Alkyne-tagged PLGA allows direct visualisation of nanoparticles in vitro and ex vivo by stimulated Raman scattering microscopy

Citation for published version:

Digital Object Identifier (DOI):
10.1021/acs.biomac.9b01092

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Peer reviewed version

Published In:
Biomacromolecules

General rights
Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy
The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.
Alkyne-tagged PLGA allows direct visualisation of nanoparticles \textit{in vitro} and \textit{ex vivo} by stimulated Raman scattering microscopy

\textit{Sally Vanden-Hehir} \textit{a}, \textit{Stefan Cairns} \textit{a}, \textit{Martin Lee} \textit{b}, \textit{Lida Zoupi} \textit{c}, \textit{Michael P. Shaver} \textit{a}†, \textit{Valerie G. Brunton} \textit{b}, \textit{Anna Williams} \textit{c} and \textit{Alison N. Hulme} \textit{a}*

\textit{a}EaStCHEM School of Chemistry, University of Edinburgh, David Brewster Road, Edinburgh, EH9 3FJ, UK

\textit{b}Edinburgh Cancer Research Centre, University of Edinburgh, Western General Hospital, Crewe Road South, Edinburgh, EH4 2XR, UK

\textit{c}MRC Centre for Regenerative Medicine, The University of Edinburgh, Edinburgh BioQuarter, 5, Little France Drive, Edinburgh, EH16 4UU, UK

\textbf{ABSTRACT}

Polymeric nanoparticles (NPs) are attractive candidates for the controlled and targeted delivery of therapeutics \textit{in vitro} and \textit{in vivo}. However, detailed understanding of the uptake, location and ultimate cellular fate of the NPs is necessary in order to satisfy safety concerns, which is difficult because of the nanoscale size of these carriers. In this work, we show how small chemical labels
can be appended to poly(lactic acid-co-glycolic acid) (PLGA) in order to synthesise NPs that can then be imaged by stimulated Raman scattering (SRS) microscopy, a vibrational imaging technique which can elucidate bond specific information in biological environments, such as the identification of alkyne signatures in modified PLGA terpolymers. We show that both deuterium and alkyne labelled NPs can be imaged within primary rat microglia, and the alkyne NPs can also be imaged *ex vivo* cortical mouse brain tissue. Immunohistochemical analysis confirms that the NPs localize in microglia in the mouse brain tissue, demonstrating that these NPs have the potential to deliver therapeutics selectively to microglia.

**KEYWORDS**

Nanoparticles, drug delivery, alkynes, Raman imaging, bioorthogonal polymers, stimulated Raman scattering

**INTRODUCTION**

The use of polymeric NPs for drug delivery has become increasingly popular in order to achieve controlled release, targeted delivery and increased lifetime of therapeutics *in vivo*. NP drug delivery can have wide ranging applications such as the targeting of cancer therapeutics,\(^1,2\) delivery of drugs to the brain,\(^3,4\) and encapsulation protein therapeutics, which are sensitive to certain biological environments and may degrade to an inactive form *in vivo*. PLGA micro and nanoparticles have been used extensively in drug delivery research as they are FDA approved, biocompatible and biodegradable.\(^5\) However, due to the nanoscale size of these drug carriers,
imaging their uptake, bio-distribution and ultimate cellular fate \textit{in vitro} and \textit{in vivo} is challenging.

Previous efforts to image polymer NPs have mainly focused on the encapsulation of fluorescent dyes inside the NPs to allow imaging by fluorescence microscopy,\textsuperscript{6} or by various Raman-based methods, which we have recently reviewed.\textsuperscript{7} The degradation of PLGA NPs in macrophages has been monitored by quantifying the loss of fluorescent signal over time.\textsuperscript{8} PLGA NPs degrade by hydrolysis under physiological conditions,\textsuperscript{9} thus the loss of fluorescent signal could correspond to dye being released from the NPs, and not give direct information about the location and state of the NP. Also, fluorophores are generally unstable in biological environments, and photobleaching can also cause signal degradation of the dye (Figure 1).\textsuperscript{10}

