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## Designed ankyrin repeat proteins for detecting prostate-specific antigen expression *in vivo*†

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Late-stage prostate cancer often acquires resistance to conventional chemotherapies and transforms into a hormone-refractory, drug-resistant, and non-curative disease. Developing non-invasive tools to detect the biochemical changes that correlate with drug efficacy and reveal the onset of drug resistance would have important ramifications in managing the treatment regimen for individual patients. Here, we report the selection of new Designed Ankyrin Repeat Proteins (DARPs) that show high affinity toward prostate-specific antigen (PSA), a biomarker used in clinical monitoring of prostate cancer. Ribosome display and *in vitro* screening tools were used to select PSA-binding DARPs based on their binding affinity, selectivity, and chemical constitution. Surface plasmon resonance measurements demonstrated that the four lead candidates bind to PSA with nanomolar affinity. DARPs were site-specifically functionalised at a unique C-terminal cysteine with a hexadentate aza-nonamacrocyclic chelate (NODAGA) for subsequent radiolabelling with the positron-emitting radionuclide <sup>68</sup>Ga. [<sup>68</sup>Ga]GaNODAGA-DARPs showed high stability toward transchelation and were stable in human serum for >2 h. Radioactive binding assays using streptavidin-loaded magnetic beads confirmed that the functionalisation and radiolabelling did not compromise the specificity of [<sup>68</sup>Ga]GaNODAGA-DARPs toward PSA. Biodistribution experiments in athymic nude mice bearing subcutaneous prostate cancer xenografts derived from the LNCaP cell line revealed that three of the four [<sup>68</sup>Ga]GaNODAGA-DARPs displayed specific tumour-binding *in vivo*. For DARPin-6, tumour-uptake in the normal group reached  $4.16 \pm 0.58\%$  ID g<sup>-1</sup> ( $n = 3$ ; 2 h post-administration) and was reduced by ~50% by competitive binding with a low molar activity formulation (blocking group:  $2.47 \pm 0.42\%$  ID g<sup>-1</sup>;  $n = 3$ ;  $P$  value = 0.018). Collectively, the experimental results support the future development of new PSA-specific imaging agents for potential use in monitoring the efficacy of androgen receptor (AR)-targeted therapies.

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## Introduction

Prostate cancer (PCa) is the second most common cancer diagnosed in men world-wide.<sup>1</sup> Cellular signalling events derived from activation of the androgen receptor (AR) pathway play a prominent role in PCa development and progression.<sup>2,3</sup> Consequently, first-line treatment options for PCa patients have focused on androgen deprivation therapy (ADT) which

includes androgen ablation and AR inhibition.<sup>3–6</sup> Despite the many advances in chemotherapy, most PCa tumours develop resistance over time. The mechanisms that lead to drug resistance are not fully known and in spite of numerous second-generation drugs to treat castrate-resistant prostate cancer (CRPC), no curative treatment exists.<sup>7,8</sup>

The expression of several key protein biomarkers of PCa, including prostate-specific membrane antigen (PSMA) and prostate-specific antigen (PSA; also called gamma-seminoprotein or kallikrein 3 [KLK3]), have been shown to depend on AR activity (Fig. 1). In some PCa phenotypes, AR acts as a transcription suppressor of PSMA, and androgen deprivation therapy can lead to an increase in PSMA levels.<sup>9–11</sup> In contrast, in some patients PSA levels correlate directly with AR transcription activity, where a decrease in the measured concentration of PSA provides a diagnostic readout of the efficacy of AR inhibition.<sup>9–11</sup>

Molecular imaging techniques such as positron emission tomography (PET) can visualise spatiotemporal distribution

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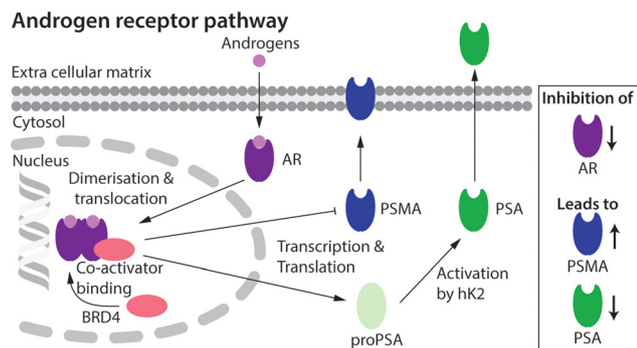


Fig. 1 Schematic showing the putative relationship between androgen-induced AR signalling and gene expression of the key PCa biomarkers, including PSMA and PSA.<sup>2,3</sup>

and quantify changes in biological processes at the tissue level, allowing disease progression or treatment efficacy to be monitored on a lesion-by-lesion basis.<sup>12,13</sup> As PSMA and PSA levels have strong links to the AR pathway<sup>9–11</sup> they are promising targets for the use in molecular imaging and therapy. Numerous PET imaging agents that delineate PSMA expression have been tested in the clinic, and clever design of the radiotracers has facilitated the transition toward PSMA-targeted radionuclide diagnosis and therapy (<sup>68</sup>Ga-PSMA-11, <sup>68</sup>Ga-gozetotide,<sup>14,15</sup> <sup>177</sup>Lu-PSMA-617, Locametz<sup>®</sup>, Novartis, Pluvicto<sup>™</sup>, Novartis<sup>16</sup>).<sup>17,18</sup> Whilst high expression levels in most PCa phenotypes mean that PSMA is the current biomarker of choice when developing radiotracers for imaging and therapy, several drawbacks remain. First, PSMA expression is not entirely PCa-specific and high levels have been identified in the salivary glands, kidneys, nerve endings, and other sites of neoangiogenesis.<sup>19,20</sup> This off-target expression is less problematic for imaging where low chemical and radiochemical doses are administered, but can cause severe, dose-limiting side-effects when using high-dose targeted radionuclide therapy. Second, the high baseline levels of PSMA expression, and the purported inverse relation between AR signalling and PSMA expression is sub-optimal in the context of companion diagnostics where PSMA-biomarker quantification is used to monitor chemotherapeutic efficacy.<sup>18</sup> Due to the high baseline, the increase in PSMA concentration upon drug treatment is potentially too ambiguous to measure drug efficacy reliably using PET.

In contrast to the expression profile of PSMA, PSA is almost exclusively found in prostate epithelial cells.<sup>21,22</sup> PSA catalyses the proteolysis of gel proteins<sup>23</sup> and is mainly excreted into the seminal fluid where concentrations can range from 0.1 to 2.2 mg mL<sup>-1</sup>.<sup>24</sup> In the absence of disease, only a small fraction of PSA leaks into the circulatory system (~0.5 ng mL<sup>-1</sup> of total PSA in males ≤ 50 years old),<sup>24</sup> where high concentrations of protease inhibitors, like the serpins  $\alpha$ 1-antichymotrypsin (ACT) and  $\alpha$ 2-macroglobulin (A2M), form a complex with the catalytically active PSA (“free” PSA or fPSA). Serpin binding blocks the enzymatic site, inactivates PSA, and produces complexed PSA (cPSA).<sup>21,25,26</sup> Measurements of both fPSA and total PSA (tPSA) in the blood are used in the clinic as both predictive and

prognostic biomarkers to monitor PCa progression.<sup>27–29</sup> However, serum measurements of PSA are considered an imperfect biomarker of PCa, because blood pool secretion does not always reflect local expression levels in the tumours<sup>21</sup> and has been shown to vary with age, body mass index, and the presence of co-morbidities.<sup>21,30</sup> The mismatch between the comparatively low levels of circulatory PSA and the true levels expressed in PCa lesions can lead to errors in patient diagnosis and management. Therefore, complementary diagnostic techniques are required to enable a more accurate assessment of PCa progression. Non-invasive PET imaging of fPSA expression in PCa lesions has been proven to be a potential route toward monitoring treatment response in patients treated with front-line AR inhibitors.<sup>30–32</sup> However, a current limitation in utilising non-invasive detection of PSA as a companion diagnostic is that very few drugs<sup>33–35</sup> or radiotracers<sup>31,32</sup> have been developed to target PSA with high affinity and specificity.

