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Multi-functionalised graphene nanoflakes as tumour-targeting theranostic drug-delivery vehicles†

Jennifer Lamb,^a Eliane Fischer,^a Martin Rosillo-Lopez,^b Christoph G. Salzmann ^b and Jason P. Holland ^{*a}

Graphene nanoflakes (GNFs) consist of a graphene sheet approximately 30 nm in diameter with a pristine aromatic system and an edge terminated with carboxylic acid groups. Their high water solubility and relative ease of functionalisation using carboxylate chemistry means that GNFs are potential scaffolds for the synthesis of theranostic agents. In this work, GNFs were multi-functionalised with derivatives of (i) a peptide-based Glu-NH-C(O)-NH-Lys ligand that binds prostate-specific membrane antigen (PSMA), (ii) a potent anti-mitotic drug (*R*)-ispinesib, (iii) the chelate desferrioxamine B (DFO), and (iv) an albumin-binding tag reported to extend pharmacokinetic half-life *in vivo*. Subsequent ⁶⁸Ga radiochemistry and experiments *in vitro* and *in vivo* were used to evaluate the performance of GNFs in theranostic drug design. Efficient ⁶⁸Ga-radiolabelling was achieved and the particle-loading of (*R*)-ispinesib and Glu-NH-C(O)-NH-Lys was confirmed using cellular assays. Using dose–response curves and FACS analysis it was shown that GNFs loaded with (*R*)-ispinesib inhibited the kinesin spindle protein (KSP) and induced G₂/M-phase cell cycle arrest. Cellular uptake and blocking experiments demonstrated that GNFs functionalised with the Glu-NH-C(O)-NH-Lys ligand showed specificity toward PSMA expressing cells (LNCaP). The distribution profile and excretion rates of ⁶⁸Ga-radiolabelled GNFs in athymic nude mice was evaluated using time–activity curves derived from dynamic positron-emission tomography (PET). Image analysis indicated that GNFs have low accumulation and retention in background tissue, with rapid renal clearance. In summary, our study shows that GNFs are suitable candidates for use in theranostic drug design.

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Introduction

The underlying concept of a ‘theranostic’ drug is to combine chemotherapeutic and diagnostic capabilities in a single compound. Such an agent would allow the distribution of the drug to be tracked in real time and the dose delivered to the lesion to be determined accurately.^{1–3} In recent years, graphene-based nanomaterials have been studied for potential use in various biological applications.^{4–9} For instance, modified graphene nanomaterials and carbon nanotubes show potential in anti-cancer treatments, as drug delivery vehicles, and in diagnostic imaging.^{10–14} If these different applications can be combined, then graphene nanomaterials could be used in the synthesis of ‘theranostics’.¹⁵

The covalent modification of graphene sheets and carbon nanotubes is chemically challenging. Disrupting the delocalised π -bonding network of the graphene sheet is thermodynamically unfavourable due to steric and electronic factors.¹⁶ Most reactions that lead to graphene or nanotube functionalisation overcome these issues by exposing the material to harsh reagents and conditions.¹⁷ As a result, the activation chemistry often forms a heterogeneous surface consisting of different graphene oxide (GO) and reduced graphene oxide (rGO) groups.^{18–20} Modification of the oxygen-containing functional groups on GOs and rGOs is relatively simple and the increased polarity improves water solubility. For example, Shi *et al.* modified rGO with polyethylene glycol (PEG), a tumour targeting vector, and a ⁶⁴Cu-radiolabelled complex, creating a targeted positron emission tomography (PET) radiotracer.¹²

While GO and rGO are promising scaffolds for drug design, the heterogeneous chemical nature of the graphene materials is an issue. Specifically, standard GO and rGO materials contain varying amounts of oxide, hydroxide and epoxide groups scattered throughout their structures. This makes it challenging to control the chemo- and regioselectivity of functionalisation reactions. In addition, heterogeneity in the starting materials

^aDepartment of Chemistry, University of Zurich, Winterthurerstrasse 190, CH-8057, Zurich, Switzerland. E-mail: jason.holland@chem.uzh.ch; Web: <http://www.hollandlab.org>; Tel: +41 44 63 53 990

^bDepartment of Chemistry, University College London, 20 Gordon Street, London WC1H 0AJ, UK

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hinders batch-to-batch reproducibility by increasing variability in the product.

In an effort to circumvent some of these issues, we recently synthesised edge-carboxylated graphene nanoflakes (GNFs) *via* the oxidative breakdown of chemical vapour deposition (CVD) multi-walled carbon nanotubes.^{21,22} Unlike GO and rGO materials, GNFs consist of a single graphene sheet approximately 30 nm in diameter with a pristine aromatic system. The GNF edge is terminated with carboxylic acid groups that can form carboxylic anhydrides (Fig. 1A).²² An atomic force microscopy image of GNFs spin-coated from aqueous dispersion onto highly oriented pyrolytic graphite is shown in Fig. 1B. The high solubility in water, negative zeta potential,²² and relative ease of functionalisation using carboxylate chemistry means that GNFs are potential platforms for the design of theranostic agents, where reactions can be restricted to the carboxylated edge.

