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Publisher's version available at: http://dx.doi.org/10.4172/2157-7439.1000363

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Design, Synthesis, Characterization and Toxicity Studies of Poly (*N-Iso*-Propylacrylamide-*co*-Lucifer Yellow) Particles for Drug Delivery Applications

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Nanomedicine & Nanotechnology

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Abstract

Journal of

A novel fluorescent temperature/pH responsive particle was designed, synthesized, characterised and tested for toxicity. Poly (*N-iso*-propylacrylamide-*co*-5%-lucifer yellow) (p(NIPAM-*co*-5% LY)) was prepared using a surfactant free emulsion polymerisation technique. The particles were negatively charged and were approximately 250 nm at 15°C. When the particles de-swell following an increase in temperature, a particle size around 100 nm is obtained. The toxicity of different concentrations of the new particles (p(NIPAM)-*co*-5% LY, as well as the 100% p(NIPAM) and the main monomer NIPAM was tested on two cell lines (Hela and Vero). The toxicity was tested in comparison to a positive control (dextran sugar) and a negative one (poly(ethylenimine)) (PEI). The results show that the two particles show cell viability over 80% (for both cell lines Hela and Vero) up to a concentration of 3 mg/mL while NIPAM monomer showed cell viability over 80% at a concentration equal to or less than 0.3 mg/mL. The fluorescence property of the novel particles make them traceable. Combining this property (tracing) to the ability of the particles to release their content in response to temperature and pH change can be a potential drug delivery system for cancer treatment.

Keywords: Fluorescent p(NIPAM); Lucifer yellow; Toxicity of p(NIPAM); Drug delivery; Diagnostics

Introduction

Poly (N-isopropyl acrylamide) (p(NIPAM)) particles are characterised by their ability to change their conformation and physico-chemical properties (such as particle size and surface charge density) according to the change in the surrounding environment such as temperature, pH and ionic strength [1-5]. This is thought to be a property that can be used in different applications such as drug delivery [6].

Below the volume phase transition temperature (VPTT), the polymer-polymer and polymer-solvent interactions are weak, the interstitial spaces in the nanogel particles are filled with water and the nanogel particles are swollen. Above the VPTT, the polymer-polymer interactions become stronger than the polymer-solvent ones and so the polymer particles tend to collapse (Figure 1). When the temperature is increased, the polymer chains tend to interact with each other rather than with the solvent molecules and the hydrogen bonding between the polymer chains and the solvent molecules is reduced. This causes most of the solvent entrapped within the interstitial spaces of the polymer matrix to be expelled; forming hard like sphere particles and the nanogel tends to shrink. During this collapsed state, the particles are dispersed; this is due to the repulsion between the similar ionic charges on the particle surface caused by the ionic initiators. This process (deswelling) is reversible, so if the temperature goes back below the VPTT, the interactions between the polymer and the solvent improve and the nanogel swells again.

The fact that the volume phase transition temperature (VPTT) of NIPAM based nanoparticles (32-34°C) [3] is very close to the human body temperature (37°C) makes them a good choice for biomedical applications. As an example of suggested drug delivery applications based on temperature trigger, the particles are swollen at room temperature (around 25°C which is below the VPTT), so, if they incorporate a drug in its matrix, when administered in the human body VPTT, it will deswell and release the drug. Many researchers suggested drug delivery [7-9], diagnostics [10] and protein binding [11-13] applications of NIPAM-based nanoparticles.

Nanogels sensitive to both temperature and pH can be prepared by incorporating acidic or basic co-monomers into poly (NIPAM) nanogel matrix. An example of this is the poly (NIPAM)-co-acrylic acid nanogel [1]. Figure 1 shows the effect of temperature and pH on the temperature/pH sensitive p(NIPAM)-co-acrylic acid particles. At temperature lower than the VPTT, the particles take in solvent molecules (water) and hydrogen bonds are formed between them (the particles and the water molecules). When the temperature exceeds the VPTT, the hydrogen bonds are broken and the particles collapse (Figure 1) [1]. The pH conformational transition point depends upon the logarithm of the acid dissociation constant (pK_{a}) of the functional groups incorporated. At pH below the pK_a of acrylic acid, the nanogel particles are collapsed (due to low electrostatic repulsion forces (E.R.F) but when the pH is more than the pK_2 , the E.R.F. between the ionised acrylic acid molecules within the polymer matrix increase causing the gel to acquire a more swollen configuration (Figure 1). In Figure 1, the blue colour represents low temperature (below the VPTT) and high pH (higher than the pK_{a}), the red colour represents high temperature and low pH. Both cases (low temperature/low pH and high temperature/ high pH) are represented as light blue.

The incorporation of a fluorescent dye such as Lucifer yellow (LY) to p(NIPAM) particles adds a new function (colour and fluorescence) that makes it traceable. This suggests new applications especially in the field of diagnosis of different diseases either *in vivo* or *in vitro*.

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Received March 03, 2016; Accepted March 21, 2016; Published April 02, 2016

Citation: Mohsen R, Alexander BD, Richardson SCW, Mitchell JC, Diab AA, et al. (2016) Design, Synthesis, Characterization and Toxicity Studies of Poly (*N-Iso*-Propylacrylamide-co-Lucifer Yellow) Particles for Drug Delivery Applications. J Nanomed Nanotechnol 7: 363. doi:10.4172/2157-7439.1000363

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Combining the previously suggested drug delivery applications of nanogels [7-9] to its ability to being traced in the human body can be a promising system for combined diagnosis and delivery of drugs to suitable environments [14,15].