A first step toward fPSA targeting for non-invasive detection and radioimmunotherapy was achieved by utilising the humanised monoclonal antibody (mAb) hu5A10 to image (with <sup>89</sup>Zr) and treat (with <sup>90</sup>Y or <sup>225</sup>Ac) PCa tumours in mice, and in macaques.<sup>31,32</sup> Hu5A10 binds selectively and with a high specificity and affinity to the catalytic cleft of fPSA.<sup>36,37</sup> The <sup>89</sup>Zr-DFO-hu5A10 radiotracer showed high and specific uptake in LNCaP xenografts (22.75 ± 8.2% ID g<sup>-1</sup>) which was observed to be dependent on AR-drug treatment.<sup>31</sup> These results confirmed the biochemical relationship between AR-signalling and PSA expression in the LNCaP model (Fig. 1). Therapeutic effects for groups of mice treated with <sup>90</sup>Y- or <sup>225</sup>Ac-labelled hu5A10 were observed.<sup>32</sup>

In this work, our experimental goal was to identify new Designed Ankyrin Repeat Proteins (DARPin)s for selective, high-affinity binding of PSA. DARPins are small (11–20 kDa), highly stable synthetic consensus proteins.<sup>38–40</sup> Ankyrin repeat proteins are composed of stacked ankyrin motifs, consisting of a  $\beta$ -turn followed by anti-parallel  $\alpha$ -helices. The C-terminal and N-terminal capping repeats (caps) shield a more hydrophobic internal core of the protein.<sup>38–41</sup> Target engagement occurs primarily *via* interactions with the internal repeats, which carry randomised residues, and the sequence can be selected to provide high affinity and specificity toward a given epitope.<sup>40,42</sup> Several groups have developed radiotracers based on this protein platform.<sup>40,43</sup> For example, Bragina *et al.* reported data from a phase I trial which confirmed the safety and excellent performance of a <sup>99m</sup>Tc-labelled DARPin for imaging human epidermal growth factor 2 expression in breast cancer patients.<sup>43–46</sup> Here, we demonstrate the feasibility of developing new PCa diagnostic radiotracers based on the selection of PSA-specific DARPins.

## Results and discussion

### Selection of PSA-binding DARPins

Ribosome display is an efficient technique to select high-affinity DARPin binders from a randomised library.<sup>39,40,42</sup> PSA-binding DARPins were selected in four rounds of ribosome



display. Initial selections produced a long-list of 380 binders, which were evaluated in a Förster Resonance Energy Transfer (FRET) based assay,<sup>42</sup> from which 32 FLAG-tagged DARPins (abbreviated as F-DARPins) were chosen, their sequence was determined and 27 single clones were selected for further analysis (see Methods in the ESI†). The selected F-DARPins were assessed for their binding affinity and selectivity toward PSA by using ELISA methods (Fig. S1, ESI†). The chromatographic profile of each F-DARPin measured by size-exclusion chromatography was used as a second selection criterion to refine the pool to a short-list of 12 PSA-binding F-DARPins. This short-list contained F-DARPins with a varying number of internal ankyrin repeat units ( $N = 1, 2, \text{ or } 3$ ), leading to different sizes of approximately 12, 15, or 19 kDa, respectively. In addition to the 12 different PSA-binding F-DARPins (1–12), a non-binding F-DARPin (13, E3\_5)<sup>38</sup> was used as a negative control. A summary of the exclusion criteria and associated experimental data for each of the 27 F-DARPins is presented in Table S1 (ESI†).

### Cloning and purification of DARPins with a C-terminal Cys

Cysteine functionalisation is one of the most commonly applied protein conjugation methods.<sup>47</sup> The Michael-addition reaction of a Cys sulfhydryl group with a reagent bearing an electrophilic maleimido group is a well-established, reliable, and biocompatible way to site-specifically functionalise proteins.<sup>47</sup> The DARPins libraries were previously engineered to omit cysteines from their core structure.<sup>38</sup> Here, to enable site-specific modifications by using maleimide reagents, the 12 PSA-binding F-DARPins (1–12) and control F-DARPin 13 were cloned into a vector encoding a unique Cys residue located at the C-terminus (Sequences of the 13 F-DARPins and DARPins-Cys are displayed in Fig. S2 and S3, ESI†). The vectors also contained an N-terminal His<sub>6</sub>-tag to allow protein purification by immobilised metal ion (Ni) affinity chromatography (IMAC). After ligation, the vectors were cloned into *E. coli* and the modified proteins were expressed, extracted, and purified.

An ELISA was used to evaluate the effect of introducing the Cys at the C-terminus on PSA-binding of the isolated DARPins-Cys. Data revealed that when comparing the measured absorbance ( $A_{405-540 \text{ nm}}$ ) from the ELISA for the 12 PSA-binding DARPins either with C-terminal Cys (Fig. 2, DARPins-Cys, cyan) or the initial FLAG-tag (Fig. 2, F-DARPin, light blue), the

addition of the bioconjugation handle had no impact on PSA binding. All 12 PSA-binding DARPins-Cys (1–12) proteins displayed specificity toward PSA. However, both forms of DARPins 4, 5, 7, and 11 gave significantly lower absorbance, suggesting lower affinity binding, and were excluded from further studies. We also noted that DARPins-Cys 2 showed a higher level of non-specific binding, and therefore, this construct was also excluded.

In addition to the ELISA measurements, the DARPins-Cys constructs (1–12) were subjected to SDS-PAGE analysis (ESI,† Fig. S4) and size-exclusion chromatography coupled to a multi-angle light scattering (SEC-MALS) detector (ESI,† Fig. S5, and Table S2) to assess the molecular weight and purity of the protein samples. All DARPins-Cys proteins behaved as expected on the SDS-PAGE showing a dominant band associated with the monomeric protein. Two constructs, DARPins-Cys 2 and 10, gave a major band at the anticipated molecular weight but SDS-PAGE also revealed lower purity with the presence of several higher molecular weight proteins in the samples. SEC-MALS analysis revealed that DARPins-Cys 2 contained a significant fraction of dimeric (or aggregated) protein, and DARPins-Cys 4, 5, and 7 gave major peaks corresponding to low molecular weight species that were potentially protein fragments. Collectively, these experimental data supported the exclusion of DARPins 2, 4, 5, 7, 10, and 11 from the pool.

### Functionalisation of DARPins-Cys proteins with a metal ion binding chelate

Of the remaining six PSA-binding DARPins-Cys proteins (1, 3, 6, 8, 9, and 12), for practical and economic reasons, we made a further selection to reduce the number of constructs to four. DARPins-Cys 6 (19 kDa) and 9 (12 kDa) were chosen due to their different size arising from a different number of internal ankyrin repeats ( $N = 3$ , and 1 ankyrin repeats, respectively). In addition, we selected DARPins-Cys 3 and 12, which are both of intermediate size (15 kDa with  $N = 2$  ankyrin repeats) and showed high PSA-specific binding from the ELISA measurements. DARPins-Cys proteins 1 and 8 were randomly excluded from the study. The four PSA-binding DARPins-Cys proteins 3, 6, 9, and 12, and the non-binding control DARPins-Cys (13) were then site-specifically conjugated at the Cys residue by using a hexadentate tris-aza-nonamacrocyclic chelate (NODAGA)-maleimide (details are provided in ESI,† Table S3). Prior to

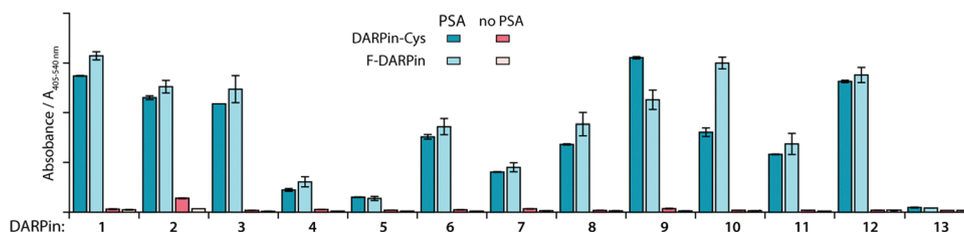


Fig. 2 Bar chart with results of ELISA measurements showing the specific binding of the 12 selected DARPins towards PSA. Data presented show the absorbance ( $A_{405-540 \text{ nm}}$ ) for PSA-binding DARPins (1–12) and the non-binding control (E3\_5, 13) either with C-terminal Cys modification (DARPins-Cys, cyan) or FLAG-tagged DARPins (F-DARPin, light blue). Red bars show the non-specific binding control where no PSA was coated on the plate. Error bars represent one standard deviation about the mean ( $n = 2$  measurements per sample).



conjugation, potentially oxidised, dimeric DARPin-Cys species were reduced by incubating protein samples with an excess of dithiothreitol (DTT, 100 eq.), thereby ensuring that the Cys sulfhydryl groups were available for the Michael addition. After removal of the excess DTT by spin filtration, the DARPin-Cys samples (3, 6, 9, 12, and control 13) were incubated with NODAGA-maleimide (2 eq.) in phosphate buffered saline (PBS, pH 7.9, 2 h, room temperature). Aliquots of the crude reaction mixtures were kept for radiochemical analysis. Excess or unreacted small-molecule reagents were removed by NAP-25 column purification, and subsequent spin concentration giving the purified chelate-functionalised NODAGA-DARPins in 21% to 56% yield (ESI,† Table S3). We note that the variation in yields reflects the different behaviour of the NODAGA-DARPins on the purification column. Successful conjugation was confirmed by high-resolution mass spectrometry (HR-MS, ESI,† Table S4 and Fig. S6–S10) and SDS-PAGE (Fig. 3).

