induce vesicular disruption by elevating the intravesicular pH of lysosomes and endosomes. We and others have shown that chloroquine induces endosomal disruption and increases the availability and stability of gold nanoparticles inside the cells (85, 86), yet the effects are not massive and the nanoparticles seem to stay in the region of broken endosomes. Maiolo et al. have shown a specific redistribution of fluorescently labeled cell-penetrating peptides from endosomes to cytoplasm using laser illumination. The phototreatment is supposed to open the endosomes without causing cell death for 24 h (115). As yet, this attractive method has not been evaluated for gold nanoparticles.

In the CPP field, the HA2 peptide has been used in combination with penetrating peptides to increase cytoplasmic delivery. HA2 is a 20 amino acid fusogenic peptide from the influenza virus. It is a highly conserved pH-sensitive sequence, which destabilizes plasma membrane at low pH (116) and is therefore expected to escape endosomal compartments (117). It has been used in combination with polyarginine to increase transfection efficiency (118). Gold nanoparticles capped with Tat-HA2 have recently been successfully used for actin filaments labeling with gold nanoparticles and live cell detection of cytoskeleton dynamics (100).

Some strategies have been developed to avoid endosomal uptake, by designing pH sensitive liposomes combined with fusogenic peptides (119) or with the bacterial toxin listerolysin O for DNA-nanoparticles delivery (119). Interestingly, successful mitochondrial targeting of 10 nm gold colloids was achieved using functionalized liposomes containing octa-arginine on the surface, which enter cells via macro-pinocytosis and then fuse with the mitochondrial membrane to deliver the cargo (MITO-Porter) (120).

**Measurements and detection in mammalian cells**

To understand the fate of gold nanoparticles in live cells and their interactions with intracellular molecules or compartments, it is necessary to combine complementary methods. This can be achieved by combining different technologies, such as fluorescence microscopy, electron microscopy, and atomic force microscopy. Table 1 provides a comparison of gold nanoparticles subcellular localization using different methods of cellular delivery.

### Table 1. Comparison of gold nanoparticles subcellular localization using different methods of cellular delivery

| Method         | Gold core size (nm) | Capping                      | Subcellular localization                        | Toxicity | Reference |
|----------------|---------------------|------------------------------|--------------------------------------------------|----------|-----------|
| CPP            | 16                  | 94% PEG, 2% Tat, 2% NLS, 2% penetratin | Cytoplasmic and nuclear                          | Not reported | 43        |
| CPP            | ~20                 | Nuclear localization sequence and receptor-mediated endocytosis peptides | Nucleus                                           | ~5% death | 78        |
| CPP            | 12                  | Sweet arrow peptide          | Endosomal                                        | Not reported | 72        |
| CPP            | 2.8                 | Tat peptide                  | Cytosolic around the mitochondria and in nucleus | Low cytotoxicity below 10 μM | 71; 68    |
| CPP            | 20                  | Biotinylated Tat-HA2, PEG-SH, anti-actin antibodies | Cytoskeleton (cytoplasm)                          | Not reported | 100       |
| PEI            | 4                   | PEI                          | Mainly endosomal and some nuclear localization    | 20–30% death | 83        |
| SLO toxin      | 5 and 10            | 90% CALNN – 10% CALNN-PEG    | Endosomal and cytosolic                          | Toxicity controlled by protocol optimization Shaheen et al., unpublished data |
| Liposomes      | 1.4                 | Phospholipid                 | Lysosomes near the nuclear membrane              | Not reported | 89        |
| Transferrin    | 14 – 100            | Tranferrin                   | Endosomal                                        | Non-toxic | 57; 81    |
| Ultrasound     | 100                 | DDPE                         | Endosomal                                        | Non-toxic | 101       |
| Microinjection | 11 – 32             | Nucleoplasmin                | Nuclear and cytoplasmic                          | Not reported | 77        |

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techniques to obtain a detailed view, which includes quantitative assessment of uptake, nanoparticle intracellular localization, biochemical environment, and the fate of the nanoparticle capping layer.

**Direct measurements of the gold content: TEM, ICP-AES, and ICP-MS**

TEM is widely used for intracellular detection of metal nanoparticles as it allows their direct visualization due to their high electron density. In conventional TEM (as opposed to high resolution TEM), nanoparticles above 5 nm can easily be detected inside cells because of the good contrast provided by the high electron density of metal nanoparticles. It is a powerful technique that permits the visualization of single nanoparticles in cellular compartments and organelles. However, it has rather low throughput since it necessitates time-consuming processing of the samples (cell fixation and resin embedding). It also requires many images taken from a large number of sliced cells to obtain significant results about localization. Many studies reporting TEM experiments show a single representative image, but important statistical information, such as number of nanoparticles found, number of images analyzed, number of different cells imaged, are often missing (for review on quantification of intracellular gold nanoparticles, see Reference 10).