In this work, a novel nanoparticle, p(NIPAM-co-5% LY), is synthesized and characterized using DLS (particle size and electrophoretic mobility), infrared spectroscopy, scanning electron microscopy and fluorescence microscopy. Being a candidate for biomedical applications, the toxicity was tested using two cell lines: kidney cells (Vero) and breast cancer cells (HeLa).

Lucifer yellow VS

Lucifer yellow VS dilithium is a water soluble fluorescent dye of molecular weight 550.39 g/mol [16]. Due to the presence of 3 benzene rings, it is hydrophobic in nature. It has two excitation wavelengths (430 and 280 nm) and emission wavelength at 540 nm [17]. Lucifer yellow is used for different biomedical applications including marking cells, neurons, albumin, cholesterol, phospholipids, collagenase, polynucleotides, progesterone derivatives, testosterone, estriol [16], and tracers of cell-cell fusion [18]. It is also selectively binds to copper [19] which can be used in industrial and environmental applications.

The logarithm of acid dissociation constant (pK_a) of LY was calculated using the online database ACD/labs [20]. The presence of more than one ionisable group in the molecular structure of LY (Figure 2) results in more than one pK_a , these are; -2.2 (0.6), -0.7 (0.5), 1.4 (0.8) and 5.7 (0.8) where standard deviations are shown between brackets. Figure 2 shows the protonation state of LY versus pH. The most acidic form is coloured red and the most alkaline one is coloured blue. The two negative pK_a values are mainly caused by the protonation/ deprotonation of the sulphonate groups (groups 1 and 2 (coloured in red)) (Figure 2). The first positive pK_a is caused by the protonation/ deprotonation of the nitrogen in the piperidine ring (group 3 (coloured in light blue)) while the second one is caused by the amino of the aniline (group 4 (coloured in blue)) [1]. An overview of the protonation state of lucifer yellow at different pH values is shown in Figure 3 [1].

From polymerization point of view, lucifer yellow VS dilithium salt is a perfect choice. This is due to the presence of a pendant vinyl group (coloured in grey) (Figure 2) which makes it readily polymerizable with other monomers under suitable conditions. This is important for synthesizing monodispersed particles with the similar electrophoretic mobilities. On the other hand, a disadvantage of using lucifer yellow is its relatively low molar extinction coefficient (ε) of 12000 M⁻¹ cm⁻¹ [17,21,22] when compared to other fluorophores such as fluorescein isothiocyanate (FITC) with a molar excitation co-efficient of 75000 M⁻¹ cm⁻¹ [23], texas red (ε = 85000 M⁻¹ cm⁻¹ [24] and tetramethyl rhodamine isothiocyanate (TRITC) with an ε of 100000 M⁻¹ cm⁻¹ [23]. It is also characterised by a relatively low quantum yield (21%) [25] compared to other fluorophores such as rhodamine 6G (95%), fluorescein (95%) and quinine sulphate (58%) [26].

Materials and Methods

Materials

*N-iso*propylacrylamide (NIPAM) 97% (Aldrich), lucifer yellow (LY) (Aldrich), *N*, *N*'-methylenebisacrylamide 99% (Aldrich), potassium persulphate 98% (BDH Laboratory Suppliers), branched poly(ethylenimine) (PEI) of molecular weight 25 kD (Cat. 18197-8) (Aldrich) were obtained from commercial suppliers and used without further purification. Vero CCL-81 (normal kidney of *Cercopithecus aethiops*) and HeLa CCL2 (human cervical epithelial cells) from ATCC company. Media were purchased from Gibco Life Technologies. For Vero cells, Dulbecco's Modified Eagle Medium (DMEM) was used while for HeLa cells, Minimum Essential Medium (MEM) was used. Penicillin-Streptomycin-Glutamine (PSG) (100x) and fetal Bovine Serum (FBS) were purchased from Gibco life technologies. (3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) was purchased from Sigma Aldrich UK.

Methods

Synthesis of p(NIPAM-co-5% LY): In a 1 L reaction vessel, 0.5 g of the initiator (potassium per sulphate) was dissolved in 800 mL of







distilled water. A three-necked lid was clamped to the reaction vessel, which was then heated to 70° C with continuous stirring. Monomer (4.75 g NIPAM), co-monomer (0.25 g lucifer yellow (LY)) and cross linker (0.5 g *N*,*N*'-methylenebisacrylamide) were stirred in distilled water (200 mL) and transferred into the reaction vessel containing the initiator and continuously stirred at 70° C for six hours under an inert atmosphere of nitrogen. When the reaction was complete, the nanogel dispersion was allowed to cool to room temperature. The nanogel was dialyzed in fresh de-ionised water for a week (changed daily), centrifuged and freeze dried.

Dynamic light scattering (DLS): Dynamic light scattering (DLS) measurements were carried out using a Malvern Zetasizer Nano ZS. The hydrodynamic diameter of the particles was measured in response to temperature in the 15 to 60°C range. All samples were diluted with deionized water (1 mL of dialysed microgel dispersion was diluted with 2 mL of water) before measurements. A quartz cuvette with two polished windows (Starna Type 1) was used for all the measurements and the temperature of the dispersion was controlled by a Peltier thermocouple. Data for p(NIPAM-co-5% LY) measurements were collected every 1°C and the samples were equilibrated for 2 min before each data collection point. Three measurements, each consisting of 13 subruns, were taken at each temperature to obtain an average hydrodynamic diameter. Experiments were also conducted where the pH of the samples was adjusted to different pH values using HCl and NaOH to test for the effect of changing pH upon the size of the particles. The electrophoretic mobility of p(NIPAM-co-5% LY) in a salt concentration of 1 x10⁻⁴ M NaCl was measured in response to temperature change across the range of 15-60°C using a disposable folded capillary cell.