The chemical purity of all isolated PSA-binding NODAGA-DARPins (3, 6, 9, 12) was estimated to be >95% by SDS-PAGE and >98% by SEC (Fig. 3 and ESI,† Fig. S12 light blue trace). From the SDS-PAGE data, shifts were observed in the band positions to slightly higher molecular weights for the NODAGA-DARPin products when compared with the non-functionalised DARPin-Cys constructs (Fig. 3). SDS-PAGE also illustrated the different size of the NODAGA-DARPins (12, 15, and 19 kDa). Unlike the DARPin-Cys constructs, no bands associated with the protein dimerisation were observed in the SDS-PAGE of the NODAGA-DARPin samples even without addition of reductant in

the loading buffer, indicating that quantitative functionalisation of all available sulfhydryl residues was achieved. These observations were also supported by the HR-MS data from which only peaks associated with the correct masses and isotopic distribution patterns of the functionalised NODAGA-DARPins were detected, with essentially no peaks present for the non-functionalised DARPin-Cys species. In all cases, covalent attachment of the NODAGA chelate to the proteins increased the molecular weight by  $m/z +497$ , concordant with the additional mass of the chelate and linker. Furthermore, upon addition of  $\text{Ga}(\text{NO}_3)_3$  the expected shifts by  $m/z +66$  (for the most abundant natural isotope:  $^{69}\text{Ga}$ , 60.11%; minus protons associated with the carboxylic acid groups of the NODAGA chelate) were measured in the HR-MS data of the  $^{nat}\text{Ga}$ NODAGA-DARPin samples, confirming the successful metalation with  $^{nat}\text{Ga}^{3+}$  (ESI,† Table S4 and Fig. S6–S10).

### Qualitative assessment of NODAGA-DARPin PSA-binding by native PAGE and size exclusion chromatography

PSA-binding of the functionalised NODAGA-DARPins was assessed to confirm that the addition of the chelate did not interfere with target engagement. Aliquots of the NODAGA-DARPins (3, 6, 9, 12, and control 13) were incubated with equimolar amounts of PSA at 24 °C for 30 min. For each of the NODAGA-DARPins (3, 6, 9, 12), native PAGE analysis showed a discrete shift in the protein band associated with formation of the PSA-DARPin complexes (Fig. 4(A)). For the non-binding control NODAGA-DARPin 13, no change was observed, and discrete bands were present for both free NODAGA-DARPin 13 and PSA.

The same reaction mixtures were also analysed by SEC (Fig. 4(B) and ESI,† Fig. S11). As an example, NODAGA-DARPin 6 gave a major peak with a retention time  $R_t$  of 6.66 min. (Fig. 4(B), cyan trace), which under the chromatography conditions overlapped with the peak observed for the free PSA protein (Fig. 4(B), black trace, peak  $R_t = 6.67$  min.). After incubating NODAGA-DARPin 6 with PSA, the peak observed in the SEC chromatogram (Fig. 4(B), purple trace) shifted to shorter retention time ( $R_t = 6.17$  min.), which is characteristic of the formation of the higher molecular weight NODAGA-DARPin-PSA binary complex (total  $M_w = \sim 52$  kDa). The SEC data also suggest that the NODAGA-DARPins bind to PSA with high affinity to produce complexes that are kinetically stable under the chromatographic conditions. For the non-binding control NODAGA-DARPin 13, no shift in the  $R_t$  of the peaks was observed, which also provided support that the chromatographic shifts observed for NODAGA-DARPins (3, 6, 9, and 12) reflect specific, high-affinity binding to PSA (ESI,† Fig. S11).

### Measurement of PSA-binding affinities by surface plasmon resonance

The PSA-binding affinity for the various DARPin species was evaluated by using surface plasmon resonance (SPR) single-cycle kinetic measurements. SPR was used to estimate  $k_{on}$  (rate constant of association,  $\text{M}^{-1} \text{s}^{-1}$ ),  $k_{off}$  (rate constant of dissociation,  $\text{s}^{-1}$ ), and  $K_D$  (dissociation constant,  $\text{M}^{-1}$ ) of the

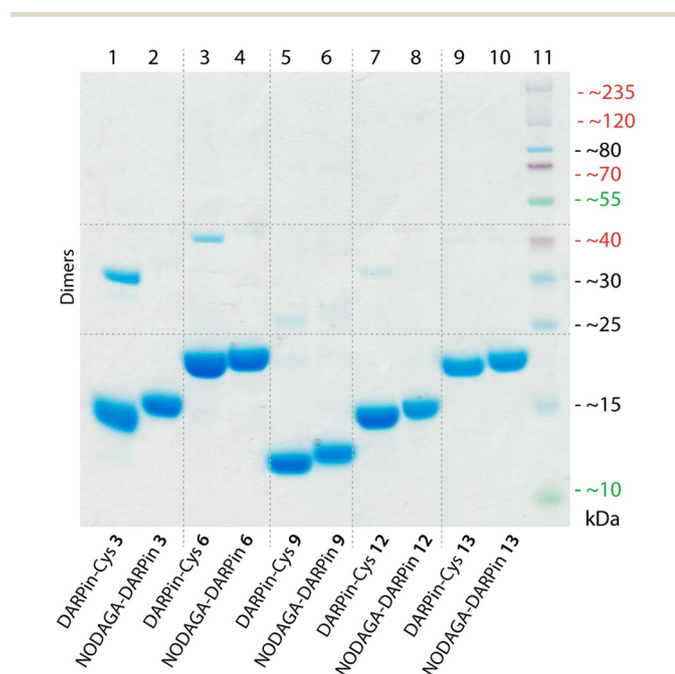
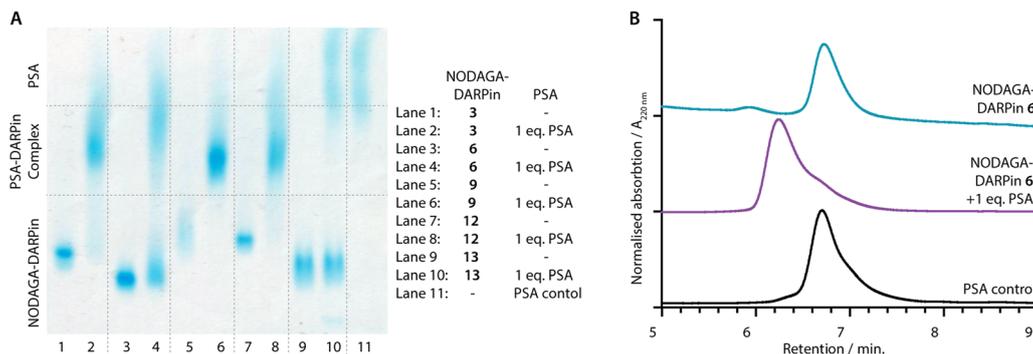


Fig. 3 SDS-polyacrylamide gel showing DARPin-Cys 3, 6, 9, 12, and 13 (lanes 1, 3, 5, 7, and 9, respectively) and the equivalent functionalised NODAGA-DARPins (lanes 2, 4, 6, 8, and 10, respectively). Note: protein bands are stained with Coomassie blue, and the molecular weight ladder is shown in lane 11.