TEM has been used to quantify the number of citrate-coated gold nanoparticles in vesicles after uptake in mammalian cells (49). Here, TEM results have been further corroborated by inductively coupled plasma atomic emission spectroscopy (ICP-AES). This technique enables a count of the number of gold atoms, thereby providing a more precise quantification of the number of gold spheres, though following the same trend as the TEM estimate. As for ICP-AES, inductively coupled plasma mass spectrometry (ICP-MS) gives an elemental analysis and, therefore, an estimate of the number of nanoparticles in the sample can be calculated. It has been successfully used to study the uptake of Gd

**Fluorescence**

To detect gold nanoparticles by fluorescence, two options can be used: a fluorescent label attached to the gold core or a fluorescent gold core.

**Fluorescent molecules**

The use of fluorescent molecules attached to the gold core is the most widely used. However, attention must be drawn to the quenching effect of the gold core. Depending on the distance between the fluorescent label and the gold core (122), the size of the nanoparticle, and the loading of the fluorescent dye on the nanoparticle (123, 124), energy transfer may prevent photons from being emitted (or enhance their emission). The molecule should therefore be far enough from the gold core, and if the fluorescent signal is to report on the localization of the particle, independent proof that the conjugate is intact should be provided. It has been shown that a ligand can be removed on or after cell entry either by ligand exchange (125) or proteolysis (86). The fluorescence can be measured by wild field microscopy (126) or, for a better resolution, by laser scanning confocal microscopy. The latter technique provides an image of a thin section of the sample, thereby providing better localization information. For example, the uptake of 3.5 nm lysine and FITC functionalized gold nanoparticles was investigated in RAW264.7 macrophage cells by laser confocal scanning microscopy (127).

**Fluorescent gold clusters**

Gold clusters have been shown to exhibit intrinsic fluorescence that varies with the size of the cluster (128). Chang’s group reported the synthesis of intrinsically fluorescent gold nanoclusters (AuNC@DHLA) of less than 4 nm. These particles displayed less photo-bleaching than organic fluorophores and successful conjugations with PEG, PEG-biotin, and streptavidin were achieved (129). They have been used to specifically label endogenous biotin of HepG2 fixed cells and, in a preliminary study, unconjugated AuNC@DHLA have been imaged inside live HAEC cells as a ‘proof of principle’ of their potential in live cell imaging (129). Two other studies have reported internalization of fluorescent gold clusters in living cells (130, 131). However, controls in the absence of nanoparticles are needed (131), and in both cases fluorescence microscopy images indicate endosomal localization, although Lin et al. claim nuclear localization.

**Scattering and absorption**

Several recent papers have reviewed the optical properties of metal nanoparticles and their applications in biosensing (2-4). Here, we focus on selected examples of reports that use these techniques in the context of live cells.

**Dark field**

Curry et al. studied the epidermal growth factor receptor (EGFR) by dark-field microspectrometry and 60 nm gold nanoparticles conjugated to anti-EGFR (132). Here, the reflected dark-field imaging allowed visualization in live cells of the receptor-mediated uptake of gold nanoparticles and spectrometry allowed recording of the scattering spectra of gold nanoparticles. Yet, the fact that individual
nanoparticles of this size can be distinguished in a scattering environment like a cell requires further evidences. Anti-EGFR conjugated and unconjugated gold nanoparticles were compared and different distributions of peak wavelengths were found from the scattering data. The unconjugated gold nanoparticles had a red-shifted and broader distribution as compared to the conjugated ones. The red-shift can be explained by the aggregation of the nanoparticles, and indeed TEM images showed that unconjugated gold nanoparticles were in endosomess in close proximity to one another, when the anti-EGFR conjugated particles were isolated. As mentioned in Assisted endosomal escape section, Kumar et al. have used 20 nm gold nanoparticles functionalized with an anti-actin antibody for labeling actin in live cells (100).

