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DNA-encoded chemical libraries enable the discovery of potent PSMA-ligands with substantially reduced affinity towards the GCPIII anti-target†

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Prostate-specific membrane antigen (PSMA) is a tumor-associated protein that has been successfully targeted with small organic ligands and monoclonal antibodies. Pluvicto™ is a PSMA-targeted radioligand therapeutic (RLT) recently approved by the FDA for the treatment of metastatic castration-resistant prostate cancer (2022 FDA marketing authorization). Although a large Phase III clinical trial (VISION trial) demonstrated clinical benefits in patients treated with Pluvicto™, the therapeutic window of the drug is narrowed by its undesired accumulation in healthy organs. Glutamate carboxypeptidase III (GCPIII), an enzyme sharing 70% identity with PSMA, may be responsible for the off-target accumulation of PSMA-RLTs in salivary glands and kidneys. In this work, we designed and synthesized affinity and selectivity maturation DNA-encoded chemical libraries (ASM-DELS) comprising 18'284'658 compounds that were screened in parallel against PSMA and GCPIII with the aim to identify potent and selective PSMA ligands for tumor-targeting applications. Compound A70-B104 was isolated as the most potent and selective ligand (K_D of 900 pM for PSMA, K_D of 40 nM for GCPIII). ¹⁷⁷Lu-A70-B104-DOTA, a radiolabeled derivative of compound A70-B104, presented selective accumulation in PSMA-positive cancer lesions (i.e., 7.4% ID g⁻¹, 2 hour time point) after systemic administration in tumor-bearing mice. The results of autoradiography experiments showed that ¹⁷⁷Lu-A70-B104-DOTA selectively binds to PSMA-positive cancer tissues, while negligible binding on human salivary glands was observed.

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Introduction

Radioligand therapeutics (RLTs) are innovative radiopharmaceutical products that combine the use of radionuclide payloads with small organic ligands that specifically bind to proteins expressed on cancer cells or in the tumor microenvironment.¹ This approach enables the selective accumulation of biocidal radiation within the tumor mass, thus maximizing the therapeutic effect of radioactivity while minimizing unwanted healthy tissue damage.¹ Lutathera™,² Xofigo™,³ and Pluvicto™⁴ are examples of approved products for the treatment of specific solid tumor entities. Treatment with RLTs is

usually paired with companion diagnostics that enables the selection of patients with high ligand uptake in tumor masses.⁵

Prostate-specific membrane antigen (PSMA)⁶ is one of the few examples of tumor-associated antigens which have been validated through nuclear medicine studies for the development of targeted anti-cancer RLTs.^{7,8} While the physiological function of PSMA in prostate cells is not yet fully understood,⁹⁻¹² its pronounced upregulation within prostate cancer lesions led to the development of high-affinity monoclonal antibodies and small organic ligands,¹³⁻²⁰ including (2*S*,2'*S*)-2,2'-(carbonyldiimino)dipentanedioic acid (DUPA, $IC_{50,PSMA} = 47$ nM).²¹ Further chemical elaborations on DUPA prompted the identification of PSMA-11²² and PSMA-617²³ as lead compounds for diagnostic and therapeutic applications (respectively) with excellent biodistribution profiles in tumor-bearing mice^{24,25} and patients.^{26,27}

Pluvicto™, also known as ¹⁷⁷Lu-PSMA-617, is the most recent RLT product gaining marketing authorization for the treatment of metastatic castration-resistant prostate cancer (mCRPC; FDA approval in 2022).⁴ Pluvicto™ consists of PSMA-617 conjugated to a DOTA-lutetium-177 complex as the

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radioactive payload (beta-emitter). The registrational phase III clinical trial VISION showed that treatment with Pluvicto™ in combination with the best standard of care (BSoC) enhances the overall response rate from 2% (BSoC alone) to 30%. Only a marginal benefit was observed in overall survival (15.3 months *vs.* 11.3 months with BSoC alone).⁴ Despite the efficacy demonstrated, around 12%²⁸ of individuals discontinued Pluvicto™ due to treatment-related adverse events, including xerostomia (“dry mouth”) and renal and urinary disorders. Nuclear medicine studies indicate that these side effects are due to the accumulation of the drug in healthy structures, including the salivary and lacrimal glands, kidneys, spleen, and liver.^{29,30} The undesired uptake of Pluvicto™ in healthy organs limits the possibility to increase the administered dose, thereby narrowing its therapeutic window.³¹

Our group has recently hypothesized that the off-target accumulation of Pluvicto™ may be linked to the presence of glutamate carboxypeptidase III (GCPIII) in healthy tissues.³¹ GCPIII shares 70% sequence identity with PSMA,³¹ along with a comparable enzymatic activity and analogous structure of the active site.³² We have previously shown that a fluorescent surrogate of Pluvicto™ (PSMA-617-FITC) strongly binds to recombinant GCPIII with a K_D of ~ 1 nM. This antigen is abundantly expressed in the salivary glands and kidneys (by immunofluorescence studies),³¹ and this molecular event may be responsible for the accumulation of biocidal radiation in healthy structures. Despite documented efforts to identify ligands specific to PSMA over GCPIII,^{33,34} PSMA-targeted RLTs which show high tumor-to-salivary gland and tumor-to-kidney ratios have not yet been reported.