**Fig. 4** (A) Native PAGE analysis of PSA-binding: NODAGA-DARPin **3**, **6**, **9**, **12**, and **13** (lanes 1, 3, 5, 7 and 9, respectively) show bands in the lower third of the gel. NODAGA-DARPin **3**, **6**, **9**, and **12** incubated with 1 eq. PSA (lanes 2, 4, 6, and 8, respectively) form higher molecular weight NODAGA-DARPin-PSA complexes which appear in the centre of the gel with a broad band. The non-binding control NODAGA-DARPin **13** incubated with 1 eq. PSA (lane 10) does not form a complex and two distinct bands consistent with unbound proteins were detected. Note: Protein bands are stained with Coomassie blue, and PSA as a control is displayed in lane 11. (B) Analytical SEC analysis of PSA-binding: The NODAGA-DARPin **6** (cyan) was incubated with 1 eq. of PSA (black) giving a higher molecular weight complex with slightly shorter retention time (purple). Chromatographic profiles were measured by electronic absorption at 220 nm.

DARPin-Cys and NODAGA-DARPin species with either PSA (Table 1), the PSA- $\alpha$ 1-antichymotrypsin (PSA-ACT) complex, or ACT alone (ESI,† Tables S5, S6 and Fig. S12–S15). The DARPin-Cys and NODAGA-DARPin were immobilised *via* their N-terminal His<sub>6</sub> tags.

When fitting the acquired data, we observed deviations from expected 1:1 binding kinetics. In most cases, modelling the experimental data with a sum of two exponentials (Sum2Exp) gave satisfactory fits. The human PSA analyte was obtained from natural sources and is known to contain at least 5 isoforms.<sup>10,21,48</sup> Moreover, PSA carries a single N-linked glycosylation site at Asp69 that is known to be heterogeneous,<sup>49</sup> as can also be seen in the diffuse band in Fig. 4(A). We suggest that the higher order kinetic profile observed in these measurements arises from these inhomogeneities of the analyte.

As summarised in Table 1, all DARPin-Cys and NODAGA-DARPin (**3**, **6**, **9**, and **12**) displayed comparable, nanomolar

affinity for PSA and PSA-ACT. The measured  $K_D$  values ranged from 0.9 to 43 nM (full data are presented in Tables S5 and S6 (ESI,†); representative sensorgrams with fitted curves are presented in Fig. S12–S15, ESI,†). The SPR data revealed that, in general, small differences were observed between the respective DARPin-Cys and NODAGA-DARPin pairs where in all cases  $K_D$  values for PSA binding remained in the desired low nanomolar range. For NODAGA-DARPin **3** and **6**, functionalisation with the chelate appears to improve protein binding affinity, but this conclusion is drawn with caution. Overall, we concluded that the derivatisation did not impinge on the potential use of NODAGA-DARPin (**3**, **6**, **9**, and **12**) as radiotracers.

SPR measurements were also performed using the PSA-ACT complex and ACT alone as the analytes (ESI,† Tables S5, S6 and Fig. S14, and S15). No binding was observed for any of the DARPin constructs with the ACT protein (Tables S5 and S6, ESI,†). All DARPin-Cys proteins and two NODAGA-DARPin

**Table 1** Binding affinities measured in SPR experiments<sup>a</sup>

Probe	Analyte	Model	$k_{on}/10^5 \text{ M}^{-1} \text{ s}^{-1}$	$k_{off}/10^{-3} \text{ s}^{-1}$	$K_D/\text{nM}$	$RU_{(\max)}$	$\text{Chi}^2/RU^2$
DARPin-Cys <b>3</b>	PSA	Sum2Exp	$1.515 \pm 0.003$	$2.440 \pm 0.001$	$16.11 \pm 0.06$	$117.1 \pm 0.3$	0.21
			$7.137 \pm 0.065$	$12.92 \pm 0.005$	$18.10 \pm 0.18$	$55.92 \pm 0.37$	
NODAGA-DARPin <b>3</b>	PSA	Sum2Exp	$8.824 \pm 0.417$	$0.8728 \pm 0.0073$	$0.9891 \pm 0.009$	$53.73 \pm 0.35$	0.72
			$2.243 \pm 0.034$	$4.046 \pm 0.043$	$18.04 \pm 0.33$	$25.98 \pm 0.30$	
DARPin-Cys <b>6</b>	PSA	Sum2Exp	$8.646 \pm 0.038$	$13.88 \pm 0.045$	$16.06 \pm 0.09$	$34.67 \pm 0.11$	0.066
			$1.201 \pm 0.003$	$2.498 \pm 0.004$	$20.79 \pm 0.06$	$61.15 \pm 0.09$	
NODAGA-DARPin <b>6</b>	PSA	Sum2Exp	$5.399 \pm 0.027$	$0.9626 \pm 0.0062$	$1.783 \pm 0.015$	$67.94 \pm 0.30$	0.62
			$1.432 \pm 0.041$	$6.184 \pm 0.130$	$43.18 \pm 1.54$	$16.40 \pm 0.20$	
DARPin-Cys <b>9</b>	PSA	Sum2Exp	$1.104 \pm 0.006$	$0.1460 \pm 0.0064$	$1.322 \pm 0.059$	$70.29 \pm 0.27$	0.44
			$8.924 \pm 0.632$	$2.484 \pm 0.007$	$2.783 \pm 0.021$	$105.3 \pm 0.3$	
NODAGA-DARPin <b>9</b>	PSA	Sum2Exp	$46.95 \pm 0.39$	$16.04 \pm 0.098$	$3.416 \pm 0.036$	$31.94 \pm 0.07$	0.30
			$2.428 \pm 0.010$	$2.536 \pm 0.011$	$10.44 \pm 0.06$	$29.05 \pm 0.07$	
DARPin-Cys <b>12</b>	PSA	Sum2Exp	$9.484 \pm 0.063$	$3.022 \pm 0.009$	$3.186 \pm 0.023$	$41.51 \pm 0.09$	0.069
			$0.8821 \pm 0.0043$	$0.2977 \pm 0.0057$	$3.375 \pm 0.067$	$33.91 \pm 0.08$	
NODAGA-DARPin <b>12</b>	PSA	Sum2Exp	$16.05 \pm 0.08$	$9.007 \pm 0.030$	$5.613 \pm 0.039$	$28.05 \pm 0.09$	0.21
			$1.683 \pm 0.006$	$1.438 \pm 0.005$	$8.543 \pm 0.045$	$34.69 \pm 0.09$	

<sup>a</sup> Summary of the results of single-cycle kinetic surface plasmon resonance (SPR) measurements of the DARPin-Cys and NODAGA-DARPin **3**, **6**, **9**, and **12** using PSA as analyte (full data sets and data using PSA-ACT, and ACT as analytes are given in the ESI, Tables S5 and S6). Measurements were performed on a NiHC30 chip at 20 °C in HBS buffer.



(6 and 12) displayed similar or higher affinity for the free PSA analyte *versus* the PSA-ACT complex, which suggest a small degree of selectivity for the catalytically active form of PSA. Conversely, for NODAGA-DARPin 3 and 9, similar or slightly higher binding to the PSA-ACT complex were observed. However, the degree of selectivity displayed by our current pool of NODAGA-DARPin (3, 6, 9, and 12) is likely insufficient to discriminate PSA from PSA-ACT using non-invasive imaging methods. Even though selective discrimination of free PSA and complexed (inactive) PSA-ACT could help stratify patient response to therapy, fPSA concentrations in cancerous prostate tissue are several orders of magnitude higher than total PSA levels found in circulation.<sup>24</sup> This should lead to a sufficiently high contrast, even without selectivity for fPSA.

