Advancing the management of adult solid tumours in 2023 and beyond:
Unlocking the potential of radiopharmaceuticals



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# Addressing diagnostic and treatment challenges with radiopharmaceuticals: Learnings from theranostic approaches in GEP-NETs

### **Dr Jason Starr**

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How have radiopharmaceuticals impacted the diagnosis and treatment of GEP-NETs in recent years?

### FDA-approved radiopharmaceuticals in GEP-NETs

<sup>68</sup>Ga-DOTATATE 64Cu-DOTATATE <sup>68</sup>Ga-DOTATOC



177Lu-DOTATATE as second-line therapy for SSTR-positive **GEP-NETs** 



<sup>68</sup>Ga for PET imaging (diagnostic)/177Lu as **B**-emitter (therapeutic)





### Advantages

- High disease control4
- Overall limited toxicity<sup>4</sup>



#### Limitations

- Possible long-term side effects Nephrotoxicity, 4 haematotoxicity, 4,5 possible hepatotoxicity<sup>6</sup>
- Defining eligible patients<sup>7</sup> Tolerability to PRRT depends on patient's SSTR avidity, tumour burden, organ function and the patient's functional status

FDA, US Food and Drug Administration; GEP-NET, gastroenteropancreatic neuroendocrine tumour;

PET, positron emission tomography; PRRT, peptide receptor radionuclide therapy; SSTR, somatostatin receptor.

- 1. NCCN. Neuroendocrine and adrenal tumors. 2022. Available at: https://bit.ly/3OGRDtk (accessed 17 May 2023);
- 2. Brugarolas P, et al. J Nucl Med Technol. 2020;48(Suppl. 1): 34S-9S; 3. Barca C, et al. Pharmaceuticals (Basel). 2021;15:13; 4. Telo S, et al. Clin Transl Imaging. 2021;9:423-38;
- 5. Bergsma H, et al. J Nucl Med. 2018;59:452-8; 6. Riff BP, et al. Clin Nucl Med. 2015;40:845-50; 7. Burkett BJ, et al. Radiology. 2021;298:261-74.



What are the key efficacy outcomes for radiopharmaceuticals in the treatment of GEP-NETs?

### Pivotal clinical trials: Efficacy outcomes



#### NETTER-1<sup>1</sup>

SSTR positive advanced midgut NETs (N=229)



Phase III trial evaluating <sup>177</sup>Lu-DOTATATE 7·4 GBq (200 mCi) every 8 weeks (four cycles) + long-acting octreotide 30 mg vs long-acting octreotide 60 mg every 4 weeks

**PFS: 28.4** months vs **8.5** months (HR 0.21, 95% CI 0·14–0.33; p<0.0001)<sup>2</sup>

**mOS: 48.0** months vs **36.3** months (HR 0.84, 95% CI 0.60–1.17; two-sided p=0.30)<sup>3</sup>



Clinically and statistically significant improvement in PFS and clinically relevant longer mOS of 11.7 months with <sup>177</sup>Lu-DOTATATE<sup>4</sup>



#### **NETTER-R**<sup>5</sup>

Pancreatic NETs (N=62 assessed by RECIST v1.1)



Retrospective real-world data (multiple sites) from patients treated with  $^{177}$ Lu-DOTATATE 7.4 GBq at 8  $\pm$  1-week intervals; median follow-up after first cycle: 24.5 months (range 20–123.4 months)

median PFS: 24.8 months (95% CI 17.5–34.5) median TTP: 29.5 months (95% CI 21.4–67.6)

ORR: 40.3% (25/62; 95% CI 28.1-53.6)

median DoR: 60.7 months (95% CI 13.1–62.1)

**mOS (n=110): 41.4** months (95% CI 28.6–50.2)



Study reinforces the role of <sup>177</sup>Lu-DOTATATE for treatment of patients with SSTR-positive pancreatic NETs

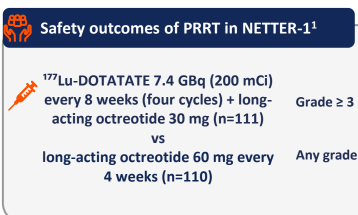
CI, confidence interval; DoR, duration of response; HR, hazard ratio; mOS, median overall survival; NET, neuroendocrine tumour; ORR, objective response rate; PFS, progression-free survival; RECIST v1.1, Response Evaluation Criteria in Solid Tumors version 1.1; SSTR, somatostatin receptor; TTP, time to progression.