Stimulated Raman scattering (SRS) microscopy has emerged as a powerful, label-free optical imaging technique for biological systems that can provide quantitative and bond-specific cellular information.\textsuperscript{11,12} Spontaneous Raman scattering can be used to probe the chemical signature of cells and biological tissues due to the inelastic scattering of light when the sample is irradiated with a laser, but recording an image may take many minutes to hours. With coherent Raman techniques like SRS, a specific vibrational bond can be probed using two lasers tuned to the bond’s energy difference. This drives a coherent resonance allowing video rate imaging to be achieved.\textsuperscript{13}
Figure 1. Illustrative summary of different strategies for intracellular NP imaging. Fluorescent dyes have been encapsulated in NPs to allow visualisation. Disadvantages of this approach are that dye can leach out of the NPs and suffer from photobleaching. In this work we directly label the polymer with a bioorthogonal Raman tag. Image created with Biorender.com.

In contrast to infrared, Raman spectroscopy has a low background interference from water making it ideally suited to study biological samples. Due to the absence of any cellular vibrational peaks in the spectroscopic region between 1800 and 2800 cm\(^{-1}\), known as the cell-silent region, small, bioorthogonal chemical tags (for example C-D, C≡C and C≡N) which resonate in this region can be exploited to give imaging free of cellular background.\(^{14-19}\) As such, poly(phenylene ethynylene) NPs, which have an intrinsic alkyne, have been visualised in HeLa cells with spontaneous Raman imaging.\(^{20}\) More recently, SRS has been used to image NP polymer dots bearing alkyne (2163 cm\(^{-1}\)), nitrile (2232 cm\(^{-1}\)) and carbon-deuterium (2293 cm\(^{-1}\)) bonds, which all produce peaks in the cell silent region.\(^{21}\)

In this work, we have explored two analogues of PLGA, with deuterium and alkyne labels which are Raman active in the cell-silent region. These polymers were then formulated into NPs and
imaged firstly in cultured primary microglia and then in *ex vivo* brain tissue by SRS microscopy. As opposed to imaging a fluorescent payload, our approach allows direct imaging of the nanoparticle itself, and can be used to monitor the cellular location of the NPs. SRS microscopy gives a signal enhancement over spontaneous Raman imaging, and also a 1000 fold improvement in image acquisition speed.\(^{22}\) We have therefore used SRS to image NP analogues of PLGA, a highly relevant biocompatible and biodegradable polymer.

**MATERIALS AND METHODS**

**Synthesis of PLGA:** Under a nitrogen atmosphere D,L-lactide (8.0 g, 55 mmol) and glycolide (2.8 g, 24 mmol) were added to a preheated round bottom flask containing stannous octanoate (0.16 g, 0.40 mmol) and dodecanol (0.07 g, 0.40 mmol) at 130°C. The reaction mixture was heated and stirred for a further hour until complete conversion was reached. After the reaction was cooled to room temperature, it was dissolved in dichloromethane (DCM) (5 mL) and precipitated in cold methanol (400 mL). The solvent was decanted off, leaving a colorless solid which was dried *in vacuo* until constant weight. \(^1\)H NMR (500 MHz, Chloroform-d) \(\delta 5.30 – 5.11 \text{ (9H, m, CH)}\), 4.97 – 4.58 (7H, m, CH\(_2\)), 1.64 – 1.49 (23H, m, CH\(_3\)); GPC \(M_n = 5700 \text{ g/mol, D} = 1.11\).

**Synthesis of PLGA-D:** D,L-Lactic acid (0.10 g, 0.94 mmol; 85% in water), L-lactic acid-d\(_3\) (0.10 g, 0.91 mmol; 85% in water) and glycolic acid (0.06 g, 0.79 mmol) were dehydrated at 150°C for 8 h in a flask fitted with an air condenser containing molecular sieves to form oligomers. Tin chloride (0.5 wt%) and *para*-toluene sulfonic acid (0.5 wt%) were then added to the oligomers followed by heating to 180°C for a further 8 h. The crude polymer was dissolved in DCM (2 mL)
and precipitated into a 50:50 mixture of ice-cold hexane/ether (100 mL) to yield a colorless solid which was dried in vacuo until constant weight. $^1$H NMR (400 MHz, Chloroform-d) $\delta$ 5.30-5.13 (1H, m, CH), 4.93-4.56 (2H, m, CH$_2$), 1.64-1.51 (3H, m, CH$_3$). GPC $M_n$ = 4553 g/mol, D = 1.84.