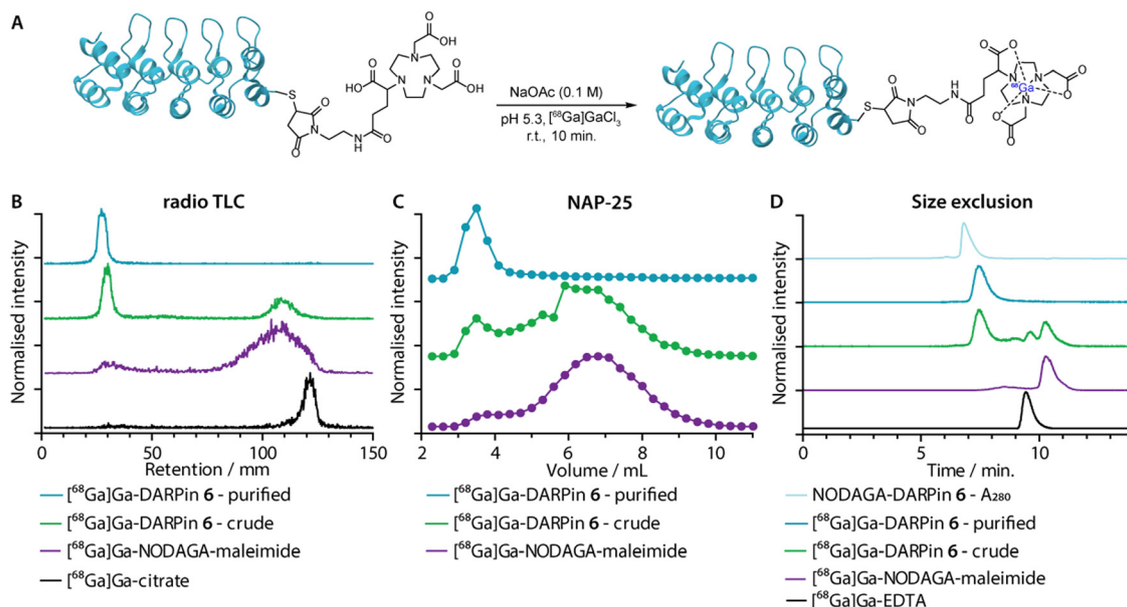
### Radiosynthesis of [<sup>68</sup>Ga]GaNODAGA-DARPin

Gallium-68 (half-life,  $t_{1/2} = 67.7$  min.; positron emission intensity,  $I(\beta^+) = 88.9\%$ )<sup>50</sup> radiolabelling of the NODAGA-DARPin was achieved by using standard radiolabelling conditions (Fig. 5(A)).<sup>51</sup> The positron-emitting radionuclide <sup>68</sup>Ga was selected because the short half-life of this radionuclide provides a complementary match with the rapid tissue uptake and excretion profiles that have been reported for other radiolabelled DARPin *in vivo*.<sup>44,46,52</sup> Briefly, an aliquot of [<sup>68</sup>Ga][Ga(H<sub>2</sub>O)<sub>6</sub>]Cl<sub>3</sub> (3–4 MBq) was added to a buffered solution of excess NODAGA-DARPin (~0.40 nmol) in NaOAc (0.1 M, pH 4.4), and the pH was adjusted to 5.3–5.5 by the addition of Na<sub>2</sub>CO<sub>3</sub> (1 M). Successful radio-metallation gave the [<sup>68</sup>Ga]GaNODAGA-DARPin species in

<10 minutes at room temperature. The [<sup>68</sup>Ga]GaNODAGA-DARPin radiotracers were obtained with radiochemical conversions >99%, and in high radiochemical purities (RCP) >99%. To determine the maximum molar activities ( $A_m$ ), NODAGA-DARPin (0.005–0.500 nmol) were titrated against a constant amount of [<sup>68</sup>Ga][Ga(H<sub>2</sub>O)<sub>6</sub>]Cl<sub>3</sub> (3–4 MBq). Radio-SEC analysis revealed  $A_m$ -values ranging from 30.7 to 54.4 MBq nmol<sup>-1</sup> of protein (ESI,† Fig. S16 and Table S7).

Radiometal ion complexation of the non-purified crude reaction mixtures as well as the NAP-25 purified NODAGA-DARPin were analysed by using three separate chromatographic methods including radioactive instant thin-layer chromatography (radio-TLC), manual NAP-25 gel chromatography, and automated radio-SEC. Representative data for <sup>68</sup>Ga-radiolabelling of NODAGA-DARPin 6 are given in Fig. 5(B)–(D), respectively. Full radiochemical data for [<sup>68</sup>Ga]GaNODAGA-DARPin radiotracers (3, 6, 9, 12 and control 13) are presented in ESI,† Fig. S16.

As an example, radio-TLC analysis showed retention of the activity at the baseline ( $R_f = 0.0$ –0.1) for both the crude and the purified samples of [<sup>68</sup>Ga]GaNODAGA-DARPin-6 (Fig. 6(B), green and cyan traces, respectively), indicative of radiolabelled protein. In the case of the crude sample, 35–45% of the activity eluted at the solvent front and is assigned to the presence of an excess of [<sup>68</sup>Ga]GaNODAGA-maleimide. After the crude NODAGA-DARPin samples were purified, subsequent quantitative <sup>68</sup>Ga-radiolabelling gave clean radio-TLC profiles whereby all the activity was retained at the baseline. For comparison, the elution profiles of the control samples confirm that the



**Fig. 5** (A) Reaction scheme showing the one-step <sup>68</sup>Ga-radiolabelling of the NODAGA-DARPin. (B) Representative radio-thin layer chromatography (radio-TLC; developed with 0.2 M, pH 4.4 aqueous sodium citrate as the mobile phase) data showing characterisation of the purified (cyan) and crude (green) samples of [<sup>68</sup>Ga]GaNODAGA-DARPin 6. The purple and black traces show control data obtained with [<sup>68</sup>Ga]GaNODAGA-maleimide and [<sup>68</sup>Ga]Ga-citrate, respectively. (C) Manual SEC analysis of the purified (cyan) and the crude (green) constructs as well as [<sup>68</sup>Ga]GaNODAGA-maleimide (purple) obtained by using NAP-25 columns eluted with PBS; (D) automated radio-SEC analysis showing a co-elution of the NODAGA-DARPin 6 UV trace (light blue) with the radiotracer of purified (cyan) and crude (green) samples of [<sup>68</sup>Ga]GaNODAGA-DARPin 6. Chromatographic profiles are also presented for the control compounds [<sup>68</sup>Ga]GaNODAGA-maleimide (purple), and [<sup>68</sup>Ga]Ga-EDTA (black), both of which show distinct retention times.



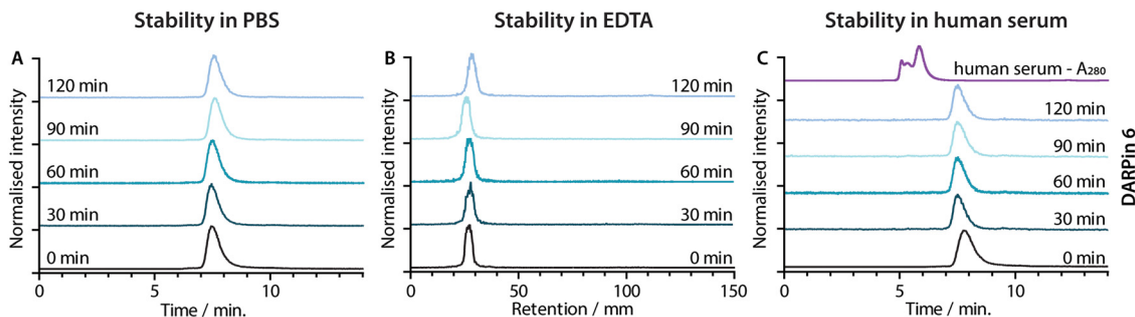


Fig. 6 Stability of [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin **6** in (A) the formulation buffer (PBS) at 37 °C assessed by radio-SEC, (B) excess EDTA at room temperature assessed by radio-TLC and (C) human serum at 37 °C assessed by radio-SEC. The black traces show the baseline data obtained at  $t = 0$  min. The fading blue traces indicate the analysis of aliquots taken at 30 min intervals from  $t = 30$ –120 min. The UV trace of human serum is shown in panel C (purple).

presence of a major peak at the baseline corresponds to the  $^{68}\text{Ga}$ -labelled protein.

Analytical size-exclusion chromatography performed by using manual NAP-25 columns showed that the purified sample of [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin-**6** (Fig. 5(C), cyan trace) gave a single peak with an elution maximum at  $\sim 3.5$  mL. These data also confirmed that the manual size-exclusion purification process successfully removed all small-molecule components from the NODAGA-DARPin conjugation step. Similarly, automated SEC chromatograms of the crude (Fig. 5(D), green trace) and purified samples of [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin **6** (Fig. 5(D), cyan trace) indicated that the radiotracer was obtained in  $>99\%$  RCP, where the peak for the product co-eluted with the electronic absorption chromatogram of NODAGA-DARPin **6** (Fig. 5(D), light blue trace, measured at 280 nm). SEC chromatograms showed the absence of any non-specifically bound 'free'  $^{68}\text{Ga}$ -ions (represented by [ $^{68}\text{Ga}$ ]Ga-EDTA), ( $R_t = 9.3$  min, Fig. 5(D), black trace) or non-conjugated [ $^{68}\text{Ga}$ ]GaNODAGA-maleimide reagent ( $R_t = 10.2$  min, Fig. 5(D), purple trace). Overall, the radiochemical data confirmed that all the  $^{68}\text{Ga}$ -labelled NODAGA-DARPins could be synthesised in sufficient radiochemical yield, chemical and radiochemical purity, and with molar activities that are suitable for applications in further biochemical studies.