Fluorescence spectroscopy: A Horiba Jobin Yvon Fluoromax 4 spectrofluorometer was used for fluorescence experiments. All samples were excited at 430 nm and the full emission spectra (450–1050 nm) were recorded with bandwidth of 5 nm at 25°C. A 10 mm path length quartz cuvette with four polished windows (Starna Type 3) was used for all the measurements.

Infrared spectroscopy (IR): A Thermo Nicolet Nexus FT-IR was used to carry out the IR experiment in this work. Nanoparticle dispersions were dialysed and freeze-dried then the solid samples (powder) were used for IR experiments using attenuated total reflection (ATR) ZnSe crystal.

Microscopy

Scanning electron microscopy: A Hitachi SU8030 cold cathode filled emission gun scanning electron microscope was used to obtain images of the particles.

Fluorescence microscopy: A Nikon Eclipse 90i microscope, fitted with a fluorescein isothiocyanate filter (FITC) and a Nikon digital sight DS-U3 camera were used to take fluorescent images of the particles. Plan Fluor 100x oil Ph3 DLL lens was used with working distance of 0.16 mm. The numerical aperture of the images was 1.3, refractive index 1.5 and exposure time 3 s. the sample was placed on a microscopic slide, left to dry in air and then immersion oil was added on it.

Toxicity studies

Sterilization: All the pipetting tips, phosphate buffered saline (PBS), glassware, magnetic stirrers, spatulas used in this experiment were sterilized used an Astell autoclave at 121°C for 85 minutes.

Preparation of media: To prepare a complete media, 50 mL PSG (100x) and 5 mL Fetal Bovine Serum (FBS) were added (without any

dilution) to a 500 mL media bottle (either DMEM or MEM). Media were kept in the fridge at 4°C.

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Cell sub-culturing: In a clean and sterile class II laminar flow hood, the media in the flask were removed using a disposable sterile 10 mL serological pipette (Fisher brand). The flask was then washed with 10 mL of sterilised phosphate buffered saline (PBS) 3 times (using a new serological pipette each time). Trypsin EDTA (0.5 mL) was added and spread among the flask which is then placed in an incubator (Thermo Scientific HERACELL 150i CO₂ incubator) at 37°C and 5% CO₂ v/v for 5-7 minutes after which the viability of cells was checked under microscope (Nikon Japan with a x10 ELWD lenswith working distance 2.62–1.8 mm). After that, 10 mL of fresh media (MEM for HeLa and DMEM for Vero cells) were added to the flask and well mixed with the cells, and then 9 mL of the media (now mixed with cells) were removed and put in an Eppendorf tube to be used for the seeding step. Fresh media (9 mL) were then added to the flask, mixed then put in the incubator. Cells were split twice weekly.

Cell seeding: Cell count (using a haemocytometer) of the cell suspension from the splitting step (9 mL removed after using trypsin) was carried out. Accordingly, a cell suspension of a concentration (1 × 10^4 cell/mL) was prepared. In a 96-well plate (excluding the peripheral cells, these are columns 1 and 12 and rows A and H (Figure 4)), $100 \,\mu$ L of the freshly prepared cell suspension were placed in each well. The plate was then put in the incubator for 24 hours.

Sample preparation: In an autoclaved beaker, the sample (either dextran, NIPAM monomer, p(NIPAM), p(NIPAM-co-5% LY) or PEI) is weighed. Sterile fresh media (MEM for HeLa and DMEM for Vero cells) was added and left stirring for an hour. In a sterile class II laminar flow hood, the sample dispersion/solution was filtered using a single use filter unit with a pore size of 2 μ m (Minisart[®] Sartorius stedium biotech). A serial dilution was then prepared and used for cell dosing.

Cell dosing: The previously seeded 96-well plates are placed in a clean and sterile fume cupboard, the 100 μ L of media (in each well) are removed using sterile pipette tips. The first two columns are used as control wells (100 μ L of fresh media is put in each well) while the rest of the columns is used for the different concentrations of the sample. In each well, 100 μ L of the pre-prepared sample-media suspension is accurately put using a sterile pipette tip. For each sample, two plates are set up; Figure 4 shows the design of each plate while Table 1 shows the concentrations used in each one. For each cell line (HeLa and Vero), a whole plate was set up to be a control one. Plates are placed in the incubator for 72 hours after which the MTT assay is carried out.



Figure 4: Structure and organisation of the 96- well plates used for toxicity studies. Blue cells were kept empty. The concentrations of the columns are shown in Table 1.

Conc.	С	С	A1	A2	A3	A4	A5	A6	A7	A8
Plate A										
mg/mL	Control	Control	3.00	1.00	0.70	0.30	0.10	0.07	0.03	0.01
Plate B										
µg/mL	Control	Control	7.00	3.00	1.00	0.70	0.30	0.10	0.07	0.03

Table 1: The design of plates A and B and the concentrations used for the toxicity testing of dextran, NIPAM monomer, p(NIPAM), p(NIPAM-co-5% LY) and poly(ethylenimine) (PEI).

The concentrations of the columns are shown in Table 1.

MTT assay: For the preparation of MTT solution, 250 mg of MTT reagent were dissolved in PBS and sterilised at 121°C for 85 minutes. In each well, 10 μL of pre-prepared of MTT solution was added. Plates were then placed in the incubator and left for 4 hours. The media was then removed (this includes 100 μL sample in media and 10 μL MTT) and 100 μL of dimethyl sulphoxide (DMSO) are added in each well and plates are left in the incubator for 30 minutes. The absorbance at 540 nm of the DMSO in each well is then determined using a Thermo Electron Corporation Multiskan Ascent plate reader.