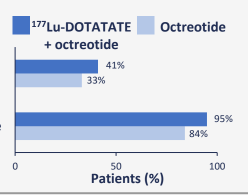


1. Strosberg J, et al. N Engl J Med. 2017;376:125–35; 2. Smith-Palmer J, et al. BMC Cancer. 2021;21:10; 3. Strosberg JR, et al. Lancet Oncol. 2021;22:1752–63; 4. Strosberg JR, et al. J Clin Oncol. 2021;39(Suppl.):4112; 5. Clement D, et al. Eur J Nucl Med Mol Imaging. 2022;49:3529–37.

What are the key safety considerations for radiopharmaceuticals in patients with GEP-NETs?

# Safety of PRRT vs targeted therapy in GEP-NETs





### <sup>177</sup>Lu-DOTATATE grade ≥3 AEs<sup>2</sup>

#### **Incidence ≥4%\***

- Lymphopenia
- ◆ ↑ AST

◆ ↑ GGT

- ALT
- Vomiting
- HyperglycaemiaHypokalaemia
- Nausea
  - **Long-term haematologic AEs**
- t-MN, mean (SD): 2.61% (4.38%)<sup>3</sup>
- Persistent haematologic dysfunction: 4%<sup>4</sup>



### Safety profile of everolimus in various clinical trials<sup>†</sup>

- Incidence ≥30%<sup>5</sup>
- Safety and tolerability was consistent in all studies in advanced NET settings (RADIANT-2, RADIANT-3 and RADIANT-4)<sup>6-8</sup>
- Frequently observed AEs were grade 1 or 2 including stomatitis, diarrhoea, fatigue, infections, rash and peripheral oedema<sup>8</sup>

AE, adverse event; ALT, alanine aminotrasferase; AST, aspartate aminotransferase; FDA, US Food and Drug Administration; GEP-NET, gastroenteropancreatic NET; GGT, gamma-glutamyl transferase; NET, neuroendocrine tumour; PRRT, peptide receptor radionuclide therapy; SD, standard deviation; t-MN, therapy-related myeloid neoplasm. 1. Strosberg J, et al. N Engl J Med. 2017;376:125–35; 2. FDA. Lutetium Lu 177 dotatate PI. Available at: <a href="https://bit.ly/3IAoNqA">https://bit.ly/3IAoNqA</a> (accessed 23 May 2023); 3. Sonbol MB, et al. J AMA Oncol. 2020;6:1086–92; 4. Bergsma H, et al. J Nucl Med. 2018;59:452–8; 5. FDA. Everolimus PI. Available at: <a href="https://bit.ly/3OKkw7E">https://bit.ly/3OKkw7E</a> (accessed 24 May 2023); 6. Pavel ME, et al. Ann Oncol. 2017;28:1569–75; 7. Yao JC. et al. New Engl J Med. 2011;364:514–23; 8. Yao JC. et al. Lancet. 2016;387:968–77.



<sup>\*</sup>With a higher incidence in <sup>177</sup>Lu-Dotatate arm; <sup>†</sup>across various tumour types.

How can clinicians best integrate radiopharmaceuticals into the clinical setting to ensure optimal outcomes for patients with GEP-NETs?

# Implementing PRRT in clinical settings

Systematic checklist to be used by the MTB for patient evaluation<sup>1–3</sup>



#### MTB<sup>1-3</sup>

- Medical/surgical oncologists
- Radiation oncologists
- Nuclear medicine specialists/radiologists
- Gastroenterologists



### Therapy appropriateness<sup>1</sup>

- Histopathologic findings
  - ✓ Proven NET
  - ✓ WHO classification
- Diagnostic imaging to confirm high somatostatin receptor expression
  - ✓ Nuclear imaging for patient selection and estimating treatment response<sup>4</sup>
  - ✓ CT/MRI