**Synthesis of PLGA-alkyne:** In a glovebox, PLGA (2.00 g, 0.56 mmol), propargyl-DOX (0.38 g, 2.78 mmol, see Scheme S2), and SnOct$_2$ (0.23 g, 0.56 mmol) were dissolved in toluene (4 mL) and charged to a Schlenk flask. The flask was removed from the glovebox and heated at 110°C for 24 h. After the reaction was cooled to room temperature, the reaction mixture was diluted further by the addition of DCM (5 mL) and the mixture was precipitated in cold methanol (400 mL). The solvent was decanted off leaving a colourless solid which was dried in vacuo until constant weight. $^1$H NMR (500 MHz, Chloroform-d) $\delta$ 5.43 – 5.11 (m, 9H), 4.97 – 4.58 (m, 7H), 2.99 – 2.77 (m, 2H), 2.14 – 2.03 (br s, 1H), 1.64 – 1.49 (m, 23H). GPC $M_n$ = 5900 g/mol, D = 1.88.

**Nanoparticle synthesis and characterisation:** A solution of PLGA-D/PLGA-alkyne (10-20 mg) in DCM (1 mL) was added to an aqueous solution of 1% polyvinyl alcohol (MW 31 000 – 50 000 Da, 98-99% hydrolysed) and 0.2% sodium dodecyl sulphate (10 mL). The two phases were emulsified using a probe tip sonicator (Soniprep 150, MSE) for 2 minutes. The resulting emulsion was stirred at room temperature overnight to allow evaporation of the DCM. The nanoparticles were collected by centrifugation at 8000 rpm for 10 minutes, and washed once with deionised water. Nanoparticle tracking analysis measurements were carried out in triplicate in 1 × PBS on a NanoSight LM10 (Malvern Panalytical) for 60 seconds with the temperature measured for each individual run. The samples were diluted until in the range of 1-8 × 10$^8$
particles mL$^{-1}$ for more accurate measurements. Dynamic light scattering (DLS) was carried out on a Zetasizer Nano ZS (Malvern Panalytical) in water at 25°C.

**Spontaneous Raman:** Spontaneous Raman spectra were acquired on a confocal Raman spectrometer (inVia Raman microscope, Renishaw). A 297 mW (206 mW after objective) 785 nm diode laser, or a 200 mW 532 nm laser excitation source was used to excite the sample through a 20× or 50× objective. All spectra were background subtracted using the background correction algorithm available on the Wire 4.4 software.

**SRS setup:** Images were acquired using a custom-built multi-modal microscope. A picoEmerald (APE, Berlin, Germany) laser gave a tuneable pump laser (720-990 nm, 7 ps, 80 MHz repetition rate) and a spatially and temporally overlapped Stokes laser (1064 nm, 5-6 ps, 80 MHz repetition rate). Back scattered RFP two-photon fluorescence signals were filtered using the following series of filters: FF552-Di02, FF440/520-Di01 (Semrock) and HQ610/75m (Chroma). For SRS measurements, the Stokes beam was modulated with a 20 MHz EoM. Forward scattered light was collected by a 20× Olympus XLUMPLFLN Objective, 1.00 NA lens and filtered using ET890/220m filter (Chroma). A telescope focused the light onto an APE silicon photodiode connected to an APE lock-in amplifier which was fed into the analogue unit of the microscope.

The pump laser was tuned to 810.5 nm (2930 cm$^{-1}$, protein), 816 nm (2850 cm$^{-1}$, lipid), 867.5 nm (2128 cm$^{-1}$, C≡C) and 858.3 nm (2252 cm$^{-1}$, C-D), and laser powers after the objective were measured up to 40-70 mW for the pump laser, and up to 70 mw for the Stokes laser. All images
were recorded at 512 × 512 or 1024 × 1024 pixels with a pixel dwell time between 2 and 20 μs by FV10-ASW software (Olympus).