In this work, we used pristine GNFs to synthesise a series of theranostic agents for targeted drug delivery and PET imaging. With this in mind, GNFs were decorated with several different components. Desferrioxamine B (DFO) is a well-known chelate for Fe³⁺, Zr⁴⁺ and Al³⁺ ions (among others) but also binds gallium with high affinity (formation constant, $\log \beta = 28.2$).²³ ⁶⁸Ga is a positron emitting radionuclide suitable for PET

imaging and is readily accessible *via* commercially available ⁶⁸Ge/⁶⁸Ga generators. (*R*)-Ispinesib is an extremely potent anti-mitotic, anti-cancer drug that targets the kinesin spindle protein (KSP); a motor protein that plays a critical role in mitosis by mediating centrosome separation and bipolar spindle formation.²⁴ Inhibiting KSP causes cell cycle arrest at the mitotic (M) phase, eventually leading to apoptosis.²⁵ Alongside its potency, (*R*)-ispinesib is a convenient choice of drug for GNF modification due to its free primary amine which lies outside of the KSP binding pocket and is available for chemical modification.²⁴ An additional goal was to incorporate a biologically active vector that targets a characteristic protein on prostate cancer cells. For this purpose, we selected the urea-based Glu-NH-C(O)-NH-Lys group, which has a high affinity and specificity toward prostate-specific membrane antigen (PSMA). Finally, we also functionalised GNFs with a pharmacokinetic modifying group (an albumin binding tag) that alters the interaction with systemic proteins and can modulate distribution profiles *in vivo*. In summary, the synthesis, *in vitro* and *in vivo* evaluation of theranostic GNF vehicles with up to four different functionalities was explored.

Experimental

Full experimental methods and materials are reported in the ESI.†

Results and discussion

Functionalisation of GNFs

Four different GNF constructs (**GNF-1**, **-2**, **-3** and **-4**, Fig. 2) were synthesised to evaluate the use of GNFs as a platform for the construction of theranostic agents. **GNF-1** was produced by pre-activating the carboxylated edge using HATU, and sequential conjugating to DFO *via* amide bond formation. The DFO chelate can be radiolabelled readily with ⁶⁸Ga³⁺ ions using established methods, allowing the use of radiochemistry to determine the success of the chemical reaction.²⁶ Radiolabelling (see below) confirmed that the DFO functionalisation reaction and the purification step were successful. With this information, we then used the same conjugation chemistry to functionalise GNFs with DFO, Glu-NH-C(O)-NH-Lys, and (*R*)-ispinesib to produce **GNF-2**. The free amine on all components allowed functionalisation *via* a one-pot, multi-component reaction. Purification was achieved on silica gel. Interestingly, when water is used as the mobile phase, the modified GNFs ran at the solvent front, allowing efficient separation from other compounds in the reaction mixture which were retained on silica.

GNF-3 and **GNF-4** were synthesised using the same chemistry. We hypothesised that by creating some distance between the GNF and Glu-NH-C(O)-NH-Lys, PSMA binding would be improved, and hence, increasing the specific uptake in target cells (LNCaP [PSMA +ve]). Therefore, **GNF-3** included a PEG₄ linker between the GNF and the PSMA ligand. The synthesis of Glu-NH-C(O)-NH-Lys-NH-C(O)-PEG₄-NH₂ (Scheme S1†) allowed

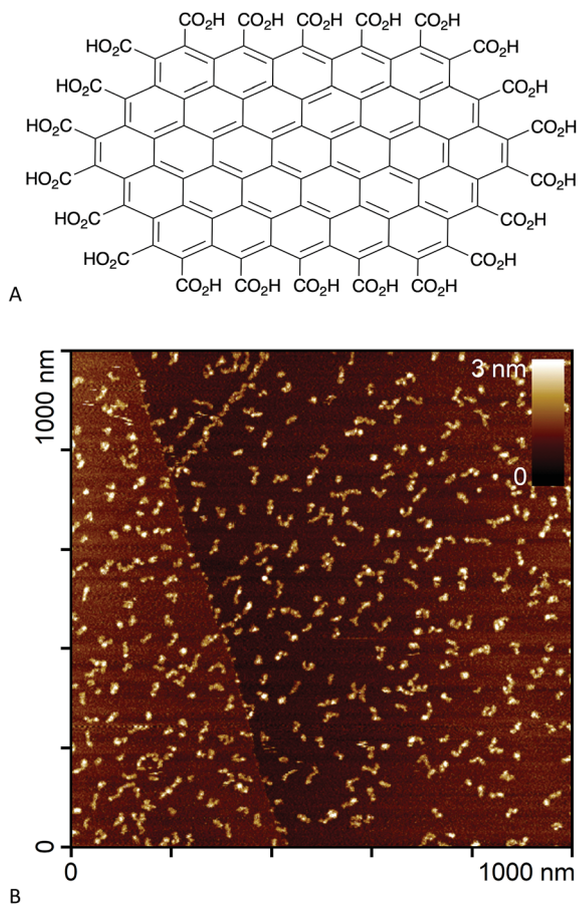


Fig. 1 (A) Chemical structure of a small GNF (not to scale). (B) Atomic force microscopy image of GNFs spin-coated onto highly oriented pyrolytic graphite.