DNA-encoded chemical library (DEL) technology is a powerful tool for the discovery of tumor-targeting ligands.^{33,35–37} DELs are large collections of small organic compounds, individually encoded by unique DNA strands that act as amplifiable identification barcodes. Binders against immobilized proteins of interest (targets) are typically isolated by affinity capture, followed by PCR amplification and high-throughput DNA sequencing (HTDS). The DEL technology has been proven reliable for *de novo* discovery and affinity maturation strategies.^{35,38–40} In the latter, a known binder for the target protein is coupled to an existing DEL with the aim of increasing its affinity to the target protein.^{38–40} In this article, we report the design, screening, and hit-validation of affinity and selectivity maturation DELs (ASM-DELs), aiming at improving ligand affinity to PSMA of previously identified HIT compounds, together with their selectivity against GCPIII (the biological “anti-target” of Pluvicto™).

Results

The structure and design of novel affinity and selectivity maturation-DELs (ASM-DELs) are shown in Fig. 1. PSMA-selective HIT₁ and HIT₂ compounds chosen for the construction of the final ASM-DELs were derived from *de novo* DEL selections previously reported by our group³³ [Fig. 1B and S33A, B†]. The design of HIT₃ (DUPA) was inspired by the seminal work of Kozikowski *et al.*²¹ [Fig. 1B]. This compound displays

high potency towards PSMA ($K_{D,PSMA} = 1$ nM), but lacks selectivity for GCPIII ($K_{D,GCPIII} = 1$ nM) [Fig. S33C†]. A number of ASM-DELs were used in our study. Some libraries featured a linear arrangement of amino acid building blocks⁴¹ (LL1, LL2, Fig. 1B). This design may be particularly attractive for the discovery of protease inhibitors, considering the peptidic nature of protease substrates. By contrast, MB libraries featured a central aromatic core with a free carboxylic acid and up to three amino acids as diversity elements (MB1, MB2, and MB3) [Fig. 1B]. Finally, the PRA DEL design³⁵ displayed a linear arrangement of two amino acids coupled to a propargyl glycine scaffold. All these libraries feature a terminal amino moiety suitable for derivatization with the HITs by amide bond formation. Detailed procedures (split-and-pool) followed during the synthesis of DEL-precursors and the final ASM-DELs are reported in ESI Sections S2 and S3.† Eighteen highly pure ASM-DELs were synthesized (>95% purity, calculated from the HPLC peak area after purification of the final pool), comprising 18'284'658 molecules. Underivatized DELs and ASM-DELs (capped with HIT₁, HIT₂, or HIT₃) were screened in parallel against site-specifically biotinylated human recombinant PSMA and GCPIII (immobilized on streptavidin beads), following state of the art screening protocols.⁴² All DEL selections were performed in duplicate.

Fig. 2 shows fingerprints of DEL selections, obtained by PCR amplification and high-throughput DNA sequencing of ASM-DEL LL2-HIT₁ against PSMA and GCPIII. The combination of A158-B51 building blocks was preferentially enriched in the PSMA selection, with a ~ 40 -fold enrichment over the background. The combination was resynthesized as a fluorescein lysine derivative (compound 1) and its binding properties to PSMA and GCPIII were studied by fluorescence polarization [Fig. 2]. Compound 1 displayed an increased affinity against PSMA ($K_{D,PSMA} = 260 \pm 5$ nM) as compared to a fluorescein lysine derivative of HIT₁ (compound 2, $K_{D,PSMA} > 2$ μ M), but a loss of PSMA-selectivity against the GCPIII anti-target was observed ($K_{D,GCPIII} = 600 \pm 3$ nM). PSMA and GCPIII selections performed with other ASM-DEL LL (LL1 and LL2) did not result in the identification of compounds with preferential PSMA binding properties [see DEL-selection fingerprints reported in ESI Section S5 and Appendix 15.1].†

The affinity maturation libraries ASM-DELs MB1, MB2 and MB3 were coupled with HIT₁, HIT₂, and HIT₃ and the resulting compound collections were screened against PSMA and GCPIII. Ligands with preferential PSMA binding were discovered only from ASM-DEL MB1-HIT₂ [Fig. 3]. The combination of HIT₂ with building block A169 was identified as selective PSMA HIT (enrichment factor of ~ 220), while the same building block combination was not enriched in GCPIII selections [Fig. 3A]. Fluorescence polarization studies revealed that combinations of MB-A169-HIT₂ (compounds 3, 5, and 7) recognized PSMA with a higher affinity compared to the unmodified HIT₂ precursor (compounds 4, 6, and 8). On-DNA fluorescence polarization studies showed that the DNA-MB-A169-HIT₂ conjugate (compound 3) bound to PSMA with a dissociation constant $K_D = 95$ nM, which was substantially improved compared to the K_D of DNA-HIT₂ and compound 4 [Fig. 3C]. Binding studies