**General procedures for cell culture:** Microglia cultures were prepared from neonatal Sprague-Dawley rats. Mixed glia cultures were isolated from postnatal day 1-2 rat pups, and after 10 days microglia were isolated by shaking the culture flasks and relying on differential adhesion of oligodendrocyte precursor cells, microglia and astrocytes. Microglia were cultured in Dulbecco’s Modified Eagle Media (Gibco) + 10% fetal bovine serum + 1% penicillin streptomycin, and were incubated at 37°C in a humidified atmosphere with 7.5% CO₂ and media changes every 2-3 days. To improve cell adherence, all culture dishes were coated in poly-D-lysine before plating the cells by covering dishes in a 1 μg/mL aqueous solution and incubating at 37°C for one hour, before removing the solution and washing once with sterile water. Cell fixation was achieved by covering the cells with a solution of 4% formaldehyde (2 mL) for 10 minutes at room temperature before washing three times with 1 × PBS (3 × 2 mL).

**SRS microglia experiments:** Microglia were plated at 3 × 10⁵ cells per well in FluoroDish Cell Culture Dishes (World Precision Instruments) and left overnight to adhere to the dish. NPs (alkyne or deuterium) were then added at 2 × 10⁶ particles mL⁻¹ followed by a 24 hour incubation. The media was then removed and the cells were washed twice with 1 × PBS (2 × 2 mL) before fixation and imaging with the SRS set-up described above.

**Slice culture experiments:** Brains were isolated from P5 C57BL6/N mice pups in cold Hibernate-A medium (Thermo Fisher A12475-01) on ice and cut into 300 μm cortical slices using a vibratome (Leica). The slices were then transferred to Millipore milli cell-CM
organotypic inserts (30 mm, hydrophilic PTFE, 0.4 µm, Merck-Millipore, PICM0RG50) in slice media (50% MEM (Invitrogen, 32360-026) with 25% Earle’s Balanced Salt Solution (Invitrogen, 24010-043), 25% heat-inactivated horse serum (Invitrogen, 26050-088), 1% glutamax supplement (Invitrogen, 35050-038), 1% penicillin–streptomycin, 0.5% Fungizone (Invitrogen, 15290-018) and 6.5 mg/ml glucose (Sigma G8769)), and were maintained at 37 °C and 7.5% CO₂ with media changes every two days. After 6 days in culture, alkyne NPs were added at $2 \times 10^8$ and $2 \times 10^9$ particles mL$^{-1}$ for 24 hours. The slices were then washed twice with $1 \times$ PBS (2 × 2 mL) before fixation. Slices were fixed with 4% formaldehyde (2 mL) for one hour at room temperature before washing three times with $1 \times$ PBS (3 × 2 mL).

**Immunostaining of slices:** Fixed slices in a 6 well plate were incubated with blocking solution (3% HIHS, 2% BSA, 0.5% Triton in PBS) (1 mL per well) for 2 h before adding the primary antibodies. Rabbit polyclonal anti-IBA1 (1/500, Abcam, AB178846) and mouse monoclonal anti-OLIG2 (1/500, Merck Millipore, MABN50) were diluted in blocking solution and 500 µL was added to each well. The slices were then shaken at 4 °C for 1.5 days before washing three times for 1 h with blocking solution (3 × 2 mL). The secondary antibodies, goat anti-rabbit 568 (1/1000, Life Technologies, A11011) and donkey anti-mouse 488 (1/1000, Thermo Fisher Scientific, A21202) were diluted in blocking solution and then incubated with the slices (500 µL per well) with gentle agitation overnight, in the dark at 4 °C. The slices were then washed once for 10 minutes followed by three 1 h washes with $1 \times$ PBS (4 × 2 mL) and finally imaged using the SRS set-up described above.
Image processing: All images were processed with the Fiji image processing package (https://imagej.net/Fiji). False colour assignments and scale bars were added to images. Image overlays and orthogonal views were also processed with Fiji.