reactions per compound). Radiochemical reactions were monitored by using radioactive instant thin-layer chromatography (radio-iTLC) and size-exclusion chromatography (SEC) (Fig. 3). Radiochemical conversions (RCCs) for the synthesis of ^{68}Ga -labelled **GNF-1**, **-2**, **-3** and **-4** were typically >97% (measured by radio-iTLC), with specific activities ranging from 8.7 ± 2.4 GBq mg^{-1} to 10.4 ± 3.0 GBq mg^{-1} (Table 1). Extending reaction times and heating the mixtures to 70°C did not improve the percentage of radiochemical conversion. Chelate-free radiolabelling of rGO with $^{64}\text{Cu}^{2+}$ has been reported.²⁸ A chelate-free approach can be beneficial as the addition of chelates to a nanoparticle (or other biological vector) can often alter the properties of the graphene, hence changing results *in vivo*.²⁹ The intrinsic radiolabelling of the GNFs with ^{68}Ga was attempted, but no labelling was observed, which also provided further confirmation that the DFO functionalisation was successful.

By using the experimentally measured molar activity of the $^{68}\text{Ga}[\text{Ga}(\text{H}_2\text{O})_6\text{Cl}_3(\text{aq.})]$ stock solutions, radioactive titrations allowed the quantification of the amount of DFO loaded per unit mass of GNF nanoparticle. Titration experiments showed that the functionalised compounds **GNF-1**, **-2**, **-3**, and **-4** contained similar amount of DFO (ranging from ~44–53 nmol DFO per mg GNF) indicating that the conjugation reactions proceeded with similar efficiency for each compound and that adding multiple components to the different GNF constructs did not compromise reaction efficiency (Table 1, Fig. S1†).

Estimates of the lipophilicity of the samples were obtained by measuring distribution coefficients ($\log D$) using standard *n*-octanol/PBS (pH 7.4) partition experiments (Table 1).³⁰ As expected, radiolabelled compounds [^{68}Ga]**GNF-1**, **-2**, **-3** and **-4**, were found to have negative $\log D$ values in the range -1.75 to -3.34 . Functionalisation of the GNF nanoparticles with Glu-NH-C(O)-NH-Lys and (*R*)-ispinesib (**GNF-2**) reduced the hydrophilicity ($\log D$ value of -2.48 ± 0.01 , $n = 3$), which suggests that the lipophilic quinazolinone drug was loaded onto the particles. For **GNF-3**, the presence of the PEG₄-spacer between the nanoparticle and the Glu-NH-C(O)-NH-Lys group increased the hydrophilicity ($\log D = -3.14 \pm 0.06$, $n = 3$). Also, as anticipated, loading the nanoparticles with the *N*-(4-aminobutyl)-6-(4-iodophenyl)butylamide tag made construct **GNF-4** less hydrophilic ($\log D = -1.75 \pm 0.07$, $n = 3$).

Stability studies *in vitro* confirmed that the ^{68}Ga -radioactivity remained bound to the particles when incubated in solutions of saline (2 h), PBS (2 h) or human serum (2.5 h) at 37°C (Tables S1 and S2, Fig. S2†).

Cellular studies

To determine the presence and activity of the (*R*)-ispinesib and Glu-NH-C(O)-NH-Lys on **GNF-2**, **-3**, and **-4**, a range of different cellular assays were performed. The PSMA expressing LNCaP cell line was used in all cell assays with PC-3 (PSMA negative) cells used in appropriate control experiments.

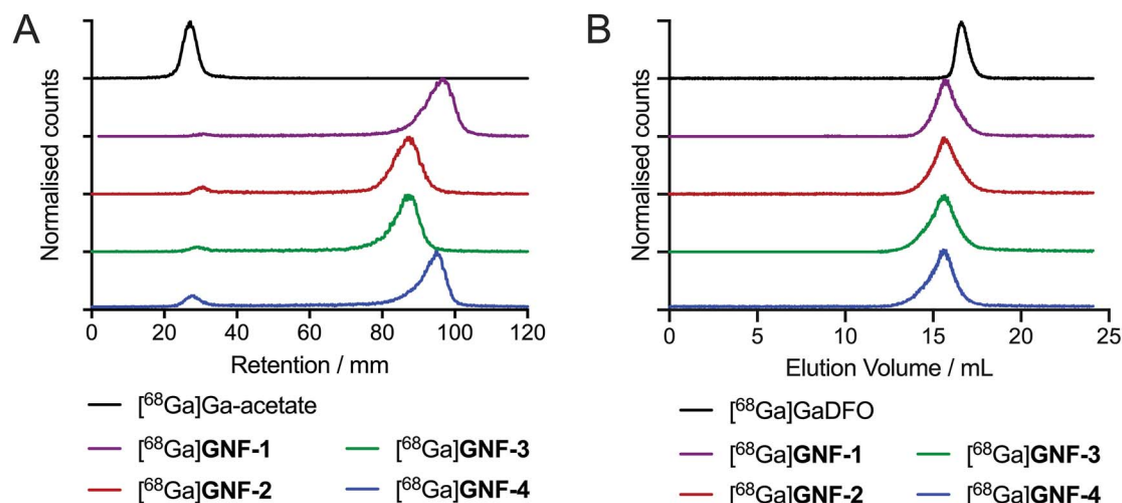


Fig. 3 Radiochemical data on the ^{68}Ga -radiolabelling of **GNF-1** to **-4**. (A) Radio-iTLC with silica gel as stationary phase and H_2O as mobile phase. (B) Radio-SEC-HPLC with 200 mM arginine in PBS as mobile phase.