Growth curves: Cells were seeded at a concentration of 1×10^4 cell/mL (as explained above) in a 96-well plate. Every day (for 10 days), MTT assay was carried out (as explained above) on one column of the plate.

Results and Discussion

Particle size

Effect of temperature on the particle size: Figure 5 shows the response of p(NIPAM-co-5% LY) to the change in temperature across a temperature range of 15-60°C both during heating and cooling. The initial particle size at 15°C at pH 7 is 250 nm which is considerably smaller than that of 100% p(NIPAM) with an initial particle size of 550 nm [3,27]. The incorporation of a hydrophobic co-monomer such as lucifer yellow [28] decreases the extent of hydrogen bonding between the particle and water (solvent) compared to 100% p(NIPAM), so, less water will be incorporated in the particle leading to a smaller particle size [29]. It also changes the reaction conditions such as the rate of the reaction, solubility of the monomer and co-monomer and the extent of reaction before termination [30]. The DLS data suggest that LY was successfully incorporated in the new particle. The Figure also shows that the novel fluorescent nanoparticle maintains the reversible temperature responsive behaviour of p(NIPAM) with a VPTT comparable to that of 100% p(NIPAM) which is 35°C [3]. The polydispersity index (PdI) value (Figure 5), which is close to zero (0.04), suggests that the majority of the particles in the dispersion acquire an even size.

Effect of pH on the particle size of p(NIPAM)-*co*- 5% LY: The effect of pH on the hydrodynamic diameter of p(NIPAM-*co*-5% LY) is shown in Figure 6. The Figure shows an increase in the particle size by increasing the pH at the same temperature (25°C). At pH (0.5-3), the particle diameter was 190 nm and increased by 20% to reach 230 nm at pH (5-10). At pH 11, the particle further swells by 9% to reach 248 nm. The change in particle size is thought to be mainly attributed to the change of the protonation state of the co-monomer lucifer yellow by the change in pH (Figure 3).

Several pH sensitive NIPAM based particles (with different comonomers such as acrylic acid) have been studied [3]. At pH lower than the pK_a of the co-monomer, the anionic groups are protonated, accordingly, they are not charged and hence the electrostatic repulsion force is minimal. When the pH increases above the pK_a of the comonomer, the anionic co-monomer groups are deprotonated, creating more surface charge on the particle, leading to more electrostatic repulsion forces and hence the increase in the particle size.

Lucifer yellow VS shows two positive pK_a values (1.4 (0.8) and 5.7 (0.9)) [1]. Figure 6 shows the change in the hydrodynamic diameter of p(NIPAM)-co-5%LY according to the change of pH. The first positive pK_a of LY (1.4) (caused by the deprotonation of the nitrogen of the piperidine ring) [1] shows a minimal effect on the particle size (Figure 6). Comparing the particle size at pH 0.5 to that at pH 3 shows minimal change in the particle size at the two cases. An increase in the particle size around the pH of 5 which is in the range of the second positive pK_a of LY (caused by the deprotonation of the amino group (Figures 2 and 3) [1] is observed. The Figure shows 23% increase in particle size by increasing the pH from 3 to 5.

At pH below 5.7 (\pm 0.9), p(NIPAM)-*co*-5% LY can be considered as a zwitterionic particle. The negative surface charge on p(NIPAM-*co*-5% LY) particles is caused by the SO₄⁻ groups from the anionic initiator used during the particle synthesis (K₂SO₄) and the fluorescent dye (lucifer yellow) with sulphonate groups (Figure 2). Having two negative pK_a values, the sulphonate groups of lucifer yellow are deprotonated when the pH is above zero (Figures 2 and 3) [1]. At pH below 1.4, the nitrogen group of the piperidine ring and the amino group of the aniline are both protonated creating positive charge that masks some of the negative charge created by the initiator and the sulphonate groups of LY. At 1.4 < pH > 5.7, the nitrogen of the piperidine is deprotonated, but the tertiary amine group of the aniline is protonated and thus charged, so, positive charge caused by the protonated amino group







can still mask some of the negative charges. When the pH exceeds 5.7 (\pm 0.9), both groups (nitrogen of piperidine and amino of aniline) are deprotonated and thus no positive charges are created. At this point, the negative surface charge on p(NIPAM)-*co*-5% LY is maximal and not masked by positive charges (the particle is not zwitterionic anymore and tends to only carry negative charges on its surface. This increases the electrostatic repulsion within the particle and hence increases the particle size (Figure 5).

It is worth mentioning that the magnitude of change in the particle diameter of p(NIPAM)-co-5% LY at different pH values is less than that of previously known temperature/pH responsive particles such as p(NIPAM)-co-5% AA [3]. The initial particle diameter of p(NIPAM)-co-5% AA at 15°C at pH 3 is 290 nm while at pH 6 (at the same temperature) is 550 nm [3]. This shows a 90% increase in the particle size by changing the pH of the particle dispersion. In the case of p(NIPAM-co-5% LY), the particles only show a 32% increase in diameter when the pH is changed from 0.5 to 11.

The magnitude of particle size change in response to pH change can be increased by incorporating another co-monomer. This has been carried out by many researchers such as the incorporation of AA (3), *N*-*t*-butylacrylamide [31].The suggested result then is the synthesis of a fluorescent temperature/pH sensitive particle with a higher magnitude of pH sensitivity.