### Radiotracer stability studies

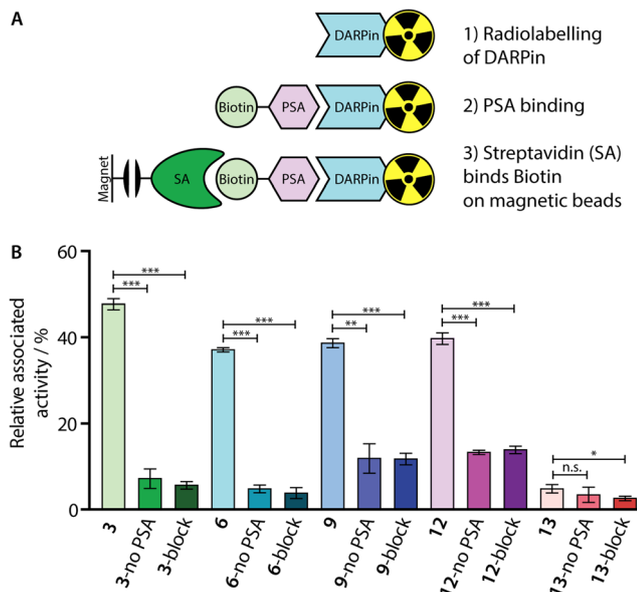
Concerns have been raised over the stability of maleimide linkages *in vivo*,<sup>53</sup> and a range of alternative bioconjugation methods have been developed.<sup>52,54–56</sup> To evaluate if the stability profiles of the [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin radiotracers were suitable for *in vivo* experiments, their radiochemical stability was measured *in vitro* over time in formulation buffer (PBS, 37 °C, pH 7.4), in chelate challenge studies using excess ethylenediamine tetraacetic acid (EDTA, 3700 eq., 37 °C, pH 7.1), and in human serum (37 °C, pH 7.4). Representative data for [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin **6** are presented in Fig. 6, and additional data for the other radiotracers are given in ESI,<sup>†</sup> Fig. S17 and Tables S8–S10. All the [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin radiotracers were found to be stable in PBS with no change in radiochemical purity over 2 h (radio-SEC data, Fig. 5(A), Fig. S17 and Table S8, ESI<sup>†</sup>). Chelate challenge experiments

using a large excess of EDTA also indicated that the [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin complex remained intact with  $<9\%$  transchelation observed after 2 h (radio-TLC data, Fig. 5(B), Fig. S17 and Table S9, ESI<sup>†</sup>). Finally, human serum challenge experiments revealed that the [ $^{68}\text{Ga}$ ]GaNODAGA-DARPins did not bind to serum proteins and were also stable with respect to loss of the  $^{68}\text{Ga}$  metal ion, or the [ $^{68}\text{Ga}$ ]GaNODAGA complex, whereby measurements of the RCP over time showed that the radiotracers remained  $>96\%$  pure after 2 h (radio-SEC data, Fig. 5(C), Fig. S17 and Table S10, ESI<sup>†</sup>). Overall, the stability studies indicated that the [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin radiotracers were suitable for further evaluation in animal models.

### Radiotracer PSA-binding *in vitro*.

To confirm that our  $^{68}\text{Ga}$ -radiolabelling conditions did not impair the specific binding of the [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin radiotracers to PSA, we developed a radioactive PSA-binding assay by using streptavidin-coated magnetic beads. An illustration of the binding assay is presented in Fig. 7(A), and a summary of the data obtained for [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin radiotracers (**3**, **6**, **9**, **12**, and control **13**) are shown in Fig. 7(B), (with further details given in ESI,<sup>†</sup> Table S11). Briefly, the [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin radiotracers were first incubated with biotinylated PSA (10 eq.) to form the [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin-PSA-biotin binary complex. Then, the mixture was incubated with streptavidin-coated magnetic beads to pull down the [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin-PSA-biotin complex. After magnetic separation, the activity associated with the beads and with the supernatant was measured, and data were normalised with respect to the total activity loaded to determine the bound fraction of [ $^{68}\text{Ga}$ ]GaNODAGA-DARPins. As a control, blocking studies were performed whereby excess DARPin-Cys (250 eq., Fig. 7(B), block) was used to saturate the available PSA-biotin. In separate controls no PSA-biotin was added to the reaction mixture (Fig. 7(B), no PSA) to allow the measurement of non-specific binding to the magnetic beads. These binding assays confirmed that the [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin radiotracers remained biochemically active with specific binding to PSA of between 37–48% of the total administered activity ( $n = 3$  independent measurements per sample). In comparison, the non-binding control [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin **13** showed only





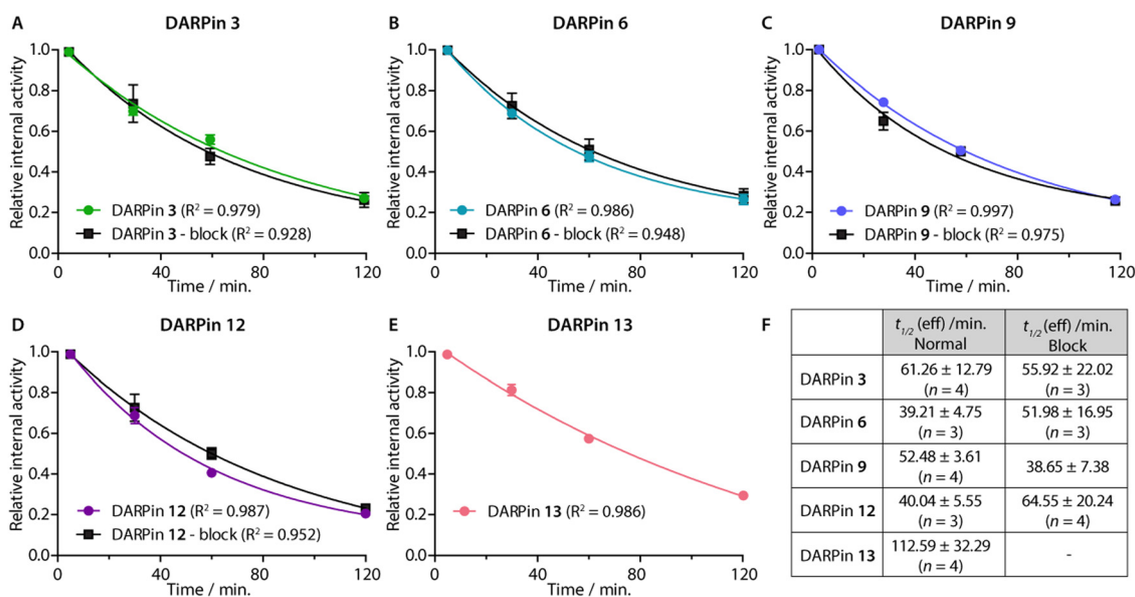
**Fig. 7** (A) Schematic representation of the *in vitro* PSA-binding assay developed to assess the biochemical viability of the  $^{68}\text{Ga}$ ]GaNODAGA-DARPin radiotracers. (B) Bar chart showing the relative activities associated with the magnetic beads for the  $^{68}\text{Ga}$ ]GaNODAGA-DARPin radiotracers with and without the addition of biotinylated PSA (no PSA), and in blocking studies containing excess non-radiolabelled DARPins (block).

$4.8 \pm 1.0\%$  non-specific binding to the streptavidin magnetic beads (Fig. 7(B), red and ESI,† Table S11). These data are consistent with the prior non-radioactive binding assays (*vide supra*) and indicate that the  $^{68}\text{Ga}$ -radiolabelling reactions did not compromise the biochemical viability or PSA-specificity of the  $^{68}\text{Ga}$ ]GaNODAGA-DARPin constructs.