Table 1 Characterisation data associated with [^{68}Ga]**GNF-1**, **-2**, **-3** and **-4**

	[^{68}Ga] GNF-1	[^{68}Ga] GNF-2	[^{68}Ga] GNF-3	[^{68}Ga] GNF-4
RCC/%	99.1 ± 0.7	98.6 ± 0.2	97.4 ± 0.6	97.3 ± 0.4
Specific activity/GBq mg^{-1}	9.08 ± 2.59	8.79 ± 0.4	10.41 ± 2.98	8.7 ± 2.44
Mole of DFO per mass of GNF/nmol mg^{-1}	46.1 ± 13.1	44.6 ± 2.26	52.8 ± 15.1	44.4 ± 12.4
$\log D$ value	-3.34 ± 0.02	-2.48 ± 0.01	-3.14 ± 0.06	-1.75 ± 0.07
Zeta (ζ) potential/mV	-36.0 ± 4.87	-39.5 ± 8.12	-60.7 ± 10.3	-42.3 ± 7.80



Anti-proliferation assays. Cellular anti-proliferation assays (MTT) were used to measure growth inhibition induced by incubation of LNCaP cells with the different GNF constructs at varying concentrations. In positive control experiments, a classic biphasic profile was observed after incubating cells for 48 h with (*R*)-ispinesib (Fig. S3†).²⁴ Analysis of the growth curves gave two distinct values for the growth-inhibition (GI_{50}) values with a $GI_{50}(1)$ of $3.32 \pm 2.31 \mu\text{M}$ and a second $GI_{50}(2)$ value of $0.7 \pm 26.7 \text{ nM}$. At high concentrations, the compound is cytotoxic to cells which corresponds to the first $GI_{50}(1)$ value. At lower concentrations, a plateau region is observed in which the cells are cytostatic with cell cycle arrest occurring in the G_2/M phase. As the drug concentration decreases into the nanomolar range, a second step is observed corresponding to the $GI_{50}(2)$ value, followed by recovery of cellular proliferation to control levels.

To determine the success of the functionalisation of the GNF with (*R*)-ispinesib, we exposed cells to **GNF-2** to observe the effect on cellular proliferation. **GNF-2** exhibited a similar biphasic profile to the drug alone, indicating that the nanoflake was successfully modified and was also a potent inhibitor of cellular growth and proliferation (Fig. 4A). To determine if further functionalisation of a graphene nanoflake with PEG₄ linkers and additional pharmacokinetic modifying groups (the albumin binder) had an impact on the anti-proliferative activity, equivalent MTT assays were conducted using **GNF-3** and **GNF-4** (Fig. 4B and C, respectively). Again, characteristic biphasic profiles were observed which suggested more extensive derivatisation of the nanoflakes did not alter the drug loading capacity or biological activity. Similar to (*R*)-ispinesib, two separate growth inhibition values could be extracted from the biphasic curves giving values centred in the micro- ($GI_{50}(1)/\mu\text{g mL}^{-1}$) and nano- ($GI_{50}(2)/\text{ng mL}^{-1}$) mass concentration ranges for **GNF-2**, -3 and -4 (Table 2). It is worth noting that anti-proliferation assays with (*R*)-ispinesib alone require the use of DMSO to solubilise the compound. The high aqueous solubility of GNFs meant that DMSO was unnecessary when using the GNF constructs. This fact provides additional support for development of nanoflakes as potential drug-delivery vehicles.

Cell cycle analysis. Although the biphasic profile in MTT assays is a characteristic feature of KSP inhibition using (*R*)-

Table 2 GI_{50} values of **GNF-2**, -3, and -4 extracted from the biphasic profiles observed in the anti-proliferation assays

	GNF-2	GNF-3	GNF-4
$\log(GI_{50}(1)/\text{mg mL}^{-1})$	-1.11 ± 0.41	-0.63 ± 0.50	-1.02 ± 0.18
$\log(GI_{50}(2)/\text{mg mL}^{-1})$	-5.70 ± 0.11	-5.73 ± 0.73	-5.52 ± 0.18

ispinesib, the mechanism of cell cycle inhibition was confirmed by using fluorescence-assisted cellular sorting (FACS). After treatment of LNCaP cells for 36 h with two different concentrations of GNF constructs, (12 ng mL^{-1} and 120 ng mL^{-1} of GNF, **GNF-2**, -3 and -4) the DNA content was measured by staining cells with propidium iodide (PI). FACS analysis showed that samples treated with **GNF-2**, -3 or -4 exhibited an increase in the proportion of cells in the G_2/M phase compared with either vehicle-treated cells or cells treated with GNF alone (Fig. 5, Table 3). For example, cells treated with 120 ng mL^{-1} **GNF-2** showed an increased number of cells in the G_2/M phase ($\sim 87\%$) in comparison to control samples which gave relative G_2/M phase populations of 9% in untreated samples, and 7% in GNF treated samples. Interestingly, the proportion of cells in the G_2/M phase was also seen by FACS to be concentration dependent. For example, for LNCaP cells treated with a lower concentration of **GNF-2** (12 ng mL^{-1}), the percentage of cells in G_2/M phase reduced to 29%. A similar trend was observed for **GNF-3** and **GNF-4**. This observation is consistent with the biphasic curve seen in the anti-proliferation assay. Collectively, the anti-proliferation assays and FACS data support the conclusion that **GNF-2**, -3 and -4 were successfully functionalised with (*R*)-ispinesib, and that the drug remained active toward KSP inhibition.