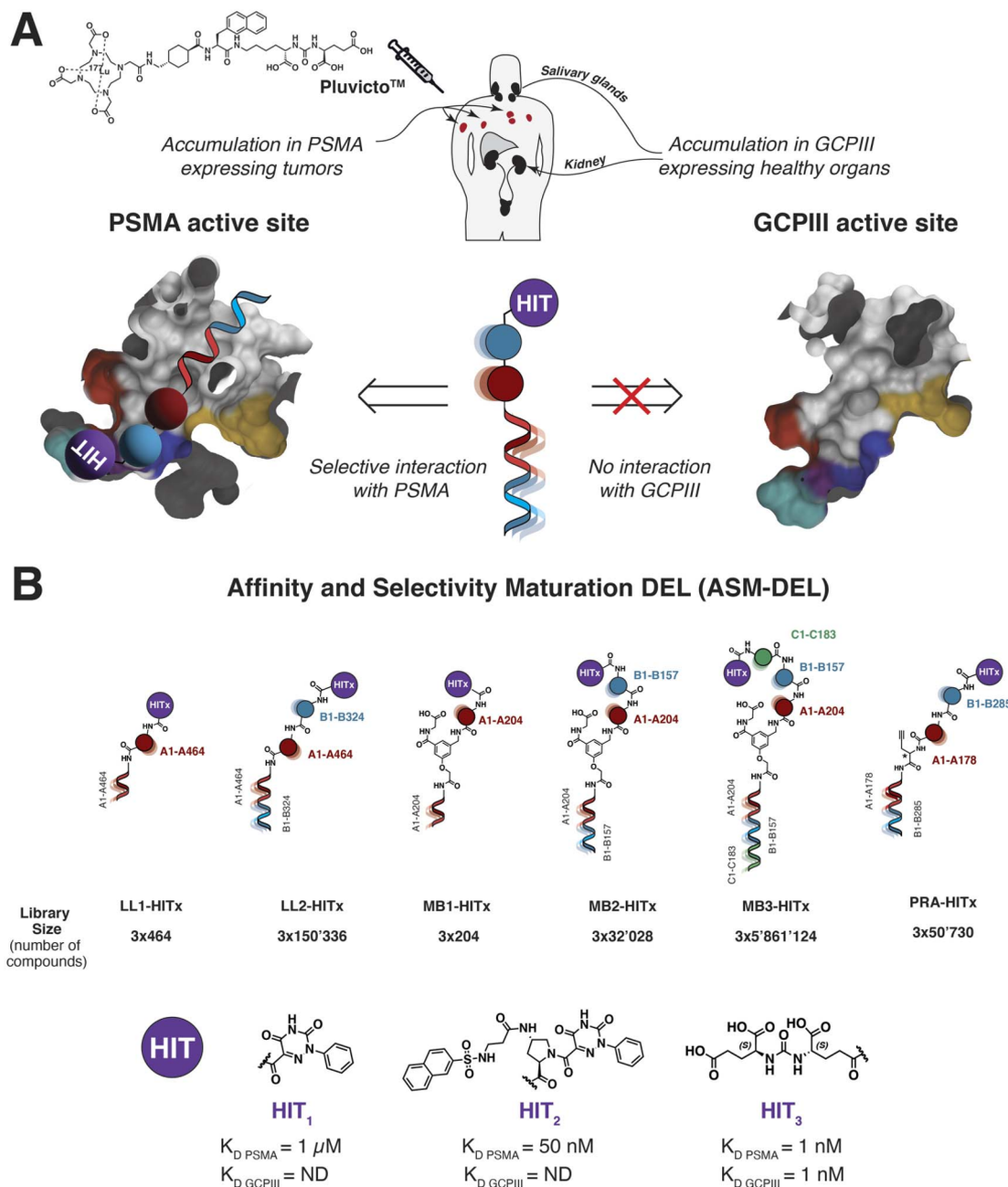


Fig. 1 (A) Pluvicto™ accumulates both in PSMA-positive tumors and in GCPIII-positive healthy organs (*i.e.*, salivary glands and kidneys). PSMA (PDB:5O5T) and GCPIII (PDB:3FED) active sites are presented in the figure as follows: S1' pockets (cyan), zinc-II active sites (purple), arginine patches (blue), S1 accessory sites (red), and arene-binding sites (yellow). (B) Design of the ASM-DELs and the chemical structure of the PSMA-HITs included in the final libraries (affinity constants measured by fluorescence polarization (FP) are reported, ESI 3.1.3, Fig. S33†). HIT₁ and HIT₂ have been discovered thanks to *de novo* DEL discovery activities,³³ while HIT₃ represents the DUPA tumor-targeting headpiece. The asterisk "*" indicates a racemic mixture.

performed by using fluorescence polarization with compounds devoid of the DNA moiety confirmed that both lysine and pegylated MB-A169-HIT₂ derivatives (compounds 5 and 7) were able to bind to PSMA with higher affinities compared to HIT₂ (compounds 6 and 8). The best binding constant was measured for Lys-MB-A169-HIT₂ (compound 5, $K_{D,PSMA}$ of 19 ± 2 nM) [Fig. 3D]. We observed that the removal of the MB scaffold from