### Whole-body clearance *in vivo*

After demonstrating their high stability and biochemical specificity toward PSA-binding, the whole-body clearance and tumour specificity of the  $^{68}\text{Ga}$ ]GaNODAGA-DARPins were evaluated *in vivo*. Athymic nude mice bearing subcutaneous LNCaP (PSA +ve) xenografts on the right flank ( $151.3 \pm 78.5 \text{ mm}^3$ ;  $n = 35$ ) were randomised before the study with  $n = 3$  or 4 animals per group. Synthesis of the  $^{68}\text{Ga}$ ]GaNODAGA-DARPin radiotracers (3, 6, 9, 12, and control 13) was repeated for animal studies and products were purified manually by using size-exclusion NAP-5 gel chromatography to provide the purified radiotracers formulated in sterile PBS (pH7.4), with decay-corrected radiochemical yields (d.c. RCYs) between 57% and 78% ( $n = 2/\text{each}$ , ESI,† Table S12). All  $^{68}\text{Ga}$ ]GaNODAGA-DARPins were obtained with RCP >99% and with molar activities between 7.60 and 14.02 MBq  $\text{nmol}^{-1}$  ( $n = 2/\text{each}$ , ESI,† Table S12).  $^{68}\text{Ga}$ ]GaNODAGA-DARPins 3, 6, 9, 12, and 13 were administered by intravenous tail-vein injections (150  $\mu\text{L}$ , 0.20–0.21 nmol of protein, 1.56–2.88 MBq per injection; see ESI,† Table S12). In addition to the use of the non-binding control DARPin 13, competitive (blocking) inhibition studies were performed to assess the tumour specificity of the PSA-binding radiotracers. For these blocking studies, doses of the  $^{68}\text{Ga}$ ]GaNODAGA-DARPins radiotracers (3, 6, 9, and 12) were prepared with the same amount of activity, but a reduced molar activity (0.067–0.116 MBq  $\text{nmol}^{-1}$ ,  $n = 2$ , ESI,† Table S12) by co-administration of a 100-fold excess of the equivalent non-radiolabelled DARPin-Cys.

After radiotracer administration, the whole-body effective half-lives ( $t_{1/2}(\text{eff})/\text{min}$ ) of the different  $^{68}\text{Ga}$ ]GaNODAGA-DARPin radiotracers were measured in each mouse for up to 2 h by using a dose calibrator (Fig. 8 and ESI,† Table S13).



**Fig. 8** (Panels A–E) Relative internal activities plotted against time to determine  $t_{1/2}(\text{eff})$  of the five  $^{68}\text{Ga}$ ]GaNODAGA-DARPins in the normal and blocking groups of in athymic nude mice bearing s.c. LNCaP xenografts. (F) Table summarising the calculated  $t_{1/2}(\text{eff})$  values obtained from a one-phase decay model. Note: Data were analysed in GraphPad Prism.



Data indicated that the smallest radiotracer, [<sup>68</sup>Ga]GaNODAGA-DARPin 9, had a  $t_{1/2}(\text{eff})$  value of  $52.5 \pm 3.6$  min. The intermediate sized constructs, [<sup>68</sup>Ga]GaNODAGA-DARPins 3 and 12, gave  $t_{1/2}(\text{eff})$  values of  $61.3 \pm 12.8$  min and  $40.0 \pm 5.6$  min, respectively. The two largest constructs, [<sup>68</sup>Ga]GaNODAGA-DARPin 6 and the non-binding control 13 gave measured  $t_{1/2}(\text{eff})$  values of  $39.2 \pm 4.8$  min and  $112.6 \pm 32.3$  min, respectively. In addition, changing the administered protein dose in the blocking groups also gave only small changes in the measured  $t_{1/2}(\text{eff})$  values. For example,  $t_{1/2}(\text{eff})$  values of [<sup>68</sup>Ga]GaNODAGA-DARPin 3-block and 9-block decreased ( $55.9 \pm 22.0$  min, and  $38.67 \pm 7.4$  min, respectively), but measurements showed a slight increase for 6-block ( $52.0 \pm 17.0$  min), and 12-block ( $64.6 \pm 20.2$  min). In all cases, between 70–80% of the administered activity was eliminated from the mice by 2 h post-administration. These observations suggest that the elimination rates are similar, and that other factors beyond radiotracer size and administered protein dose are likely to be involved in small differences in the tissue distribution and whole-body clearance of these [<sup>68</sup>Ga]GaNODAGA-DARPin radiotracers. Overall, these measurements of whole-body clearance are consistent with previous published data which showed rapid clearance of DARPin-based radiotracers from the circulation and elimination *via* the renal pathway, indicating that proteins whose size is smaller than the glomerular filtration unit do not follow a size-dependent elimination rate.<sup>44,45,52</sup> Provided that the radiotracers also display rapid tumour uptake and specific retention of activity, the fast excretion *via* renal clearance is a desirable

property for most molecular imaging agents, which helps to reduce background activity in healthy tissue, and increase target-to-background contrast.

### Biodistribution studies

*Ex vivo* analysis of the tissue distribution of [<sup>68</sup>Ga]GaNODAGA-DARPin (3, 6, 9, 12, and control 13) was performed at 2 h post-administration (Fig. 9, and ESI,† Table S14). Tumour-associated activity varied between the four PSA-binding [<sup>68</sup>Ga]GaNODAGA-DARPins (3, 6, 9, and 12). The smallest and largest radiotracers, [<sup>68</sup>Ga]GaNODAGA-DARPin 9 and [<sup>68</sup>Ga]GaNODAGA-DARPin 6, showed the highest tumour uptake with values of  $4.72 \pm 1.07\%$  ID  $g^{-1}$  and  $4.16 \pm 0.58\%$  ID  $g^{-1}$ , respectively. For both radiotracers, significantly higher tumour uptake was observed compared to the non-binding control radiotracer [<sup>68</sup>Ga]GaNODAGA-DARPin 13 ( $2.93 \pm 0.55\%$  ID  $g^{-1}$ , Student's *t* test *P* value  $< 0.05$  for both comparisons). The intermediate-sized radiotracers ([<sup>68</sup>Ga]GaNODAGA-DARPin 3 and 12), which both feature two ankyrin repeat units, were found to have lower tumour uptake ( $2.39 \pm 0.43\%$  ID  $g^{-1}$  and  $1.80 \pm 0.59\%$  ID  $g^{-1}$ , respectively). Blocking studies were used to measure the degree of PSA-specific binding in tumours *in vivo* where [<sup>68</sup>Ga]GaNODAGA-DARPin 6 and 9 showed a significant decrease of  $\sim 50\%$  in tumour-associated uptake ( $2.47 \pm 0.42\%$  ID  $g^{-1}$  and  $2.90 \pm 0.46\%$  ID  $g^{-1}$ , respectively [both *P* values  $< 0.05$  for the normal *versus* blocking groups]). [<sup>68</sup>Ga]GaNODAGA-DARPin 3 also showed a specific tumour-associated uptake where tumour-associated uptake in the blocking group decreased by  $\sim 39\%$