Confocal microscopy. As a final confirmation that the mechanism of G_2/M phase arrest was associated with KSP inhibition, confocal microscopy images of cells that were incubated with GNF, **GNF-2**, -3 or -4 (and controls) were obtained (Fig. 6). Untreated samples and slides treated with the control GNF sample showed predominantly cells in interphase region of the cell cycle with DNA (stained with Hoechst 33342) concentrated in a well-defined nucleus. These control

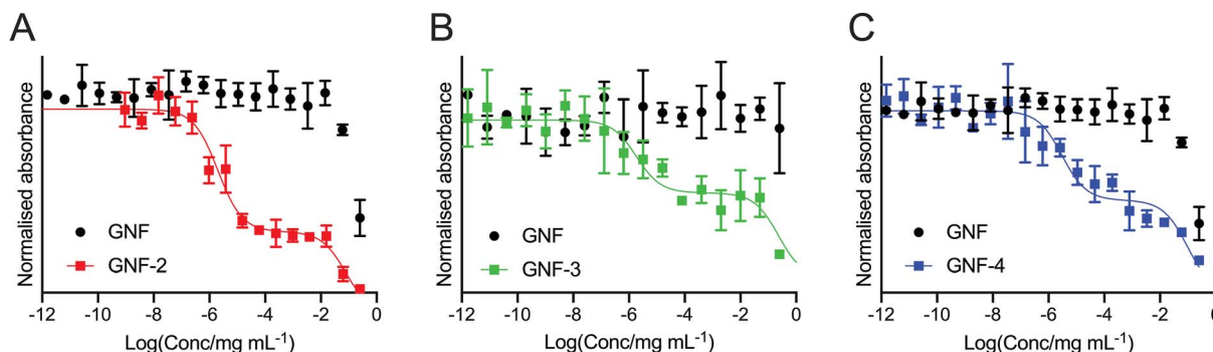


Fig. 4 Cellular anti-proliferation (MTT) curves showing the response of LNCaP (PSMA +ve) cells to treatment with (A) **GNF-2**, (B) **GNF-3**, and (C) **GNF-4**. In each plot, the black data points correspond to treatment of the LNCaP cells with the native (non-functionalised) GNF in the same formulation as the active compounds.



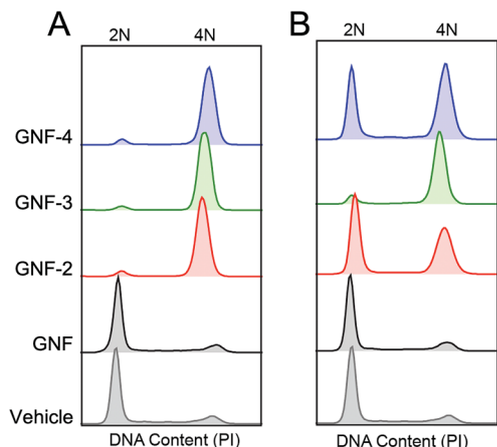


Fig. 5 Cell cycle analysis derived from FACS analysis with concentrations (A) 120 ng mL^{-1} and (B) 12 ng mL^{-1} of compounds. 2N refers to cells in the G_0/G_1 phase whilst 4N refers to cells in the G_2/M phase.

Table 3 Percentage of cells in the G_0/G_1 , S and G_2/M phase. Data extracted from FACS cell cycle analysis using the Flowjo software (Watson model)

Sample	Mass concentration/ ng mL^{-1}	Cells associated with each cell cycle phase/%		
		G_0/G_1	S	G_2/M
Vehicle	N/A	74.1	15.8	8.97
GNF	120	71.2	10.6	7.24
	12	69.2	11.6	10.2
GNF-2	120	4.92	5.21	87.3
	12	38.2	9.51	29.3
GNF-3	120	4.15	4.06	89.6
	12	4.43	10.8	78.7
GNF-4	120	5.79	6.49	77.6
	12	36.4	12.5	36.7

experiments showed no evidence of G_2/M phase arrest and confirmed the results from the FACS which showed that GNF particles (or the vehicle) alone do not interfere with the cell. In comparison, cells treated with GNF-2, -3 or -4 exhibited a characteristic punctate pattern of DNA distributed throughout the cell with no defined nucleus. The phenotype observed in these confocal cellular fluorescence microscopy data is consistent with previous reports on the mechanism of action of (*R*)-ispinesib in various cancer cells.^{24,31,32}

Cellular association assays. PSMA (also known as glutamate carboxypeptidase II or *N*-acetyl-L-aspartyl-L-glutamate peptidase I) is a membrane-bound zinc metalloenzyme which is upregulated in many prostate cancer cells.³³ The Glu-NH-C(O)-NH-Lys group is a well-documented inhibitor of PSMA and many urea-based, small-molecule radiotracers have tested *in vivo* and in human trials.^{26,34–37} To determine if GNF-2, -3 and -4 were successfully functionalised with the PSMA binding ligand, radioactive cellular binding and blocking assays were conducted using LNCaP cells (PSMA +ve) (Fig. 7A). [^{68}Ga]GNF-2, [^{68}Ga]GNF-3 and [^{68}Ga]GNF-4 showed an increased uptake in LNCaP cells compared with [^{68}Ga]