Lys-MB-A169-HIT₂ slightly reduced the PSMA affinity of the compound (from $K_D = 19$ nM to 41 nM) [ESI 7, Fig. S120†]. Interestingly, the affinity matured MB-A169-HIT₂ derivatives (featuring a lysine or a PEG2 linker, respectively) retained the GCPIII selectivity observed for HIT₂³³ [Fig. 3E]. Fluorescein derivatives of HIT₂ and MB-A169-HIT₂ were tested by flow cytometry on PSMA-positive and PSMA-negative tumor cells



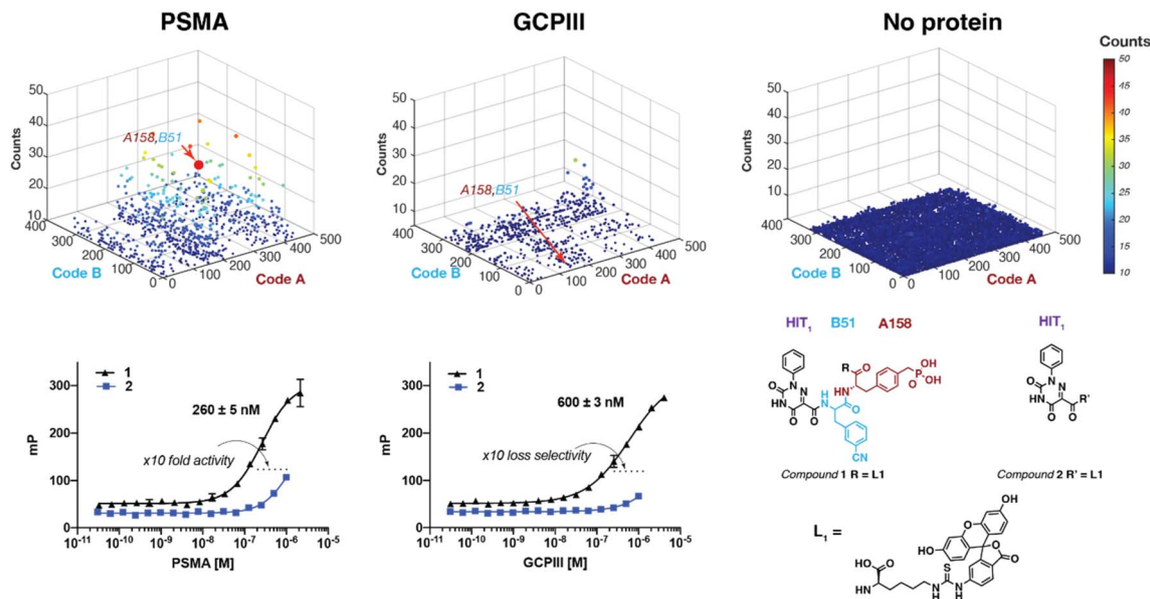


Fig. 2 LL2-HIT₁ fingerprints against PSMA, GCPIII, and underivatized beads (no protein control) used for the selection process. The selection results are displayed as three-dimensional fingerprints, where the x and y axes are the combinations of diversity elements A and B of LL2-HIT₁. Meanwhile, the vertical z-axis indicates the frequency at which a particular combination is detected. A158-B51-HIT₁ was enriched against PSMA (EF = 40) and displayed $K_D = 260 \pm 5$ nM against PSMA and $K_D = 600 \pm 3$ nM versus GCPIII, while the original compound **2** has $K_D > 1$ μ M for both target proteins.

(HT1080.hPSMA and HT1080.wt). The results, which are shown in Fig. 3F, indicated that both HIT₂ and MB-A169-HIT₂ were not able to interact with PSMA on tumor cells expressing the human antigen on their cellular membrane, as no significant fluorescence shift between HT1080.hPSMA and HT1080.wt cell samples was observed by flow cytometry. For this reason, the MB-A169-HIT₂ combination was not further investigated in *in vivo* biodistribution studies.

PRA-DEL was also functionalized with HIT₁, HIT₂, and HIT₃ and subjected to affinity-capture selections on PSMA and GCPIII [See ESI Section S5 and Appendix 15.1†]. The selection fingerprint of the PRA-HIT₃, displayed in Fig. 4A, revealed A70-B104 and A76-B104 as the most enriched combinations for PSMA, compared to selections performed with GCPIII or in the absence of a target protein. A plot of the enrichment factors towards GCPIII and PSMA (Fig. 4B) revealed that compounds A70-B104 and A76-B104 were preferentially recovered in PSMA selections (bottom right side of the graph in Fig. 4B). Fluorescent derivatives of the diastereoisomers *S* and *R* of A70-B104 (compounds **9** and **10**) were synthesized and tested by FP against recombinant PSMA and GCPIII. Similar affinity constants against PSMA were observed for the two compounds (compound **9** $K_{D,PSMA} 0.9 \pm 0.5$ nM and compound **10** $K_{D,PSMA} 1.3 \pm 0.7$ nM) and the original HIT₃ (compound **12** $K_{D,PSMA} 0.9 \pm 0.3$ nM) [Fig. 4C]. By contrast, the affinity-matured compounds displayed ~ 40 -fold higher dissociation constants towards GCPIII (compound **9** $K_{D,GCPIII} 38 \pm 6$ nM and compound **10** $K_{D,GCPIII} 37 \pm 5$ nM) as compared to the original HIT₃ (DUPA, $K_{D,GCPIII} 1.2 \pm 0.3$ nM) [Fig. 4E], possibly as a result of a steric clash in the GCPIII binding pocket [Fig. 4F]. Flow cytometry