The paper does not report single particle detection (and the TEM shows a very high density of labels), but clearly demonstrates the potential of dark field to improve the understanding of nanoparticle interaction with cells. Louit et al. have reported single particle imaging, tracking, and spectroscopy using dark-field microscopy on live cells (133). For this purpose, they used large 80 nm nanoparticles, which allowed overcoming the scattering background of the cells. To extract robust information from single particle or single molecule studies, it is normally necessary to look at hundreds of traces and to carry out rigorous statistical analysis. Louit et al. did not present such analysis, but their paper is a clear proof of principle demonstration of the potential of this technique to follow the interaction and uptake of single large gold nanoparticles in live cells. They suggested that information on the environment of the particle could be deduced from an observed modulation in the scattering spectrum.

**Plasmon coupling**

Detection of biological events at the single molecule level has been used for biosensing by exploiting the distance-dependent scattering properties of the gold nanoparticles. Using these properties, optical contrast between isolated and closely spaced gold nanoparticles can be achieved (differences in peak intensity and wavelength) (134–136). Inducing the close proximity of plasmonic particles in the presence of a specific target can therefore provide a sensor to probe molecular interactions (135). Nanoparticle plasmon resonance coupling has allowed dynamic imaging of growth factor states in living cells (137). A recent application of the plasmon coupling in living cells has been the detection of protease activation at single molecule level. The signal provided by the plasmon rulers based on optical imaging of crown gold nanoparticles allowed continuous monitoring of caspase-3 activity in live cells (138). Other ways of achieving sensor detection in cells is to exploit the fluorescence quenching properties of gold nanoparticles. This has successfully been developed to detect mRNA in living cells using nano-fores (139). Plasmonic resonance energy transfer (PRET) has been developed for ion metal sensing in living HeLa cells (140).

**Raman scattering**

Surface Raman scattering (SERS) has also been used for live cell probing. SERS can fulfill the requirements of dynamic in vitro systems by using low laser powers and short data acquisition times. Raman scattering occurs during collision of photons with molecules. During this process, photons gain or lose energy and this change in energy results in scattered photons. A Raman spectrum is generated by scattering from different molecular vibrations and provides a vibrational fingerprint of a molecule (for a review on SERS principle and applications, see Reference 141). In SERS, Raman scattering signals can be enhanced by several orders of magnitude for molecules in the vicinity of metal nanostructures. For example, gold nanoparticles can amplify the efficiencies of adsorbed molecules by $10^{14}$ to $10^{15}$-fold, allowing spectroscopic detection of single molecules. Applications are multiple and range from chemical probing in intracellular compartments such as pH (141), to subcellular detection of 20 nm gold nanoparticles decorated with large T antigen NLS (142) or in vitro tumor detection using PEGylated SERS nanotags (143). Recently, time-resolved acquisition of the SERS spectra has also allowed observation of the transport of gold nanoparticles through a living cell (144).

**Differential interference contrast**

Tkachenko et al. have studied the uptake of 20 nm gold nanoparticles functionalized with different NLS sequences using VEC-DIC. The principle of the technique seems appealing, as the gold nanoparticles are supposed to appear as colored areas together with the shape and topography of the cells provided by DIC on a single image recorded at video rate (79). However, there is no evidence of the possibility of individual nanoparticles imaging and it is unclear how much uptake is required to obtain a significant signal. More recently, a dual-wavelength DIC microscopy has been developed to image 20–80 nm silver or/and gold nanoparticles, which is based on the wavelength-dependent contrast of metal nanoparticles (145). The technique has also been used to image the entry of Tat-functionalized 40 nm gold nanoparticles in live HeLa cells at video rates (145). The principle relies on comparing two DIC images taken at the same time at two different wavelengths; one at 540 nm corresponding to the maximum contrast wavelength of gold nanoparticles to detect the nanoparticles and the cells, and one at 720 nm as a control image to detect cells only. The DIC contrast is not only depending on the size of the nanoparticles, but also on the complex refractive indexes of metal nanoparticles (145).
**Photothermal microscopy (absorption)**
In 2002, Boyer et al. introduced photothermal imaging of gold nanoparticles (17). It is an absorption-based imaging technique that allows the direct imaging of metal nanoparticles (146, 147), nanotubes (148), and quantum dots (149). The technique relies on the overlay of two laser beams, one that induces a small localized temperature change (~1–2°C) if a nano-absorber is present, the other that detects the eventual change of temperature. It has further been used to image membrane proteins using gold nanoparticles as labels (150). It allowed detection of single small nanoparticles (10 nm) for a long period of time without blinking or photobleaching. The sensitivity of the technique has since been improved to allow imaging of 5 nm gold nanoparticles in living cells and the technique was renamed laser-induced scattering around a nano absorber (LISNA). Single nanoparticle photothermal tracking (SNaPT) has been developed by the same group and is able to achieve the tracking of metal nanoparticles at video rate for long periods of time. They have recorded the lateral movement of a membrane protein AMPA receptor in live neurons via the tracking of 5 nm AMPA receptor-conjugated gold nanoparticles for more than 5 min (151). Photothermal absorption correlation spectroscopy (PhACS), the equivalent of fluorescence correlation spectroscopy (FCS) with a detection based on the absorption of metal nanoparticles, would also allow the measurement of diffusion constants of metal nanoparticles and the hydrodynamic radii of functionalyzed nanoparticles (152), but it has not yet been applied to live cells.