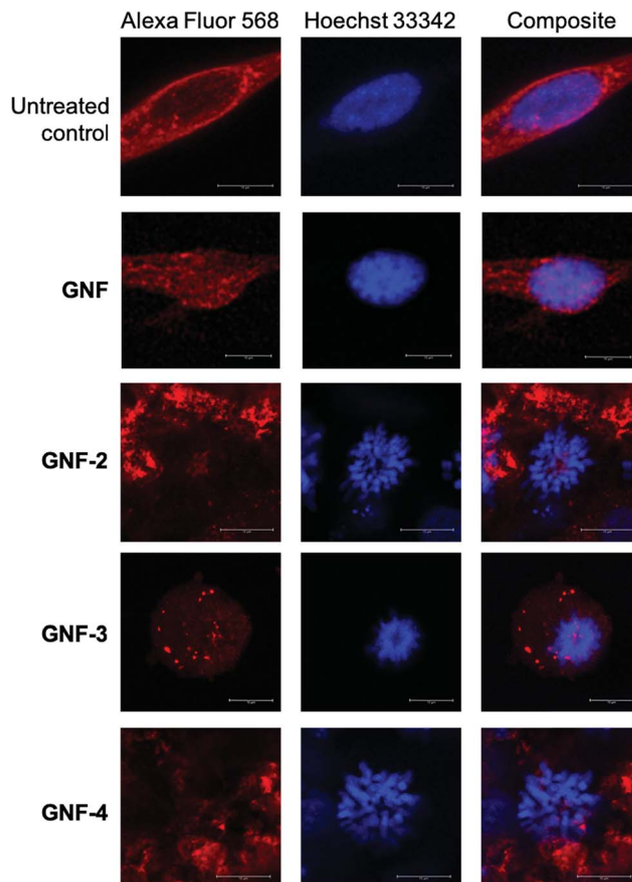


Fig. 6 Confocal microscopy images showing the effects of treatment of LNCaP cells with GNF, GNF-2, GNF-3, GNF-4. Microtubules stained with Alexa Fluor 568 (red) and DNA stained with Hoechst 33342 (blue). (Inset) Scale bars correspond to $10 \mu\text{m}$.

GNF-1. PC-3 cells (PSMA -ve) were also used as a negative control cell line. A statistically significant decrease in cellular associated activity was observed in PC-3 cells compared with the activity accumulation in the LNCaP cells for [^{68}Ga]GNF-2 ($p \leq 0.01$), [^{68}Ga]GNF-3 ($p \leq 0.001$) and [^{68}Ga]GNF-4 ($p \leq 0.0001$). These results were encouraging and suggested that functionalisation of GNFs with tumour-targeting groups was successful. However, a comparison between the LNCaP and PC-3 cell lines showed that the GNF constructs also exhibit non-specific binding which accounted for up to $\sim 50\%$ of the cell-associated activity. For instance, this non-specific binding component can be seen in the relative amount of [^{68}Ga]GNF-1 versus the 'targeted' GNF constructs. Non-specific binding is difficult to circumvent and is likely due to the interaction of the GNFs with cellular proteins, a phenomenon that has been observed for other nanographene materials.^{38,39}

We hypothesised that increasing the distance between the GNFs and the Glu-NH-C(O)-NH-Lys targeting vector using a PEG₄ linker may increase the cellular uptake and specificity for LNCaP cells. However, no difference was observed between the cell-associated uptake of [^{68}Ga]GNF-2 and that [^{68}Ga]GNF-3 (Fig. 7A). It is possible that the short PEG₄ chain is insufficient to shift the uptake mechanism away from non-specific