experiments on HT1080.hPSMA and LNCaP tumor cells were performed to assess whether compounds were able to recognize PSMA in a native cellular environment. Both compounds **11** and **12** avidly bound to PSMA, as shown by the cellular staining presented in Fig. 4G. ASM-matured compound **11** showed a more pronounced shift in flow cytometry on both HT1080.hPSMA and LNCaP cells, compared to compound **12** [Fig. 4G].

The internalization rate of compound **11** in LNCaP and PSMA-positive HT1080 cancer cells was assessed by confocal microscopy analysis. PSMA-617-FITC was used in the same experiment as a positive control. The results of this experiment indicate that both compounds internalize at a similar rate in both PSMA-positive cancer cell lines, with progressive internalization at 1, 6, and 24 hours after incubation [Fig. 5A and ESI S123†]. No staining was detected in PSMA-negative cancer cells (HT1080.wt).

Finally, we performed quantitative biodistribution studies with radiolabelled compounds to assess whether the favourable PSMA binding properties were associated with a preferential enrichment of the molecules at the tumor site. A head-to-head comparison of A70-B104-HIT₃ with Pluvicto™ was performed, as the latter product is routinely used in the clinic [Fig. 5C]. The corresponding DOTA conjugates were labeled with ¹⁷⁷Lu (compound **18** and Pluvicto™) and were intravenously injected into BALB/c nude mice ($n = 3$ per group) bearing HT1080.hPSMA tumors (single dose). Mice were sacrificed two hours after compound administration and the quantitative biodistribution of both molecules was evaluated by counting the radioactivity associated with their lutetium-177 payload.