RESULTS AND DISCUSSION

Deuterated PLGA (PLGA-D) was synthesised via a direct poly-condensation method from lactic acid, lactic acid-d$_3$ and glycolic acid, which produced a polymer with a molecular weight of 4.5 kDa and a dispersity of 1.84 (Scheme 1A).$^{24,25}$ Since Raman is an inherently weak effect, with only approximately 1 in every $10^8$ molecules experiencing Raman scattering,$^{26}$ a deuterium group was introduced into the monomer, as opposed to substitution onto a pre-made polymer, to maximise the Raman signal of the NPs.

To produce the alkyne analogue of PLGA (PLGA-alkyne), PLGA was first synthesised by a ring opening copolymerisation of lactide and glycolide, using dodecanol as the initiator and stannous octanoate as the catalyst.$^{27}$ Recent work has shown that 1,3-dioxolan-4-ones (DOX) are a versatile and sustainable family of monomers utilised for the synthesis of biodegradable polyesters by ring opening polymerisation.$^{28,29}$ To introduce the alkyne functionality, the telechelic properties of PLGA were exploited to initiate the polymerisation of propargyl-DOX (see Scheme S.2 for propargyl-DOX synthesis). Ring-opening polymerisation from the PLGA macroinitiator initially affords a blocky PLGA-P(propargyl-DOX) copolymer. However, extended reaction times lead to transesterification catalysed by stannous octanoate, scrambling the terpolymer structure to afford a statistical distribution of the alkyne units throughout the PLGA chain (Scheme 1B). This gave rise to a polymer with comparable molecular weight and dispersity to the deuterium analogue (5.9 kDa and 1.88 respectively). The incorporation of the
alkyne into the terpolymer was confirmed by a single cross peak in the diffusion-ordered spectroscopy (DOSY) NMR (Figure S2).

**Scheme 1.** Synthesis of Raman active PLGA analogues. A) Synthesis of PLGA-D. Reagents and conditions: a; 150°C, 8 h, b; SnCl$_2$, p-TSA, 180°C, 8 h. B) Synthesis of PLGA-alkyne. Reagents and conditions: c; Sn(oct)$_2$, toluene, 110°C, 24 h.

The Raman labelled polymers, along with unlabelled PLGA, were then characterised with spontaneous Raman spectroscopy (Figure 2). PLGA (black) was analysed as a control to show that there are no peaks in the cell-silent region. In contrast, the PLGA-D (red) shows multiple broad peaks between 2000 and 2300 cm$^{-1}$, and the PLGA-alkyne (blue) shows a single, intense peak at 2128 cm$^{-1}$. Notably, the alkyne signal presents a much stronger peak in the cell silent region than the deuterium, which is consistent with previous studies.$^{17}$
Figure 2. Analysis of polymers by Spontaneous Raman spectroscopy. Spontaneous Raman spectra of PLGA (black), PLGA-D (red) and PLGA-alkyne (blue), with the labelled analogues showing peaks in the cell silent region. Spectra are normalised to the carbonyl peak (C=O) at 1737 cm$^{-1}$ and offset for clarity, $\lambda_{ex}$=785 nm.

The PLGA analogues were then fabricated into NPs using the standard, and widely used, emulsification-evaporation method (Figure 3A), which can be easily modified to encapsulate both hydrophobic and hydrophilic drugs. The NPs were analysed by nanoparticle tracking analysis (NTA), a technique which uses an optical microscope equipped with a laser to visualise particles according to the light they scatter. Individual particles were then tracked and their Brownian motion measured, which gives both particle size and particle concentration data (Figure 3B). This showed that the mean size of the PLGA-D and PLGA-alkyne particles are 87 and 81 nm respectively. Visualisation of the particles by the NTA shows that they are uniform and well separated (Figure 3C). Due to the tendency of the PLGA-D NPs to aggregate over time (Figure S4), the NPs were freshly prepared before every biological experiment.
Figure 3. The synthesis and characterization of polymer NPs. A) A general representation of the emulsification-evaporation process. B) Particle size distribution acquired from the NTA of PLGA-alkyne (blue) and PLGA-D (red). Spectra are normalised between 0 and 1 and offset for clarity. C) A visual representation of the NPs detected by the NTA camera, which is proportional to the amount of light scattered by the particles.