**Optoacoustic and photoacoustic tomography (absorption)**
Optoacoustic tomography (OAT) uses a pulsed laser that produces a change of temperature localized in the vicinity of any absorbing tissue or object. It creates an acoustic wave that is detected and further reconstructed in a 2-D or 3-D image. It has been reported in in vitro experiments with 40 nm gold nanoparticles conjugated to an antibody that targets human SK-BR-3 breast cancer cells (153). Photoacoustic tomography (PAT) is also based on the absorption of a pulsed laser by an absorbing object, which creates a temperature change generating photoacoustic waves (154). Due to the dark-field configuration of its illumination, the extracted signal produces better quality images. PAT has been used with PEGylated gold nanoparticles as a contrast agent for in vivo tumor imaging in mice (155). One advantage of this technique is the ability to image deep tissue and, therefore, has a potential in ex vivo and in vivo cancer diagnostics and therapeutics.

**Integrity of the capping layer**
To go beyond proofs of principle and develop robust applications of bionanotechnology, a critical piece is currently missing: very little is known about the fate of the nanoparticle capping layer inside living cells. This knowledge gap is due partly to the technical difficulty of investigating nanoparticle surface properties after it has been exposed to the complex environment of the cell. One pioneering study in this field was the analysis of the capping composition using laser desorption/ionization mass spectrometry of cell lysates (156). This technique provides a high level of chemical information, but is time consuming and limited to fixed time points. As shown in Fig. 7, we have used real-time fluorescence confocal imaging on single living cells combined with photothermal imaging on fixed cells to simultaneously report on the localization of the gold core and on the integrity of the capping layer (in this case a peptide ligand shell) (86). We demonstrated that peptide cleavage occurs during nanoparticle uptake and we determined that Cathepsin L was the protease responsible for the degradation. This result is of general significance because one third of all human proteins are potentially cleaved at least once by Cathepsin L (86) and most applications of nanomaterials require conjugation with peptides or proteins to encode for specific recognition, targeting, or actuation. The probability that such bionanoconjugates would degrade during internalization is very high and yet has rarely been discussed in the literature.

In summary, there is a range of methods available to probe the amount of nanoparticles internalized, their intracellular localization, their capping stability, and their potential interactions. To progress towards the rational design of nanoparticle-based sensors and actuators, it is now necessary to systematically combine these tools to obtain a comprehensive view of the bionanoconjugates uptake, localization, and biochemical fate.

**Conclusion**
In this review, we have focused on intracellular delivery of nanomaterials. We observe that in spite of the increasing number of published papers, controlling the intracellular delivery, fate, and intracellular localization of nanoparticles remains a major challenge and even simple questions such as the effect of surface charge, chemical functionalization, and size are still controversial. Clearly, one of the major challenges is the endosomal trapping of the nanomaterials taken up. For in vitro applications such as nanoparticle-based imaging, disruptive methods such as temporary permeation by physical or biochemical means can be used. Even with such methods, particles are generally found in endosomes demonstrating the powerful mechanisms that cells use to protect themselves from foreign materials and organisms. Biology, however, also
teaches us that it is possible to escape or bypass the endosomes: some micro-organisms, such as viruses, are able to disrupt the endosomal membrane to reach the cytosol. Thus, there is hope that the challenge can be overcome in a robust way using astutely designed nanomaterials. This will require progress in understanding in real time what happens to the particles and to their coating immediately after incubation in cell medium, during contact with the cellular membrane, and after intracellular uptake. Combining existing and emerging methods of investigations, with strong interdisciplinary inputs from biology, chemistry and physics, we may well achieve ‘virus-like’ nanoparticles in the near future. Other issues will include control of the intracellular mobility, targeting to specific proteins and DNA targets, but it will only be possible to start addressing these challenges seriously once a robust path to the cytosol is found.

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There is no conflict of interest in the present study for any of the authors.
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