Nanotechnology applications in veterinary diagnostics and therapeutics

SM Num¹ & NM Useh²*

1. Department of Veterinary Pathology and Microbiology, University of Agriculture, Makurdi, Nigeria
2. Department of Veterinary Pathology, Ahmadu Bello University, Zaria, Nigeria

*Correspondence: Tel.: 2348126945597, E-mail: nicodemus.useh@fulbrightmail.org

Abstract
Livestock is an important component of the economy of most countries of the world, as it provides some foreign exchange earnings for social amenities and general development. Disease has been a major setback in livestock production and new diagnostic and therapeutic approaches have evolved over time to strictly identify and treat diseases of animals for the purpose of increased protein supply for human nutrition. In the search for improved diagnostic methodologies, livestock disease diagnostics and therapeutics have moved from the traditional methods to molecular and currently nanotechnology. In this contribution, the authors identified the importance of nanotechnology in veterinary diagnostics and therapeutics and suggest that nanotechnology should be combined with molecular diagnostics and therapeutics to boost the efficiency in the diagnosis and treatment of animal diseases for improved protein supply and food security.

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Introduction
Nanotechnology applies the nanoscale principles and techniques to understand and transform biosystems (living or non living) which uses biological principles and materials to create new devices and systems integrated from the nanoscale. It is engineering at the molecular (groups of atoms) level. Nanotechnology is not confined to one industry or market. Rather, it is an enabling set of technologies that cross all industry sectors and scientific disciplines. Probably uniquely, it is classified by the size of the materials being developed and used, not by the processes being used or products being produced (Chauhan et al., 2010).

Diseases of livestock such as helminthosis, trypanosomosis, tick and tick-borne diseases, influenza, clostridial infections, just to mention a few have made news headlines all over the world, because of their devastating effects on livestock population and the potential for some of these to be transmitted to human beings (WHO, 1998, 2012; Engles et al., 2002; Thontiravong et al., 2012; Useh et al., 2012). In some cases, it was difficult for several decades, to distinguish animal and human pathogens that caused similar diseases, because of unavailability of sensitive diagnostic tools and protocols (Nagano et al., 2008; Weatherhead & Tweardy, 2012). The National Science and Technology Council of United States of America (2004) defined nanotechnology as “research and development (R&D) aimed at understanding and working with - seeing, measuring and manipulating - matter at the atomic, molecular and supramolecular levels. This correlates to length scales of roughly 1 to 100 nanometers. At this scale, the physical, chemical and biological properties of materials differ fundamentally and often unexpectedly from those of the corresponding bulk materials.” Veterinary nanotechnology has the potential to improve diagnosis and treatment delivery systems, provide new tools for molecular and cellular breeding, identity preservation of animal history from birth to a consumer’s table, the security of animal food products, major impact on animal nutrition scenarios ranging from the diet to nutrient uptake and
utilization, modification of animal waste as expelled from the animal, pathogen detection, and many more (Scott, 2007).

In contemporary veterinary science, if an animal becomes infected with disease, it takes days, weeks, or even months before presence of the disease is detected by whole-organism symptoms. By that time infection may have spread with the need to destroy the entire herd. Nanotechnology operates at the same scale as a virus or a disease-infecting particle, and thus holds the potential for very early detection and eradication. Nanotechnology holds out the possibility that “smart” treatment delivery systems could be activated long before macro symptoms appear. For example, a smart treatment delivery system could be a miniature device implanted in an animal that samples saliva or other body fluid on a regular basis. Long before a fever or other symptoms develop, the integrated sensing, monitoring and controlling system could detect the presence of disease and notify the farmer and veterinarian to activate a targeted treatment delivery system. Smart treatment delivery systems are envisioned for animal systems such as drugs, nutrients, probiotics, nutraceuticals and implantable cell bioreactors (Scott, 2007).

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It is believed that in the near future, nanotechnology will play a leading role in global veterinary practice (Feneque, 2003; Scott, 2005). The use of nanotechnology devices for diagnosis of animal diseases or as animal models for the diagnosis of human diseases is great achievement in the one health initiative. Recent reports suggest the use of quantum dots (QD) for in vivo imaging in small animal models (Bentolila et al., 2009). Functionalized nanoparticles (NPs) conjugated to monoclonal antibodies have been used to rapidly and specifically detect respiratory syncytial virus in vitro and in vivo. The results suggest that functionalized NPs can provide direct, rapid, and sensitive detection of viruses and thereby bridge the gap between current cumbersome virus detection assays and the need for more rapid and sensitive detection of viral agents (Tripp et al., 2007). Several other reports show the usefulness of nanoparticles in disease diagnosis (Na et al., 2009; Jackson et al., 2011; Schlacter et al., 2011; Huang et al., 2012).

Nanotechnology is currently employed in the treatment of African animal trypanosomosis (Kroubi et al., 2010). It enabled the development of a new drug delivery system against diaminazene (DMZ) a trypanocidal drug registered for veterinary use. The porous cationic nanoparticles used improved the potential targeting of trypanosomes. Similarly, mice pre-treated with protein cage nanoparticles (PCN) independent of any specific viral antigens, were protected against both sub-lethal and lethal doses of two different influenza viruses, a mouse-adapted SARS-coronavirus, or mouse pneumovirus. Treatment with PCN significantly increased survival and was marked by enhanced viral clearance, accelerated induction of viral-specific antibody production, and significant decreases in morbidity and lung damage (Wiley et al., 2009). In another report, an aerosol formulation of biodegradable, biocompatible and nontoxic gelatin nanoparticles bound CpG-ODN 2216 was used to treat equine recurrent airway obstruction in a clinical study. Most remarkable was that regulatory anti-inflammatory and anti-allergic cytokine IL-10 expression was significantly triggered by five consecutive inhalations. Thorough assessment of clinical parameters following nanoparticle treatment indicated a partial remission of the allergic condition. The authors concluded that although employed for the first time, the treatment protocol showed the effectiveness of colloidal nanocarrier-mediated immunotherapy in food-producing animals with potential future applicability to other species including human beings (Klier et al., 2012). Nanobiotix technology and its role in cancer therapy is a novel innovation that is already gaining acceptability in diagnoses and therapeutics. It is based on the novel idea of nanotherapeutics, using nanoparticles with control diameter less than 70 nm with a therapeutic core that can be remotely activated by an external energy supply. The nanoparticles are injected into the patient intravenously or intratumoral and target tumor tissues and take 20-48 hours to accumulate selectively in them. Once the particles have been internalized by the cancer cells, an external energy field is applied to activate the nanoparticles and a local physical or chemical effect then destroys the tumor cell (Chauhan et al., 2010).