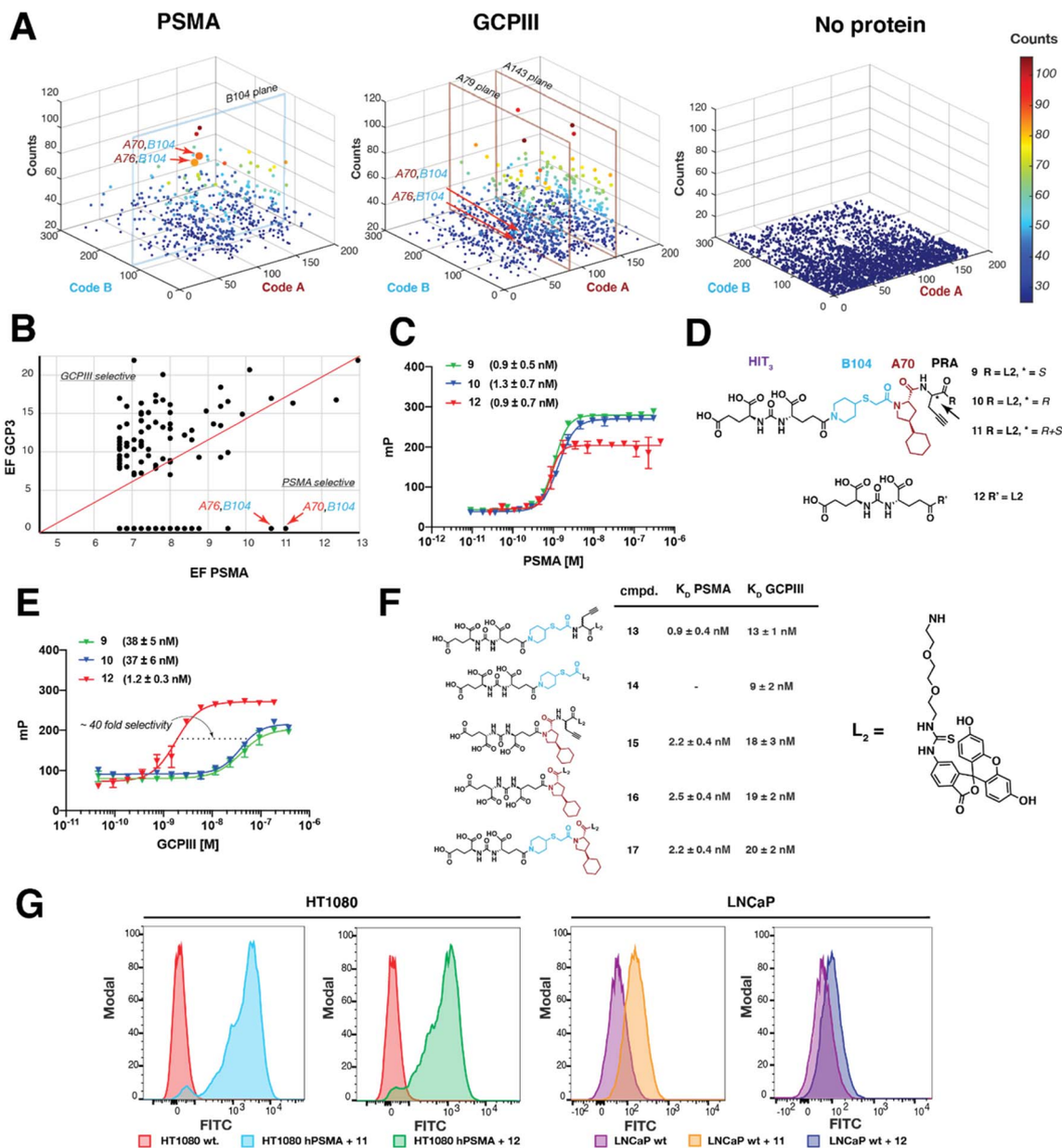


Fig. 4 (A) PRA-HIT₃ fingerprints against PSMA, GCPIII and underivatized beads (no protein) used for the selection process. (B) PSMA and GCPIII enrichment factors of the most enriched compound in PSMA screening. (C) Off-DNA FP of the diastereoisomers of the affinity matured molecules **9** and **10**, $K_D = 0.9 \pm 0.5$ nM and $K_D = 1.3 \pm 0.7$ nM, respectively, and the fluoresceinated HIT₃ **12**, $K_D = 0.9 \pm 0.3$ nM against PSMA. (D) Structure of the matured molecule PRA-A70-B104-HIT₃ and HIT₃ with their respective linker. (E) Off-DNA FP of the diastereoisomers of the affinity matured molecules **9** and **10**, $K_D = 38 \pm 6$ nM and $K_D = 37 \pm 5$ nM, respectively, and the fluoresceinated HIT₃ **12**, $K_D = 1.2 \pm 0.3$ nM against GCPIII. (F) Table with affinity against PSMA and GCPIII measured by Off-DNA FP of the fragments of the affinity matured compound. (G) Flow cytometry analysis of HT1080. hPSMA and LNCaP incubated with **11** and **12**. HT1080.wt was used as a reference.

Discussion

Pluvicto™ is the most recent RLT approved for the treatment of patients affected by metastatic castration-resistant prostate cancer. The ultra-high affinity of the drug to its cognate target antigen (prostate specific membrane antigen, expressed on the surface of prostate cancer cells) enables the efficient and selective targeted delivery of radionuclide payloads to tumors. Despite the clinical benefit demonstrated in the Phase III

VISION trial, 12% of the patients discontinued the treatment with the drug due to adverse events (*e.g.*, xerostomia)⁴⁴ which additionally limited the administration of Pluvicto™ at higher doses. Some of the undesired side effects occur due to the expression of GCPIII in salivary glands and kidneys, which is a highly homologous PSMA protein that causes an off-targeted accumulation of the PSMA-targeted RLT in healthy tissue.³¹

In this article, we have described the affinity and selectivity maturation of nanomolar and sub-nanomolar PSMA ligands, by