Next, we treated primary rat microglia with our two Raman active NPs. We chose microglia as ‘proof of principle’ cells as they are phagocytes in the brain, and therefore likely to internalise the NPs.\textsuperscript{33} NPs can cross the blood-brain barrier, and so have great potential for delivery of cargo to the brain.\textsuperscript{34,35} The microglia used were primary cultures isolated from rat brains to retain more biologically relevant \textit{in vivo} characteristics as opposed to using cell lines.
To test whether NPs were toxic to microglia, we used a luminescence cell viability assay which quantifies the amount of ATP, indicative of live cells. This assay found that PLGA, PLGA-D and PLGA-alkyne NPs were all non-toxic to microglia at all concentrations tested (1, 2 and $4 \times 10^9$ particles mL$^{-1}$) (Figure S5).

Microglia were then incubated with PLGA-D or PLGA-alkyne NPs for 24 hours before imaging with SRS microscopy (Figure 4). Tuning the energy difference between the pump and Stokes beams of the SRS microscope to 2939 cm$^{-1}$ excites CH$_3$ vibrations only, showing the protein content of the cells. Similarly, tuning to 2856 cm$^{-1}$ excites CH$_2$ vibrations, which is indicative of the cellular lipid content. Due to the bioorthogonal deuterium and alkyne chemical labels on our NPs, tuning to 2253 cm$^{-1}$ (C-D) and 2128 cm$^{-1}$ (C≡C) respectively allowed visualisation of the cellular location of the NPs, which were shown to be distributed throughout the cytoplasm and absent from the nuclei. Z-stack analysis of the cells also confirmed that the NPs had been internalised (Figure S6).

Time dependent analysis of PLGA-alkyne NP internalisation into microglia was also carried out over 24 hours. The alkyne intensity of individual cells at 0, 12 and 24 hours were quantified, and this showed that the alkyne intensity, and therefore the intensity of NPs, increased over time (Figure S7).
Figure 4. SRS imaging of NPs in microglia. Microglia were incubated with PLGA-D NPs (A) or PLGA-alkyne NPs (B) at $2 \times 10^9$ particles mL$^{-1}$ for 24 hours before fixing and imaging with SRS microscopy. Scale bars 20 µm.

As predicted by inspection of the spontaneous Raman spectra in Figure 2, the PLGA-alkyne NPs gave a stronger intracellular signal compared to PLGA-D. For this reason, we chose to carry out further studies with the PLGA-alkyne NPs only. Using a multi-modal approach to imaging, NPs encapsulating rhodamine were visualised using SRS (2128 cm$^{-1}$, NP) and two photon fluorescence (866 nm and 1064 nm, rhodamine) and the signals were shown to co-localise after
24 h incubation demonstrating that this model payload could be delivered to microglia using these novel PLGA-alkyne NPs (Figure S8).

We then went on to investigate the uptake of the alkyne NPs in *ex vivo* mouse brain slices. These slices were freshly harvested from mice pups and cultured, bridging the gap between *in vitro* cell culture and *in vivo* experiments in rodents. The slices were incubated with alkyne NPs at $2 \times 10^8$ and $2 \times 10^9$ particles mL$^{-1}$ for 24 hours before washing, fixing and imaging with SRS microscopy. SRS imaging at 2128 cm$^{-1}$ showed that there are alkyne NPs distributed throughout the tissue.