Other authors determined the efficacy of paclitaxel (Tx)-loaded biodegradable nanoparticles (NPs) on tumor inhibition in a murine model of prostate cancer (Sahoo et al., 2004). They hypothesized that NPs following conjugation to transferrin (Tf) ligand (NPs-Tf) would enhance the therapeutic efficacy of the encapsulated drug. The antiproliferative activity of NPs was determined in human prostate cancer
cell line (PC3) and their effect on tumor inhibition in a murine model of prostate cancer. NPs (approximately 220 nm in diameter, 5.4% w/w drug loading) under in vitro conditions exhibited sustained release of the encapsulated drug (60% release in 60 days). The IC50 (concentration of drug for 50% inhibition of cell growth) of the drug with Tf-conjugated NPs (Tx-NPs-Tf) was about 5-fold lower than that with unconjugated NPs (Tx-NPs) or drug in solution. Animals that received a single-dose intratumoral injection of Tx-NPs-Tf (Tx dose = 24 mg/kg) demonstrated complete tumor regression and greater survival rate than those that received either Tx-NPs or Tx-Cremophor® EL formulation. Overall, sustained release NPs demonstrated greater antitumor activity following their conjugation to Tf ligand.

The immunological properties of a novel nano-bead adjuvant in a sheep (large-animal) model were investigated (Scheerlinck et al., 2006). In contrast to alum, antigen covalently coupled to nano-beads induced substantial cell mediated responses along with moderate humoral responses. No adverse reactions were seen at the site of immunization in the sheep. The authors concluded that nano-bead adjuvants in veterinary species may be useful for the induction of immunity to viral pathogens, where a cell mediated response is required. These findings also highlight the potential usefulness of nano-bead vaccines for intracellular pathogens in human beings.

Vaccination against foot-and-mouth disease virus (FMDV) is a major problem, as current vaccines do not allow easy differentiation between infected and vaccinated animals. Furthermore, large scale production of inactivated virus poses significant risks. To address this, Greenwood et al. (2008) investigated the feasibility of using inert nano-beads that target antigen to dendritic cells (DCs) to induce immune responses against FMDV-specific synthetic peptides in sheep. It was clear that while single peptides induced responses in most sheep, the combination of multiple peptides either conjugated separately to individual nano-beads or conjugated as a mixture induced significant cell-mediated and humoral immune responses.

There are potential hazards associated with the use of nanotechnology and research is still ongoing to mitigate the hazards. For instance, lipopolysaccharide-coated NPs induced lung inflammation which subsequently changed the microenvironment leading to higher translocation rates of NPs to secondary organs (Chen et al., 2006).

Also under high environmental, occupational or chronic exposure, inhaled engineered NPs can enter olfactory and trigeminal nerve in nasal region and sensory nerve network in tracheobronchial region taking their way to the central nervous system (Simko & Mattsson, 2010). Elder & Oberdorster (2006) found that colloidal gold NPs (50 nm) translocated in the axons of the olfactory nerves to the olfactory bulb after instilled internasally in monkeys. Regarding the skin route, there is some evidence that NPs can accumulate around hair follicles and enter the deeper layers when these follicles open during hair growth (Lademann et al., 2006). Moreover, it was shown that quantum dots can penetrate the healthy skin (Ryman-Rasmussen et al., 2006). Nano-scale structures can directly reach the gastrointestinal tract via food, water, drinks, drugs or drug delivery systems. Also, NPs cleared from the respiratory tract via the mucociliary escalator can subsequently be ingested into the gastrointestinal tract. The absorption of NPs from the gastrointestinal tract is governed by both the size (Hillyer & Albrecht, 2001) and surface characteristics (Jani et al., 1989) of the particles. Jani et al. (1990) showed that absorption of 125 iradialabeled polystyrene NPs was found to be size dependent (50 nm > 100 nm) in rats and was mainly confined to the Peyer’s patches of the gut. Most of the studies have demonstrated that NPs undergo limited gastrointestinal absorption and systemic translocation following oral administration (Stem & McNeil, 2008). For example, studies of the oral absorption of 14C-radiolabeled fullerenes and 192Ir NPs in rats observed minimal systemic absorption (Yamago et al., 1995).

In developing countries, especially Nigeria where technology is at the lowest level of development, the use of this technology in veterinary diagnostics and therapeutics is almost near impossible in the near future. Moreover, the dearth of trained personnel that will handle and manipulate nanodevices at the level of the farm setting remains a great challenge indeed.

Conclusion
Although there are gaps that need to be filled, veterinary nanotechnology holds a great key in diagnostics and therapeutics of animal diseases and research is being intensified to breach the gaps. It is suggested that in the future molecular diagnostics and therapeutics should be combined with nanotechnology to boost the efficiency in the diagnosis and treatment of animal diseases for
improved protein supply and food security.

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