Electrophoretic mobility

Effect of temperature on the electrophoretic mobility: Figure 7 compares between the electrophoretic mobility of p(NIPAM) and p(NIPAM)-*co*-5% LY. p(NIPAM) has a negative surface charge due to the anionic initiator (potassium persulphate) used during synthesis while p(NIPAM)-*co*-5% LY has a negative surface charge due to both the initiator and the anionic co-monomer (LY) [32].

The Figure shows that at 15°C the electrophoretic mobility of p(NIPAM-co-5% LY) (-0.32 µmcm/Vs) is higher than that of p(NIPAM) (-0.17 µmcm/Vs). This can be attributed to two main reasons; first, the presence of a higher surface charge density on p(NIPAM-co-5% LY) particles due to the addition of an anionic co-monomer (lucifer yellow) [32] beside the anionic initiator (potassium persulphate) used during the synthesis. Second, is that the particle size of p(NIPAM-co-5% LY) is smaller than that of p(NIPAM) which causes the particles to move faster. When the temperature exceeds the VPTT (35°C), the particle deswells which results in a higher surface charge density [29] leading to a higher electrophoretic mobility.

Effect of pH on the electrophoretic mobility: The effect of pH on the electrophoretic mobility of p(NIPAM)-co-5% LY in 1×10^{-4} M NaCl at 25°C is shown in Figure 8. The data of the electrophoretic mobility versus pH shown in Figure 8 are comparable to those of the particle size shown in Figure 5. As discussed above (point 3.1.1.2), the electrophoretic mobility increases around the pK_a of LY (5.7 (0.9)) due to the deprotonation of the amino group of the aniline [1] (Figure 2) that causes an increase of the net negative charge on the particle surface.

Fluorescence spectroscopy

The fluorescence spectra of 0.01 μ g/mL LY, 0.2 μ g/mL p(NIPAM) and 0.2 μ g/mL p(NIPAM-*co*-5% LY) excited at 430 nm at 25°C and 45°C at pH 3 and 7 are shown in Figures 9 and 10.

The emission spectra of lucifer yellow at pH 7 at temperatures 25°C and 45°C (Figure 9) and at 25°C pH 3 and 7 (Figure 10) display an emission band at 532 nm which is comparable to the reported emission wavelength of lucifer yellow at 540 nm [17]. The emission spectra of

p(NIPAM)-*co*-5% LY at pH 3 and 7 and at 25°C and 45°C (Figures 9 and 10) show an emission band at 520 nm. Accordingly, a blue shift is observed when the fluorophore LY was incorporated into p(NIPAM). This is due to the change in the environment around the LY such as the electronic cloud in the polymer particle p(NIPAM) as well as the polarity of the solvent [18]. The shift of the emission band also provides evidence (together with the DLS and IR) of the incorporation of LY in the new nanoparticle.

Effect of addition of lucifer yellow: The concentration of LY used for the fluorescence spectra in Figures 9 and 10 (0.01 μ g/mL) is the theoretical concentration of LY incorporated in 0.2 µg/mL p(NIPAMco-5% LY) (5%), in other words, it is the concentration of LY added during the synthesis of p(NIPAM)-co-5% LY, and was selected to give a qualitative comparison of the emission spectrum intensity. However, the fluorescence intensity of LY is significantly higher than that of p(NIPAM-co-5% LY) which suggests that either not all the LY added during the synthesis reaction is incorporated in the new particle, or the quantum yield of LY was quenched by the p(NIPAM) particles, or other environmental conditions such as solvent polarity [18]. The second suggestion is supported by Furstenberg et al. [18] who studied the excited state dynamics of lucifer yellow and concluded that LY is an environment sensitive probe whose fluorescence dynamics change in the presence of proteins and by changing the solvent. They also concluded that the fluorescence of LY is quenched by amino acids. Given the fact that both proteins and polymer particles are considered as large complex molecules/particles, it is a possibility that polymer particles might have similar quenching effect on the fluorescence of LY.













p(NIPAM) does not show an emission band in a similar region or of a similar intensity. On the other hand, it shows some fluorescence intensity (of much lower magnitude than p(NIPAM)-co-5% LY which is thought to be due to light scattering [33]. This is a common characteristic of colloidal dispersions that is related to particle size. The turbidity of colloidal dispersions is due to the intense light scattering by its particles. The size and shape of the scattering particles are the main factors affecting the intensity of the scattered light. If a colloidal nanogel sample is placed in a UV spectrophotometer and a beam of light is passed through it, a fraction of the light will be scattered as it passes through the sample. While measuring fluorescence of p(NIPAM)and p(NIPAM)-co-5% LY, the scattered light from the particles can partially interfere with the fluorescence emission.

Effect of temperature: The effect of temperature on the fluorescence of LY, p(NIPAM) – co- 5% LY and p(NIPAM) can be compared using Figure 9. In 1852, Sir G. G. Stokes observed that fluorescence emission generally occurs at lower energies and longer wavelengths than that of the absorption [34]. In the case of LY, the emission band at pH 7 at both 25°C and 45°C are similar λ_{max} , band shape and intensity. This indicates that the effect of internal conversion (due to temperature change) on the Stokes shift is minimal. In terms of fluorescence intensity, no significant difference between the fluorescence at 25°C and 45°C is seen (Figure 9). The percentage change in fluorescence intensity between both temperatures is 4% at pH 3 and 1.6% at pH 7.