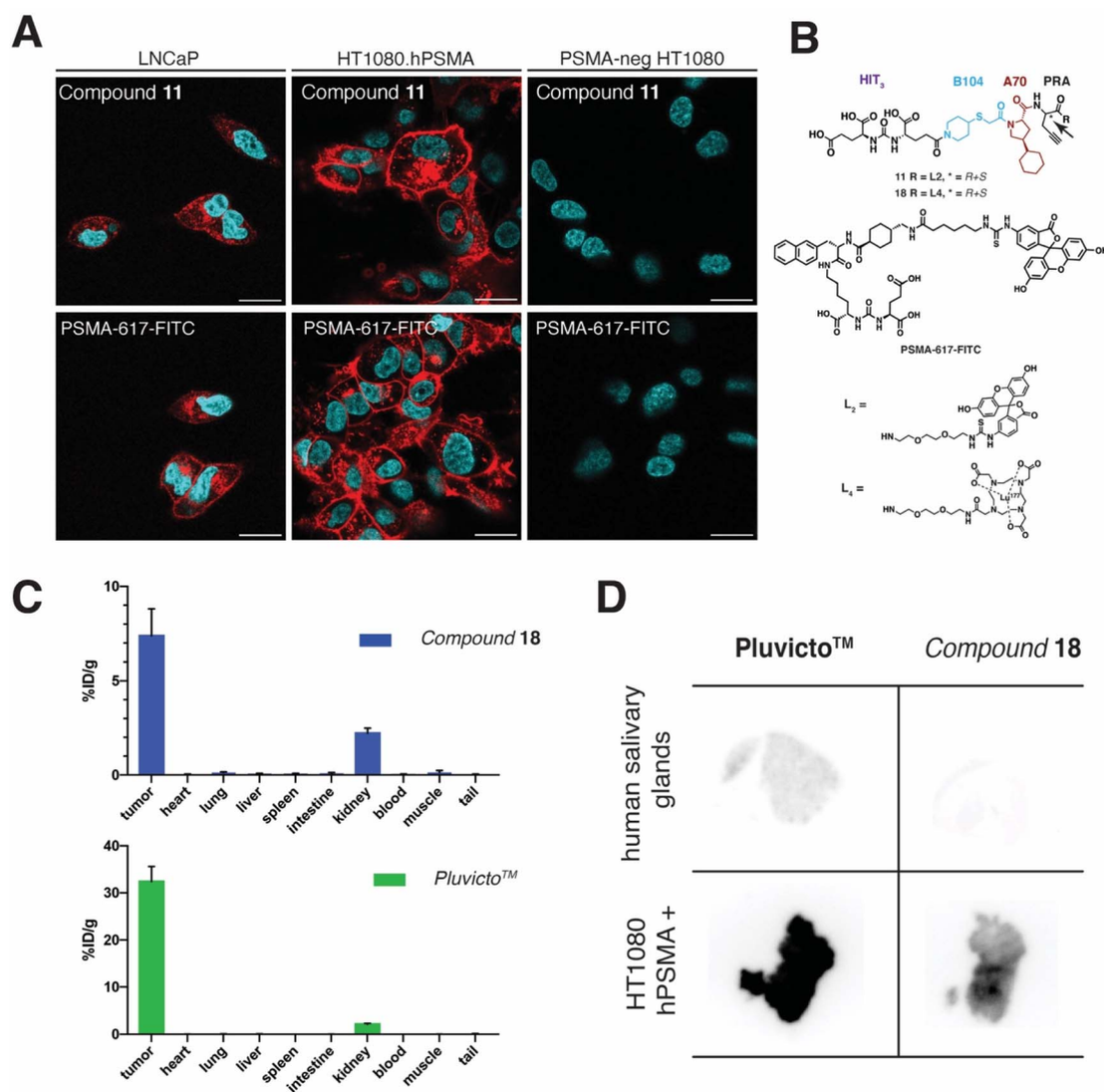


Fig. 5 (A) Confocal microscopy images after 24 h incubation of compound **11** and PSMA-617-FITC with PSMA-positive cells (LNCaP and HT1080.hPSMA). No interaction with PSMA-negative HT1080 (HT1080.wt) cells was observed. Red = fluorescein derivative staining; blue = Hoechst 33 342 staining. Scale bar, 20 μm. (B) Structure of compound **11**, PSMA-617-FITC and DOTA conjugate of A70-B104 (compound **18**). (C) Biodistribution experiment on HT1080.hPSMA tumor-bearing BALB/c nude mice with **18**, and Pluvicto™ with a dose of 1 nmol per mouse, 1 MBq for **18** and 0.5 nmol per mouse, 1 MBq for Pluvicto™. (D) Results of the autoradiography experiment on human salivary glands and HT1080.hPSMA tumor slices with compound **18** and Pluvicto™.

using DNA-encoded chemical library technology, to improve the affinity to PSMA, while enhancing ligand selectivity for the tumor-associated antigen over GCPIII. Therefore, the occurrence of selective and potent PSMA binders may prevent unwanted off-target accumulation in healthy organs, thus improving the RLT therapeutic window. We synthesized affinity and selectivity DNA-encoded chemical libraries (ASM-DELS) comprising 18'284'658 individually encoded compounds, obtained by derivatization of six different DELs with three elected HITS. Screening against PSMA and GCPIII resulted in the identification of several novel building block combinations with the three HITS. Compound **11** (A70-B104-HIT₃) was identified from the PRA-DEL as the most potent and selective PSMA ligand ($K_{D,PSMA} = 900$ pM; $K_{D,GCPIII} = 38$ nM). The compound was found to bind to and internalize into PSMA-positive cancer cells

by flow cytometry and confocal microscopy experiments. Ligand internalization of PSMA-targeted RLTs may influence their *in vivo* tumor uptake and tumor residence time, thus having an impact on their anti-cancer efficacy.

Moreover, *in vivo* biodistribution experiments in tumor-bearing mice, conducted with a lutetium-177 radioactive derivative of compound **11**, showed successful and selective accumulation in PSMA-expressing tumors (tumor uptake = ~7.4% ID g⁻¹, 2 hours after systemic administration). Furthermore, *ex vivo* autoradiography analysis provided an estimation of the ability of the ligand to selectively target the cancer lesion against human salivary glands, in contrast to Pluvicto™, which exhibited visible staining for both tissues.

The clinical maximum tolerated dose of Pluvicto™ has not yet been formally identified. The dose regimen approved for