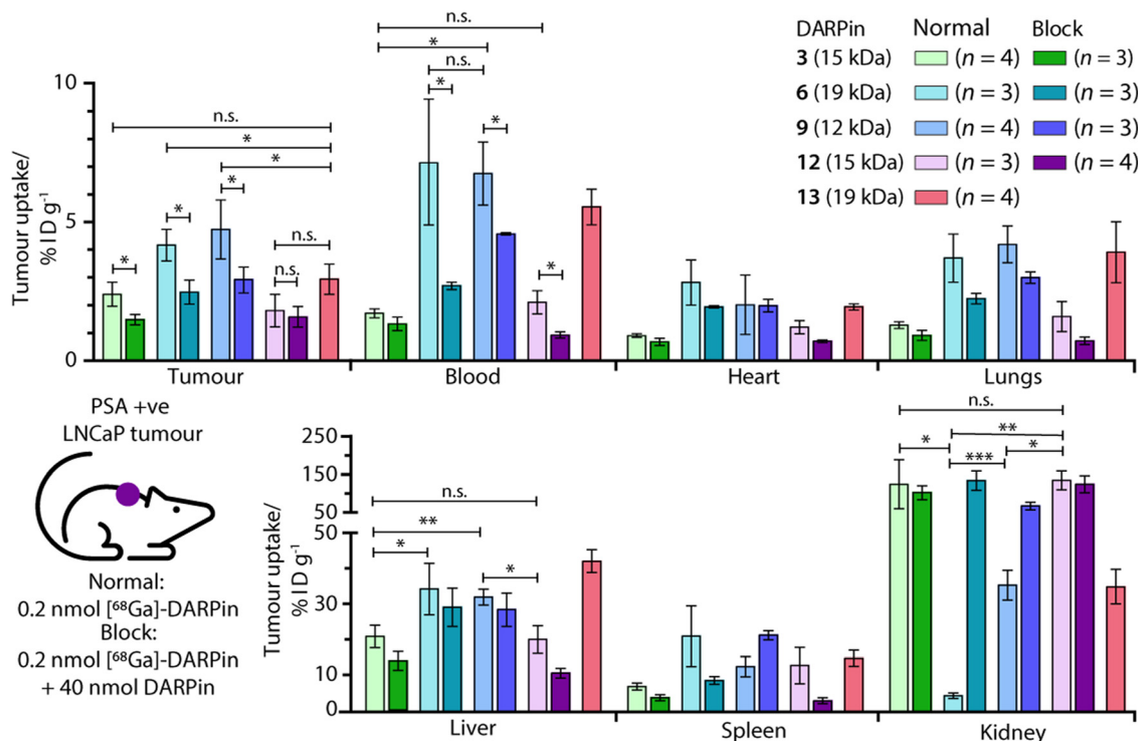


Fig. 9 Bar charts showing the *ex vivo* biodistribution analysis in selected organs for the five [<sup>68</sup>Ga]GaNODAGA-DARPin radiotracers in mice bearing subcutaneous LNCaP (PSA +ve) tumours. Statistically significant changes are displayed with \**P* < 0.05, \*\**P* < 0.01, \*\*\**P* < 0.001, and n.s. (not significant) > 0.05.



to  $1.48 \pm 0.19\%$  ID  $g^{-1}$  ( $P$  value  $< 0.05$ ). In contrast, [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin-12 gave the lowest level of tumour-associated uptake, which was found to be statistically the same as uptake observed in the blocking group ( $1.58 \pm 0.28\%$  ID  $g^{-1}$ ). The tumour associated uptake of [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin-13 was found to be  $2.93 \pm 0.55\%$  ID  $g^{-1}$ . This relatively high uptake might be explained by non-specific binding of the DARPin and is in line with tumour uptake values of 1.5–2.9% ID  $g^{-1}$  for the blocking groups of DARPins 3, 6, 9, and 12.

Unlike humans, mice do not carry functional PSA (KLK3) genes, and thus do not express PSA.<sup>57,58</sup> Moreover, in humans, PSA concentrations at prostate tissue have been reported to be  $10^6$ -fold higher in comparison with concentrations in the blood serum.<sup>24</sup> Therefore, background tissue uptake in organs due to PSA-related binding should be minimal in the reported experiments. We evaluated the radioactivity accumulation in background organs. Smaller proteins, like DARPins, are typically excreted *via* renal clearance. As expected, [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin 3 and 12, both  $\sim 15$  kDa in size, showed very similar organ distribution. One notable difference was found in the accumulation of the activity in bone, for which the uptake of [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin 3 was significantly lower at  $1.26 \pm 0.17\%$  ID  $g^{-1}$  than for 12 ( $2.35 \pm 0.39\%$  ID  $g^{-1}$ ,  $P$  value  $< 0.05$ ). For [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin 3 and 12, retention of activity in the kidneys was high ( $139.3 \pm 65.6\%$  ID  $g^{-1}$  and  $149.6 \pm 25.0\%$  ID  $g^{-1}$ , respectively). In contrast, both the larger and the smaller [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin radiotracers (6 and 9) showed considerably lower accumulation and retention of activity in the kidney ( $4.44 \pm 0.68\%$  ID  $g^{-1}$  and  $36.13 \pm 4.28\%$  ID  $g^{-1}$ , respectively). The lowest kidney uptake, and the highest tumour-to-kidney contrast ratio ( $0.94 \pm 0.19$ , Table S15, ESI<sup>†</sup>) was observed for [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin 6. Interestingly, an inverse trend was observed for liver accumulation whereby [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin 3 ( $21.31 \pm 3.24\%$  ID  $g^{-1}$ ) and 12 ( $20.41 \pm 3.99\%$  ID  $g^{-1}$ ), had lower liver retention than 6 ( $33.16 \pm 7.93\%$  ID  $g^{-1}$ ), and 9 ( $32.65 \pm 2.31\%$  ID  $g^{-1}$ ).

Blood pool activity was found to be considerably higher for [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin 6 ( $7.13 \pm 1.25\%$  ID  $g^{-1}$ ) and 9 ( $6.74 \pm 1.14\%$  ID  $g^{-1}$ ) than either 3 ( $1.71 \pm 0.16\%$  ID  $g^{-1}$ ), 12 ( $2.10 \pm 0.42\%$  ID  $g^{-1}$ ), or the non-binding control 13 ( $5.53 \pm 0.64\%$  ID  $g^{-1}$ ). Decreased clearance through the kidneys appears to correlate with prolonged circulation in the blood pool, and a concordant increase in clearance *via* the hepatobiliary pathway, contrasting DARPin 6 and 9 on the one hand with DARPins 3 and 12 on the other. All tumour-to-tissue ratios and calculated  $P$  values are given in ESI,<sup>†</sup> Tables S15–S18.

From *in vitro* data alone, it was difficult to differentiate between the four candidate PSA-binding radiotracers. *In vivo*, we observed important differences in tumour-specific uptake and tissue distribution which might arise from sequence-specific differences between the four evaluated DARPins (ESI,<sup>†</sup> Fig. S2 and S3). Prolonged circulation in the blood pool, specific tumour uptake with high contrast over most background tissues, and low levels of accumulation in the kidneys and bone, are attractive features when developing diagnostic

radiotracers. In this regard, the *ex vivo* biodistribution analysis indicated that [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin 6 and 9 are the most promising candidates for measuring specific changes in PSA-expression *in vivo*. Beyond the use of these radiotracers for non-invasive biomarker detection and quantification, these DARPin-based radiopharmaceuticals could be attractive compounds for developing molecularly targeted radiotherapeutics, provided the PSA-specific tumour-associated uptake can be further increased by a factor of  $\sim 2$ – $3$  (as previously shown).<sup>31,32</sup> Such DARPins could be attractive compounds for developing molecularly targeted radiotherapeutics using  $\beta$ -emitting radionuclides such as  $^{90}\text{Y}$ ,  $^{177}\text{Lu}$ , or  $^{188}\text{Re}$ , or  $\alpha$ -particle (or decay-chain) emitters such as  $^{211}\text{At}$ ,  $^{212}\text{Pb}$ , or  $^{225}\text{Ac}$ . Future work will focus on fine-tuning the whole-body clearance and enhancing the tumour specific uptake of DARPin-based radiotracers to facilitate diagnostic imaging and molecularly targeted radiotherapy.

## Conclusion

Here, we report on the selection and evaluation of new PSA-targeting DARPins for use in the quantification of biomarker expression profiles *in vivo*. Four PSA-binding DARPins were selected from a pool of 380 hits. The DARPins were conjugated site-specifically to a metal ion-binding chelate *via* a unique C-terminal Cys residue. *In vitro* studies, combined with radiolabelling experiments, demonstrated that all four chosen PSA-binding DARPins showed low nanomolar affinity and specific binding toward PSA. Additional radiochemical studies *in vitro* confirmed the stability, the PSA-binding specificity, and the suitability of the  $^{68}\text{Ga}$ -radiolabelled probes for use in biomarker quantification. *In vivo* experiments using tumour-bearing mice confirmed that three radiotracers showed significant and specific tumour uptake. [ $^{68}\text{Ga}$ ]GaNODAGA-DARPin radiotracers 6 and 9 performed the best, showing promising tumour uptake with high contrast over background tissue, and desirable excretion profiles with low retention of the activity in the kidneys. Collectively, these results support the development of DARPin-based radiotracers for applications in non-invasive molecular imaging and radiotherapy of PCa *via* targeting the local tissue expression of PSA.

## Data availability statement

The data that support the findings of this study are available from the corresponding author upon reasonable request.

## Author contributions

M. G. performed most of the investigation including the synthesis, radiochemistry, *in vitro* and *in vivo* testing. B. D. investigated the SEC-MALS analysis and supervised DARPin screening and selection assays which were investigated by S. F. and supported by co-workers mentioned in the acknowledgement. J. S. investigated and formally analysed the SPR



measurements. J. P. H. and A. P. supervised the project. M. G. wrote the original draft of the manuscript which was edited by B. D., A. P. and J. P. H. All authors read and approved the final version of the manuscript.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

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