In the case of p(NIPAM)–co-5% LY, the emission bands at 25°C and 45°C (both at pH 7) show similar λ_{max} and band shape. On the other hand, a change in the fluorescence intensity is observed by changing

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ISSN: 2157-7439 JNMNT, an open access journal

the temperature (Figure 9). The percentage change in the fluorescence intensity was 11.7% at pH 3 (3 times higher than that of LY) and 14.6% at pH 7 (9 times higher than that of LY). This can be attributed to different factors such as the change in particle size by changing the temperature from below to above the VPTT (Figure 5) which changes the fluorophore density and accordingly the fluorescence intensity. It is worth mentioning that the particle size of p(NIPAM)-*co*-5% LY decrease by 43% when the temperature increases from 25°C to 45°C at pH 3 and 7. The change in temperature can also cause a change in the quantum yield of fluorescence due to change in solvent properties such as solvent polarity [18,35]. For p(NIPAM), the particles deswell when the temperature exceeds the VPTT (35°C), thus at 45°C, they acquire a de-swollen configuration and the dispersion is more turbid. This boosts the light scattering effect which is shown here as higher fluorescence intensity.

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Effect of pH: Lucifer yellow is a pH insensitive fluorescent probe [36]. This agrees with the results shown in Figure 10 where the emission bands of LY at same temperature (25°C) and different pH values (3 and 7) are similar (same λ_{em}). The same result is also shown for p(NIPAM)*co*-5% LY.

The fluorescence intensity of LY is affected by the change in pH (Figure 10). This can be due to different factors that affect the quantum yield of the fluorophore lucifer yellow. These include; the change of the polarity of the surrounding polymer (p(NIPAM)) and its electrophoretic mobility which affects the polarity of the solvent [18]. Changing the solvent polarity highly affects the fluorescence lifetime which in return affects the quantum yield [18,37,38]. Also, the change in particle size (including expel and uptake of solvent) causes a dramatic change in the surrounding local environment. Other factors that can cause a change of the fluorescence intensity by changing the pH include; change in the rate of solvent relaxation and internal charge (proton) transfer [18].

Microscopy

Scanning Electron Microscopy (SEM) and Fluorescence microscopy, the nanoparticles were examined using SEM (Figure 11) and fluorescence microscopy (Figure 12). Figure 11 show that the particles are spherical, monodispersed and their average size is 120 nm which is comparable to that shown by DLS at deswollen state (Figure 5). The fluorescence microscopy (Figure 12) also confirms the results of SEM where the particles are shown to be spherical and monodispersed as well as fluorescent. Figure 12 supports the suggestion that LY is



Figure 11: SEM pictures of p(NIPAM-co-5% LY) at pH 3 (x 45,000)



incorporated in the p(NIPAM-*co*-5% LY) nanoparticles and that they are fluorescent. Accordingly, it can be suggested that these particles can be traced in the human body (given their fluorescence) which can serve as a diagnostic tool. Researches have tried incorporating fluorescent dyes into nanogels [39,40], on the other hand, there are no up-to-date data showing fluorescence microscopy images of fluorescent nanogels.

Toxicity studies

p(NIPAM) based nanoparticles have been studied for a number of years [41-45]. Many applications have been suggested for this polymer due to its ability to change its physico-chemical properties according to the change in environmental conditions [3,4]. Considering that the VPTT of p(NIPAM) nanoparticles is 33-35°C [3] which is very close to the human body temperature, many researchers investigated the possible use of p(NIPAM) nanoparticles in drug delivery [8,46-48], delivery of stem cells [49] and other biomedical applications such as tissue engineering [27], biomedical implants [50], treatment of dentinal hypersensitivity [3] and wound dressings [51,52]. A very important point to consider in this case is the toxicity of NIPAM based nanoparticles; on the other hand, not many reports are available in this context to date.

In 2010, Naha et al. [53] studied the cyto- and geno-toxicity of p(NIPAM) nanoparticles (within a concentration range of 25-1000 mg/L) on two different mammalian cell lines, SW480 (colon) and HaCaT (dermal cell line). The viability of cells (both cell lines) after 24, 48, 72 and 96 hours were shown to be 100%. Also, the geno- and ecotoxicity [54] data confirmed the results of the cytotoxicity suggesting that p(NIPAM) nanoparticles have an excellent bioavailability profile *in vitro*. Naha et al. [53] have further studied p(NIPAM) based nanoparticles inside cells and concluded that the particles enter the cells and are localised in the lysosomes.

The cytotoxicity and uptake of p(NIPAM) based nanoparticles with incorporated hyperbranched polyglycerol (HPG) was investigated by Cuggino et al. [55]. The cytotoxic effect of p(NIPAM)-*co*-HPG nanoparticles on epithelial human lung cancer cell line A549, human hematopoietic cells U-937, and human epidermoid carcinoma cells A431. Results suggest that doses above 0.5 mg/mL are required to cause adverse effects on the cells. It is also worth mentioning that the particle could penetrate the cell membrane of lung cancer cells A549 incubated at 37°C and were distributed in the perinuclear region [55].

Herein, the effect of dextran, NIPAM monomer, p(NIPAM), p(NIPAM-co-5% LY) and PEI on the cell viability of both HeLa and Vero cell lines is reported. HeLa cells are human epithelial cells of a strain maintained in tissue culture since 1951 and used in research,

especially in virology. The original cells were provided by cervical carcinoma of *Henrietta Lacks* [56]. Vero cells are taken from the kidneys of the African green monkey, Cercopithecus aethiops sabaeus, and used to grow certain viruses for vaccine production [57].