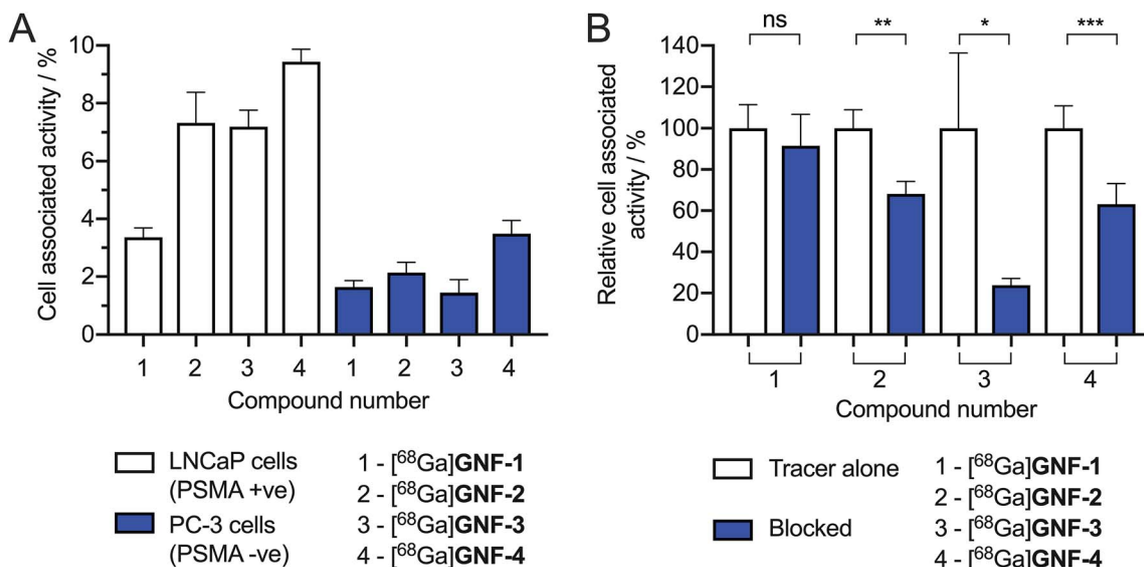


Fig. 7 (A) Cellular binding assay of [^{68}Ga]GNF-1–4 with the LNCaP (PSMA +ve) and PC-3 (PSMA –ve) cell lines. Data given as the percentage of activity bound normalised per 1 mg mL $^{-1}$ of total protein. All cell media contained 0.1% azide and protein content was measured using a BCA assay. For the LNCaP cell line: [^{68}Ga]GNF-1 vs. [^{68}Ga]GNF-2, $p \leq 0.01$; [^{68}Ga]GNF-1 vs. [^{68}Ga]GNF-3, $p \leq 0.001$; [^{68}Ga]GNF-1 vs. [^{68}Ga]GNF-4, $p \leq 0.0001$. Comparison of cell lines: PC-3 vs. LNCaP [^{68}Ga]GNF-2, $p \leq 0.01$; [^{68}Ga]GNF-3, $p \leq 0.001$; and [^{68}Ga]GNF-4, $p \leq 0.0001$. (B) Blocking assay with the LNCaP (PSMA +ve) cell line pre-treated with free Glu-NH-C(O)-NH-Lys ligand (5 μM) before addition of radiotracers. Note: Student's t -test analysis: ns = not significant, * = P -value < 0.05, ** = P -value < 0.01, *** = P -value < 0.001.

accumulation and toward specific PSMA-mediated binding. Also of note, the slightly higher cell-associated activity observed for [^{68}Ga]GNF-4 in LNCaP cells was simply attributed to an increased non-specific uptake which was also seen in the data using PC-3 cells.

Cellular blocking assays. The cellular association assays suggested that a two-state mechanism of cellular binding, involving both specific and non-specific accumulation, was operating for the targeted constructs GNF-2, -3, and -4. Therefore, to provide further confirmation of the presence of the Glu-NH-C(O)-NH-Lys group on GNF-2, -3, and -4, and to evaluate the fraction of specific binding observed using the LNCaP cells, blocking studies were performed. LNCaP cells were first incubated with ~ 1000 -fold excess of the free Glu-NH-C(O)-NH-Lys ligand (5 μM) for 1 h prior to the addition of radiotracers (Fig. 7B). The blocking studies confirmed that the radioactivity associated with the LNCaP cells could be reduced by $32 \pm 9\%$ for [^{68}Ga]GNF-2, $76 \pm 36\%$ for [^{68}Ga]GNF-3 and $37 \pm 10\%$ for [^{68}Ga]GNF-4 (Fig. 7B). Whilst the relatively high fraction of non-specific binding indicated that more work is required to optimise these constructs, the blocking experiments confirmed that GNFs can be targeted toward specific biomarkers on cancer cells.

In vivo studies

Next, to gauge the potential of using nanoflakes *in vivo*, we performed dynamic PET imaging to measure the pharmacokinetic (PK) distribution and excretion profiles of [^{68}Ga]GNF-3 and [^{68}Ga]GNF-4 in a limited number of athymic nude mice bearing subcutaneous LNCaP tumours ($n \geq 3$ mice per compound). The main goal was to measure time-activity curves

to obtain a baseline PK profile of GNFs, and to evaluate if functionalisation of the GNF constructs with an albumin binding tag could modulate blood pool concentrations, circulation times and excretion pathways.

Pilot PET imaging experiments indicated that the small size and high hydrophilicity of the [^{68}Ga]GNF-3 and [^{68}Ga]GNF-4 facilitated rapid clearance from the blood pool by a renal excretion pathway and elimination through the bladder (Fig. 8). The observation that [^{68}Ga]GNF-3 and [^{68}Ga]GNF-4 are cleared through the kidneys is an interesting finding that may help minimise radiation burden to background tissues and facilitate further optimisation of radiotracers based on GNFs. Time-activity curves (TACs) showed [^{68}Ga]GNF-3 was essentially first-pass extracted from the blood pool by the kidneys. Peak uptake in the kidney reached $38 \pm 4\%$ ID cm^{-3} within in the first two minutes post-administration. Concordantly, accumulation of ^{68}Ga -radioactivity in the bladder increased immediately after injection confirming that the [^{68}Ga]GNF-3 particles are not retained in the kidney (Fig. 8B and S4A†). In contrast, the peak in radioactivity in the kidneys for [^{68}Ga]GNF-4 occurred between 4–6 minutes post-administration with a similar time delay observed before activity was seen to accumulate in the bladder (Fig. 8C and S4B†). For both [^{68}Ga]GNF-3 and [^{68}Ga]GNF-4, blood pool activity (as measured by drawing volumes of interest over the left ventricle of the heart) showed a peak in the first frame after bolus injection of the radiotracer, and a rapid decrease over the imaging time window. In comparison to [^{68}Ga]GNF-3, the slight delay in extraction of the [^{68}Ga]GNF-4 activity from the blood pool by the kidney, gave a higher blood pool peak activity of $48 \pm 5\%$ ID cm^{-3} . Kinetic analysis of the TACs found that the effective half-life of [^{68}Ga]GNF-3 in the