To determine if the NPs were inside cells in the brain tissue, and what type of cells these were, a multimodal imaging technique of both immunofluorescence and Raman imaging was employed. Immunofluorescence uses primary antibodies which bind to markers specific to a certain cell type, and then secondary fluorescent antibodies so that the cells can be visualised by fluorescence microscopy. We used a primary antibody to the protein IBA1, which is found specifically on microglia, and also an antibody to OLIG2 to label oligodendroglia, which are glia that would not be expected to phagocytose NPs. Figure 5A shows the immunostained tissue with microglia in red, oligodendrocytes in green and NPs in magenta. This clearly shows the association of the NPs with microglia at both concentrations tested. Z-stack analysis of the tissue also allowed depth analysis of the NP location, with Figure 5B showing a Z-stack with orthogonal views of the tissue. It is clear from this image that the NPs have been internalised by microglia, although there are many other microglia that do not contain NPs, suggesting that uptake is not as homogeneous as it was with the *in vitro* microglia cultures.
Figure 5. Imaging alkyne NPs in *ex vivo* brain slices. A) Alkyne NPs were added to *ex vivo* mouse cortical brain slices at $2 \times 10^8$ and $2 \times 10^9$ particles mL$^{-1}$ for 24 hours before washing, fixing and immunostaining. Microglia are shown in red (using antibodies against IBA1), oligodendrocytes in green (using antibodies against OLIG2) and NPs in magenta. Scale bars 20 µm. B) Orthogonal views of Z-stack images show that the NPs are inside microglia. Scale bar 20 µm.

These experiments have highlighted the clear capability of SRS to image our novel PLGA-alkyne NPs in *ex vivo* brain tissue, and show that they are preferentially taken up by microglia over oligodendroglia, which give them potential for targeted drug delivery.

CONCLUSIONS

Raman imaging, and specifically SRS microscopy, have the potential to give non-invasive information about the location and state of polymeric nanocarriers in biological environments. We have developed two Raman active analogues of PLGA, one of the most commonly used
biodegradable polymers for drug delivery research with a long history and proven safety record, and have used the location of these deuterium and alkyne tags in the cell-silent region to enhance contrast of the NPs in a biological environment. As neither deuterium nor alkyne bonds are present in biological samples, the signal is certain to derive from the NPs. These NPs have been shown to be non-toxic to living cells and SRS imaging has shown that both PLGA-D and PLGA-alkyne NPs are internalized in primary rat microglia, and additionally the PLGA-alkyne NPs were imaged in mouse cortical ex vivo brain slices. We believe that the strong Raman signal obtained from the alkyne NPs gives them an especially wide ranging imaging potential in many biological applications, including live animal studies which will be a focus of our future work.

ASSOCIATED CONTENT

Supporting Information. Supporting information is available which shows the synthetic route to propargyl DOX, selected NMR spectra including DOSY NMR, aqueous stability of the NPs, time dependent analysis of PLGA-alkyne NPs in microglia, results from the cell viability assay, Z-stacks of alkyne NPs inside microglia and images showing encapsulation of rhodamine inside PLGA-alkyne NPs.

AUTHOR INFORMATION

Corresponding Author

*Alison N. Hulme, Alison.Hulme@ed.ac.uk

Present Addresses
†Michael P. Shaver, School of Materials and Henry Royce Institute, University of Manchester

**Author Contributions**

ANH, AW, VGB, MPS, SC and SVH designed the experiments. SVH, SC, ML and LZ carried out the experiments. SVH, ANH and MPS wrote the manuscript. SVH, SC, ML, LZ, MPS, VGB, AW and ANH read and edited the manuscript.

**Funding Sources**

This work was funded by the BBSRC [EASTBIO studentship to SVH; grant ref. BB/M010996/1] the EPSRC [grant to MPS, grant ref. EP/P026095/1], Cancer Research UK [grant ref: C157/A25140 and C157/A15703], and the MS Society UK.

**ACKNOWLEDGMENTS**

We thank Dr Colin Campbell for use of the spontaneous Raman microscope [UK Regenerative Medicine Platform Niche Hub, MRC grant ref. MR/K026666/1]. We thank Dr Marie Bechler, Dr Matthew Swire and Monica Kim for the gift of microglia.

**ABBREVIATIONS**

DCM, dichloromethane; NPs, nanoparticles; PBS, phosphate buffered saline; PLGA, Poly(lactic acid-co-glycolic acid); SRS, stimulated Raman scattering.

**REFERENCES**


TABLE OF CONTENTS GRAPHIC

For table of contents only.