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Figure 13 shows the effect of different concentrations (concentration range between 0.03 µg/mL to 3 mg/mL) of dextran, NIPAM monomer, p(NIPAM), p(NIPAM-co-5% LY) and PEI on the cell viability of HeLa cells. The Figure shows that the effect of dextran (used as a negative control) on the cell viability was minimal as the cells showed over 90% viability across the whole range of concentrations. On the other hand, PEI (used a positive control) shows cell viability above 80% at concentrations less than 0.003 mg/mL. When the concentration further increases, the cell viability significantly decreases till it reaches 20% at a concentration of 0.03 mg/mL. In spite of its reported toxicity [58,59]. PEI is widely researched as one of the main non-viral vectors for gene therapy [60]. These results are comparable to those of Zhizhong et al. [61] who studied the cytotoxicity of linear 25 KD PEI (known to be less toxic than branched PEI [58,60,62]) on OS-RC-2 cell line and reported that cell viability under 30 µg/mL of 25 kD PEI was 50%. For 2 kD PEI, half of the cells were viable when the concentration was 200-300 μ g/ mL. Similar results were also shown by Thomas et al. [63] who reported that less than 50% of COS-7 cells were viable when the concentration of 25 kD PEI was more than 20 µg/mL while less than 5% of the cells were viable when its concentration was above 35 µg/mL.

Despite its reported cytotoxicity [58,60-64], the use of PEI has been studied for biological applications such as gene delivery [59,61,65,66], nucleic acid delivery [62,63,67-69], drug delivery [64], cosmetics [70] and tissue engineering [71]. In 2012, Wegmann et al. [72] claimed that PEI is a potent mucosal adjuvant for human use and suggested that further investigation for improvement of potency and safety of PEI as an adjuvant suggests promising results. Many attempts to decrease the cytotoxicity of PEI by conjugation with other molecules have been varied out. These aim at turning PEI into a bioequivalent delivery system [61-63,68,69]. Similarly, it can be suggested that the conjugation of acrylamide and its derivative NIPAM into p(NIPAM) based particles can yield particles with significantly lower toxicity. It is also worth mentioning that PEI is currently commercially available (in the form of PEI-HCI) in a drug in the German market known as 'Epipak' used for mouth disorders [73].

The main challenge for the use of NIPAM based particles as drug delivery systems is its toxicity. The fact that the main monomer NIPAM includes acrylamide group which is known for its toxicity [74], suggests that the polymerised particle and/or its metabolic products (such as NIPAM monomer and/or acrylamide) might acquire the same properties. Accordingly, the toxicity of the new particle p(NIPAM*co*-5% LY), p(NIPAM) and the monomer was tested. In Figure 13, p(NIPAM) shows cell viability over 80% up to a concentration 3 mg/mL while NIPAM monomer showed a high cell viability (over 80%) for concentration equal to, or less than, 0.3 mg/mL. Above this concentration, NIPAM monomer shows considerably less viability untill it reaches 10% at a dosing concentration of 3 mg/mL.

For p(NIPAM-*co*-5% LY) particles, the results show that the cell viability is around 80% or higher up to 3 mg/mL. This suggests that p(NIPAM-*co*-5% LY) is suitable for medical and/or pharmaceutical use when used within the specified concentrations. Similar results for the effect of the same concentrations of dextran, NIPAM monomer, p(NIPAM), p(NIPAM-*co*-5% LY) and PEI on the cell viability of Vero cells. Figure 14 shows that dextran showed cell viability over 80% across all the dosing concentrations. PEI showed cell viability over 80%



Figure 13: Percentage cell viability (after 72 hrs) versus log concentration (across conc. Range of 0.03 μ g/mL to 3000 μ g/mL) of dextran, NIPAM monomer, p(NIPAM), p(NIPAM-co-5% LY) and PEI on 1 x 104 HeLa cell/mL.



Figure 14: Percentage cell viability (after 72 hrs) versus log concentration (across conc. range of $0.03 \ \mu$ g/mL to $3000 \ \mu$ g/mL) of dextran, NIPAM monomer, p(NIPAM), p(NIPAM-co-5% LY) and PEI on 1 x 104 Vero cell/mL.

at concentrations equal to or less than 0.001 mg/mL. Increasing the dosing concentration decreased the cell viability till it reached 15% at 0.01 mg/mL. NIPAM monomer showed high cell viability (between 80 and 100%) at concentrations equal to, or less than, 0.3 mg/mL. Both p(NIPAM) and p(NIPAM-co-5% LY) showed cell viability around 80% or more for all the dosing concentrations used.

Comparing the cytotoxicity (for both HeLa and Vero cells) of p(NIPAM-co-5% LY) and p(NIPAM) to that of PEI shows that the former particles show a safer toxicity profile. While a concentration of (0.7 µg/mL for HeLa and 3 µg/mL for Vero) of PEI killed half of the cells (Table 2), 3 mg/mL of the NIPAM based particles kept more than 80% of the cells alive. Furthermore, NIPAM monomer which is expected to be one of the metabolic products of p(NIPAM) based particles has an $LC_{_{50}}$ of 700 $\mu g/mL$ (for both HeLa and Vero cells) which is considerably higher than that of PEI. Thus, the results show that p(NIPAM) based particles as well as NIPAM monomer (at the specified concentrations) have a safer toxicity profile when compared to PEI. Considering the ongoing research and use of PEI in biological and pharmaceutical applications (previously discussed) and the properties of p(NIPAM) based particles such as temperature, pH, ionic strength responsiveness, it is suggested that further research of the toxicity of p(NIPAM) based particles as well as modification of the particles can yield very useful and promising systems for biological and pharmaceutical applications such as drug delivery, gene therapy and treatment of dentinal hypersensitivity.