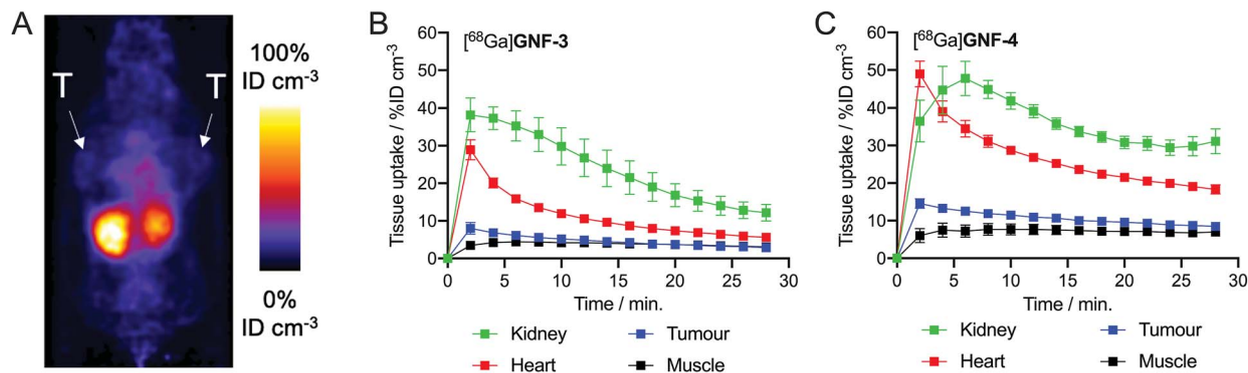


Fig. 8 PET images recorded in mice bearing LNCaP tumours, T = tumour. (A) 2 h coronal PET image recorded following injection with $[^{68}\text{Ga}]\text{GNF-4}$. TACs plotted with data extracted from 30 min dynamic PET analysis (15×2 min scans) with injection of (B) $[^{68}\text{Ga}]\text{GNF-3}$ ($n = 3$) and (C) $[^{68}\text{Ga}]\text{GNF-4}$ ($n = 4$) at $t = 2$ min VOI analysis was performed using VivoQuant™.

blood pool (heart) was 3.85 ± 0.35 min whereas for $[^{68}\text{Ga}]\text{GNF-4}$ the half-life increased to 5.22 ± 0.61 min ($p < 0.05$). These TAC data indicate that modification of the GNF particles with the albumin binding tag has a small but significant effect on prolonging the circulation time and reducing the elimination rate of $[^{68}\text{Ga}]\text{GNF-4}$.

For both $[^{68}\text{Ga}]\text{GNF-3}$ and $[^{68}\text{Ga}]\text{GNF-4}$, tumour-associated radioactivity was found to be low and non-specific. The TAC profiles of VOIs drawn over the tumours showed that the PK profile followed the same trend as the blood pool, indicating that the constructs do not show specific accumulation in the tumours on the time scale of the experiment. Factors leading to this lack of specific tumour uptake include the relatively high component of non-specific binding found for GNFs on LNCaP cells, the rapid excretion profile, and potentially, a reduced access or affinity of the Glu-NH-C(O)-NH-Lys binding ligand for the PSMA target when bound to the GNF particles. In spite of the absence of tumour uptake, the PET images provide valuable information which will aid further optimisation of the GNFs as potential drug delivery vehicles and theranostic agents. In particular, modification of the particles with higher affinity albumin tags or an increased number of tags per particle may increase circulation times, and more work on improving the cellular binding profile will likely allow increased cellular uptake and specificity.

Conclusions

GNFs were used as design scaffolds for the synthesis of targeted theranostic agents. Multi-functionalised GNFs can be prepared with a diverse array of compounds including metal-binding chelates, tumour-targeting ligands, cytotoxic drugs and pharmacokinetic modification groups. Derivatisation of GNFs with the potent drug molecule (*R*)-ispinesib gave constructs that remained pharmacologically active, and cellular assays confirmed that the mode of action was consistent with mitotic phase arrest induced by KSP motor protein inhibition. An advantage of using GNFs as a drug carrier was that it helped to solubilise (*R*)-ispinesib in water. In addition, GNF particles functionalised with the Glu-NH-C(O)-NH-Lys group exhibited

enhanced cellular uptake. Although a significant fraction of cellular association was assigned to non-specific interactions, it was found that targeted GNFs showed specificity toward PSMA expressing cells. Dynamic PET imaging found that functionalised GNFs have rapid blood pool clearance and renal excretion, but experiments also demonstrated that functionalising with an albumin-binding tag resulted in a significant modulation of the pharmacokinetic profile. Further optimisation is required to enhance the uptake kinetics and distribution *in vivo* but our study supports the conclusion that GNFs are a highly versatile platform for theranostic drug development.

Conflicts of interest

There are no conflicts to declare.

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