treating metastatic prostate cancer patients (*i.e.*, 7.4 GBq per dose, for up to 6 administrations) has been established by the early clinical use of the drug under compassionate treatment and similar regulatory frameworks in Germany and Australia. However, adverse events including xerostomia are observed in a good proportion of patients treated at the approved Pluvicto™ dose, thus leading in ~12% of cases to discontinuation of the treatment.⁴⁴ Although in the past, medical strategies have been proposed to somewhat alleviate these undesired effects (*e.g.*, cooling of the throat^{45,46}), the generation of PSMA ligands that do not interact with GCPIII in healthy organs could solve the unpleasant burden of salivary gland toxicity.⁴⁴ Beforehand, medicinal chemistry efforts have highlighted the benefit of introducing “rigid linkers” on DUPA to obtain PSMA-selective ligands.³⁴ Unfortunately, while the described derivatives were presented as highly potent and selective, their ability to target PSMA-positive tumors after *in vivo* administration was never formally demonstrated. We have utilized our high-throughput DEL platform to obtain potent and selective PSMA-targeting ligands with flexible and linear structures from a large collection of compounds. DEL technology has been previously applied as an effective tool to mature the affinity of small organic ligands for tumor-targeting purposes. Our group has recently published the implementation of DEL strategies to mature the affinity of nanomolar ligands against carbonic anhydrase IX and fibroblast activation protein to obtain ultra-potent binders with affinities in the low picomolar range and with an improved tumor-residence time.^{35,37} In this new article, we have shown how DELs can be instrumental in enhancing in parallel the affinity to a target protein and the selectivity against an anti-target protein of available HIT compounds. Selectivity is a fundamental property of targeted therapies, with a great impact on minimizing healthy organ uptake and preventing systemic side effects. The PSMA-selective compounds resulting from this DEL selection campaign are “portable” moieties that can be alternatively coupled to DNA-barcodes, fluorophores, or radionuclide chelators without losing their affinity and selectivity for the cognate target.

One of the limitations of this study is the lack of an *in vivo* preclinical model that allows the assessment of the selectivity of DEL-derived tumor-targeting ligands. Tumor-bearing mice are routinely used to assess the tumor-targeting properties of antibodies and small organic ligands. However, in the case of PSMA, unwanted accumulation in salivary glands cannot be predicted in rodents or minipigs^{23,24,47} (ESI S13.3, Fig. S127†) as, in those species, GCPIII is not expressed in salivary glands. Autoradiography experiments on human salivary glands and PSMA-positive tumors are presented here as a method of choice to quantify the selectivity of novel PSMA ligands and to benchmark them against Pluvicto™ (Fig. 5D).

Autoradiography studies show no binding of compound **18** on human salivary glands. Our work relies on the assumption that salivary gland uptake of urea-based PSMA ligands is mainly caused by GCPIII expression in the tissue. According to alternative studies, salivary gland accumulation appears to be non-PSMA-related⁴⁸ or partly non-specific.⁴⁹ It is evident that the underlying mechanism is still under investigation, and

additional work is necessary to advance in the field of PSMA radioligand therapeutics.

Given its high potency, preferential binding to PSMA compared to GCPIII, internalization properties and excellent *in vivo* tumor-targeting performance, compound A70-B104-HIT₃ is a promising HIT compound for the generation of imaging and therapeutic constructs based on radionuclides (*e.g.*, gallium-68 and lutetium-177) and cytotoxic drugs (*e.g.*, monomethyl auristatin E). The anti-cancer efficacy of radioligand therapeutics is directly determined by their tumor uptake and residence time in target neoplastic masses. These two features are strongly influenced by the ligand affinity for the cognate target and by the compounds' physicochemical properties. Medicinal chemistry optimization of A70-B104-HIT₃ tumor uptake is needed to increase the tumor-to-kidney ratio and improve the drug's therapeutic window. Nuclear medicine studies that demonstrate the ability of the compound to target tumors in prostate cancer patients with PSMA-positive lesions while sparing healthy organs are still needed to warrant its clinical application.

Conclusion

Salivary gland toxicity remains a critical problem that limits dose escalation of PSMA-targeted RLTs in patients affected by prostate cancer. At the same time, the high similarity between PSMA and GCPIII, expressed in salivary glands, represents a molecular challenge for conventional medicinal chemistry. As demonstrated in this work, the use of DEL technology can be of great utility for the maturation of binders to generate new potent and selective PSMA ligands. Affinity and selectivity matured compound A70-B104-HIT₃ is a portable PSMA-selective compound that can be used for the generation of theranostic small molecules resulting in therapeutic agents that are better tolerated and more efficacious in patients with antigen-positive prostate cancer lesions.

Data availability

The datasets supporting this article have been uploaded as part of the ESI.†

Author contributions

Laura Lucaroni: writing – review & editing, writing – original draft, visualization, validation, methodology, investigation, formal analysis, data curation, conceptualization. Sebastian Oehler: investigation, validation, methodology. Tony Georgiev: investigation. Marco Müller: investigation, validation, methodology. Matilde Bocci: investigation. Roberto de Luca: investigation. Nicholas Favalli: investigation. Dario Neri: writing – review & editing, writing – original draft, visualization, validation, supervision, resources, project administration, methodology, funding acquisition, formal analysis, data curation, conceptualization. Samuele Cazzamalli: writing – review & editing, writing – original draft, visualization, validation, supervision, resources, project administration, methodology,



investigation, funding acquisition, formal analysis, data curation, conceptualization. Luca Prati: writing – review & editing, writing – original draft, visualization, validation, supervision, project administration, methodology, investigation, formal analysis, data curation, conceptualization.

Conflicts of interest

D. N. is co-founder, CEO, CSO and President of the Scientific Advisory Board of Philogen. L. L., S. O., T. G., M. M., R. D. L., N. F. and S. C. are employed by Philochem AG, and the research and development unit of Philogen. L. P. is employed by Philogen S.p.A.

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