The biocompatibility of p(NIPAM), p(NIPAM-co-AA) and p(NIPAM-co-BA) with skin has been studied by Abu Samah et al. [75]. The effects of the three nanoparticles on the expression of cyclooxygenase-2 (COX-2) was determined using western blotting

which showed that the particles have penetrated the skin and the keratinocytes of the vital epidermis. p(NIPAM)-*co*-BA showed 67% higher COX-2 expression than the control group which indicates a pro-inflammatory response. For p(NIPAM)-*co*-AA, no significant modulation in the expression of COX2 was shown, indicating the compatibility of the particles with skin. Comparable results were obtained for nanoparticles loaded with citric acid solution. Accordingly, the authors support the use of the multiresponsive p(NIPAM)-*co*-AA as a triggered or controlled topical drug delivery system.

For *in vivo* testing, Aslam et al. [76] tested the effect of different concentrations (0, 0.2, 0.4, 0.6, 0.8 and 1 mg/mL) of 150 nm p(NIPAM) on Swiss Albino mice (Mus musculus). Oral p(NIPAM) dispersions (with the previously mentioned concentrations) were given to the mice daily for 30 days. The results suggest that lower doses (0.2 mg/mL) had no lethal effect on the mice. Increasing the dose of p(NIPAM) (0.4, 0.6 and 0.8 mg/mL), half the mice died in 20-25 days. For 1 mg/mL p(NIPAM) dispersions, half the mice died in 28 days. The authors suggest that p(NIPAM) is tolerable at low doses, on the other hand, the accumulation of high doses of the particles (by daily administration of 1 mg/mL/kg) has a lethal effect in an average of 18 days.

In 1981, Tanii et al. claimed that the metabolic product of NIPAM is acrylamide [77], accordingly, it is very important to consider its toxicity while studying the toxicity of NIPAM based particles. Many researchers have claimed that acrylamide is toxic [78], some mention its neurotoxicity [78], others mention its genotoxicity [78] as well as other serious effects such as testicular degeneration, necrosis of the liver, congestion of the lungs. Acrylamides are formed when some foods including certain nutrients are cooked at certain temperatures during grilling, frying and baking [73,79]. A review studying the doseresponse relationship of acrylamide was carried out by Shipp et al. [79], the study states that the reference dose (RfD) (safe dose) of acrylamide is 0.83 µg/kg/day (considering reproductive effects and genotoxicity) and 1.2 µg/kg/day (considering neurotoxicity). The review also claims that most of the reports of acrylamide toxicity did not consider very important factors such as the smoking state of the patients which is extremely important [79]. Considering the RfD mentioned by Shipp et al. [79], an adult of average weight 70 kg can safely have 58 µg/day of acrylamide.

Most of the drugs currently available in the market can cause different kind of toxicities and can even be lethal if administered at the wrong dose. It is worth mentioning at this point, that p(NIPAM) based particles (if pharmaceutically used) are intended to be used as a drug delivery system and not a drug, this might be important while calculating the required dose.

Conclusion

A novel fluorescent temperature/pH sensitive particle (p(NIPAM)co-5%LY) was successfully synthesized and characterized using different techniques. At pH 7 and 15°C (the particle is swollen), the

	LC ₅₀ on HeLa cells (µg/ mL)	LC ₅₀ on Vero cells (μg/ mL)			
Dextran	≥ 3000				
NIPAM monomer	700 (±6)	700 (±6)			
p(NIPAM)	≥ 3000				
p(NIPAM-co-5% LY)	≥ 3000				
PEI	0.7 (±0.2)	3 (±0.9)			

Table 2: LC $_{\rm 50}$ of dextran, NIPAM monomer, p(NIPAM), p(NIPAM-co-5% LY) and PEI on HeLa Vero cells.

particle size is 250 nm. When the temperature exceeds the VPTT (35° C) the particle de-swells to reach 130 nm. These particle sizes are approximately half that of p(NIPAM) at similar conditions. The PdI and the intensity peaks of the new nanoparticle dispersion indicate a uniform particle size across the dispersion.

This study only provides initial evidence of the biocompatibility of NIPAM based particles and its metabolite at the specified concentrations. NIPAM monomer is more toxic than both p(NIPAM) and p(NIPAM-*co*-5% LY). The two nanoparticles show cell viability over 80% (for both cell lines Hela and Vero) up to a concentration of 3 mg/mL while NIPAM monomer show cell viability over 80% equal to or less than 0.3 mg/mL. At concentration of 3 mg/mL, NIPAM monomer shows 10% cell viability. These data add to the few existing reports on the toxicity of NIPAM based particles and provide some encouragement around their biocompatibility and possible use for drug delivery applications.

The advantages (temperature, pH, ionic strength, etc. responsiveness and ability to release drugs accordingly) of NIPAM based particles can be very useful medically and pharmaceutically. Such advantages can at some points be louder than other disadvantages (depends upon the case). These particles can be very useful and flexible drug delivery systems that can be used for the treatment of different diseases such as cancer and diabetes. Further studies about the ability of p(NIPAM) based particles to be used for gene and nucleic acid delivery can also be considered.

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Citation: Mohsen R, Alexander BD, Richardson SCW, Mitchell JC, Diab AA, et al. (2016) Design, Synthesis, Characterization and Toxicity Studies of Poly (*N-Iso*-Propylacrylamide-*co*-Lucifer Yellow) Particles for Drug Delivery Applications. J Nanomed Nanotechnol 7: 363. doi:10.4172/2157-7439.1000363