Nanoparticles in biomedicine: Let there be intrinsic light

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Abstract

In this commentary we point out that metal oxide and semi-metallic nanoparticles can be tracked by their own fluorescence without need to use dyes and proteins for tagging. The presence of fluorescent agents modifies the chemistry of the sample and that may be responsible for induced changes in the properties being monitored. Therefore, it is highly advantageous if these nanoparticles intrinsic fluorescence can be used for tracking without any additional tags. In this way, there are two essential advantages. One is that there is no induced change of the chemistry or dynamics of these nanoparticles. Second, we point out that their fluorescence differs from the cell’s auto-fluorescence. We believe that this comment which discusses the intrinsic fluorescence of these nanoparticles would be important for researches in biomedicine, nanobiotechnology and nanoscale bioimaging.

Commentary

In the last few years, the study of metal oxide and semi-metallic nanoparticles (NPs) has become a distinctive subject of research. This increase in interest is driven by the numerous applications of these materials in industry, medicine, information technology, energy storage, sensing and many others [1,2]. Most importantly, they are used as an additive to human and animal food and in biomedical applications.

Their fluorescence has been widely investigated within materials science and it also has widespread use in industrial applications. One example is zinc oxide (ZnO), whose fluorescence has been studied for decades. It exhibits a strong emission peak in the UV, due to recombination of free excitons, and broadband emission in the visible region, attributed to defects within the crystallographic structure [3]. Another example is semi-metallic selenium (Se), with the largest worldwide usage in glass manufacturing, due to its red and pink color [4].

The therapeutic properties of these nanoparticles (NPs) have been extensively investigated for many pharmaceutical and biomedical applications. The therapeutic properties of ZnO have been well explored as anti-cancer and antibacterial agents [5,6]. Selenium is a trace element, naturally found in the body. Recent studies point out that Se has the potential to affect human health in many ways, including improving thyroid function, cancer prevention, capture of heavy metals and antibacterial function for use in orthopaedic applications [4].

However, in biomedical research, their intrinsic fluorescence has been rarely used for particle tracking. Biomedical scientists are commonly using commercial confocal microscopes that use dyes and proteins for tagging, usually not aware of these NPs’ autofluorescence. The fluorescent agents, used in commercial confocals, modify the chemistry of the sample and they may be inducing changes in the properties being monitored such as particle transport. Additionally, for tracking purposes single photon sensitive scanning confocal fluorescence imaging is not necessary but it is a big advantage as measurement is not limited by concentration [3].

The main challenge for optical imaging probes is to satisfy all the criteria necessary for in vitro and in vivo applications. These are: biocompatibility, biodegradability, photostability, suitable wavelengths of absorbance, and fluorescence that differs from tissue auto-fluorescence. It is well known that in a biological cell, the presence of components such as collagen and flavins produces fluorescent background signals with the absorption band in the range 300-500 nm and the emission band in the range 400-550 nm. Therefore, it is essential for the NPs to absorb light at wavelengths longer than 500 nm and to emit light at wavelengths longer than 600 nm in order to be effectively detected and tracked in the cellular environment [7]. Additionally, the light emission should be in the near-infrared region (NIR), 700-900 nm, as it penetrates centimeters into tissue, whereas visible light can only travel microns [8].

Organic dyes and fluorescent proteins meet some of these requirements, but they suffer from photobleaching [9]. Quantum dots (QDs) have been investigated extensively due to their brightness but the major obstacle in the clinical use of QDs remains their toxicity [10]. In contrast, optical centres in nanodiamond (ND) are increasingly being viewed as the crucial building blocks for the development of a variety of advanced biotechnologies due to their outstanding photostability, biocompatibility and suitable wavelengths of absorbance and fluorescence, however NDs do not biodegrade [11] (Figure 1).

Previously, intrinsic fluorescence from metal oxide has been reported as unstable, and that was the main obstacle for their usage in bioimaging application [3,12]. Several metal oxide dopants were

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successfully integrated in one compound or coated with some other materials to maintain photostability [13]. However, some recent studies indicate that metal oxide and semi-metallic NPs can be tracked in cells by their own fluorescence, see Figure 1 [14,15]. The fluorescence of NPs was observed in the red to near infrared wavelength range upon excitation with the visible green wavelength. The NPs were successfully internalized in fibroblast cells and did not show any adverse effects on cellular processes when monitored for extended time intervals. In both instances the experiments were performed at body temperature, relying only on NPs’ intrinsic fluorescence. In this way, there are two essential advantages of this methodology. One is that there is no induced change of the chemistry or dynamics of these NPs. Most importantly, their fluorescence differs from the cells’ auto-fluorescence, as long they are excited at wavelength longer than 500 nm (i.e., green or red laser). This combination of properties enables the efficient and safe use of these NPs depending on concentration [14,15]. Furthermore, these NPs are biodegradable making them ideal candidates for short-term bioimaging applications [16,17].

Thus, intrinsic fluorescence of metal-oxide and semi-metallic NPs at bio-compatible wavelengths of excitation and emission, can be observed, without the use of any additional tags. This fluorescence can be used to explore the biomedical properties of these NPs and make them attractive traceable therapeutic and diagnostic agents.

References

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