#### Patient safety assessment to tolerate therapy<sup>1</sup>

- Assess adequate organ function for PRRT
  - ✓ Renal Nephrology
  - ✓ Hepatic 

    → Hepatology
  - ✓ Bone marrow 

     Haematology
- Assess tumour burden Cardiology, endocrinology, gastroenterology, pathology (NET specialist)
- Review recent/concurrent treatments
- Assess patient factors
  - ✓ BMI
  - ✓ Karnofsky or ECOG performance status score
  - Can follow radiation safety precautions
  - × Pregnancy/breastfeeding

BMI, body mass index; CT, computerized tomography;

ECOG, Eastern Cooperative Oncology Group; MRI, magnetic resonance imaging; MTB, Multidisciplinary tumour board; NET, neuroendocrine tumour; PRRT, peptide receptor radionuclide therapy; WHO, World Health Organization. 1. Burkett BJ, et al. *Radiology*. 2021;298:261–74; 2. Mejia A, et al. *Medicine (Baltimore)*. 2022;101:(e28970).261–74; 3. Hendifar AE, et al. *Pancreas*. 2022;51:213–8; 4. Puliani G, et al. *Front Endocrinol (Lausanne)*. 2022;13:861434.



How might clinical trials help address the key remaining questions about the use of radiopharmaceuticals in **GEP-NETs?** 

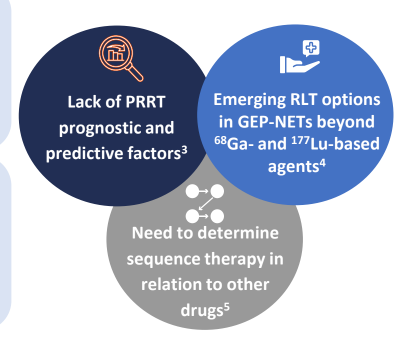
# Using the latest data to inform practice

Genetic profiling analysis design of well-differentiated aggressive grade 2 and 3 GEP-NETs in the phase III COMPOSE trial<sup>1</sup>

 Analysis may guide detection of pathogenic mutations in NET patients to inform treatment and surveillance strategies

Prognostic value of TTV with <sup>68</sup>Ga-DOTATOC PET/CT in predicting response to <sup>177</sup>Lu-DOTATOC treatment in metastatic well-differentiated NETs<sup>2</sup>

 TTV could be considered as an easily accessible and widely available prognostic imaging biomarker



Prospective evaluation of the utility of concurrent <sup>18</sup>F-FDG PET/CT and <sup>68</sup>Ga-DOTATOC imaging in GEP-NENs: The PETNET study<sup>6</sup>

A positive FDG PET was significantly associated with reduced OS

ACTION-1 phase Ib/III trial of RYZ101 in SSTR2+ GEP-NETs progressing after <sup>177</sup>Lu SSA therapy: Initial safety analysis<sup>7</sup>

- RYZ101 was well tolerated
- 120 kBq/kg declared as RP3D
- Part 2 (phase III) will compare RYZ101 with SOC in pre-treated patients with SSTR2+ GEP-NETs

CT, computerized tomography; GEPNEN, gastroenteropancreatic neuroendocrine neoplasm; GEP, gastroenteropancreatic; NEN, neuroendocrine neoplasm; NET, neuroendocrine tumour; OS, overall survival; PET, positron emission tomography; PRRT, peptide receptor radionuclide therapy; RLT, radioligand therapy; RP3D, recommended phase II dose; SOC, standard of care; SSA, somatostatin analogue; SSTR2, somatostatin receptor subtype 2; TTV, total tumour volume.

- 1. Halfdanarson TR, et al. J Clin Oncol. 2023;41(Suppl.): TPS660; 2. Vega-Zolano E, et al. J Clin Oncol. 2023;41(Suppl.):e16248;
- 3. Puliani G, et al. Front Endocrinol (Lausanne). 2022;13:861434; 4. Becx MN, et al. Cancers (Basel). 2022;14:5792; 5. Albertelli M, et al. Rev Endocr Metab Disord. 2021;22:563–79; 6. Vasconcelos JPS, et al. J Clin Oncol. 2023;41(Suppl.):4022; 7. Morris M, et al. J Clin Oncol. 2023;41(Suppl.):4132.



# Advancing outcomes in prostate cancer: Current and future perspectives on theranostics

### Prof. Jorge A Garcia

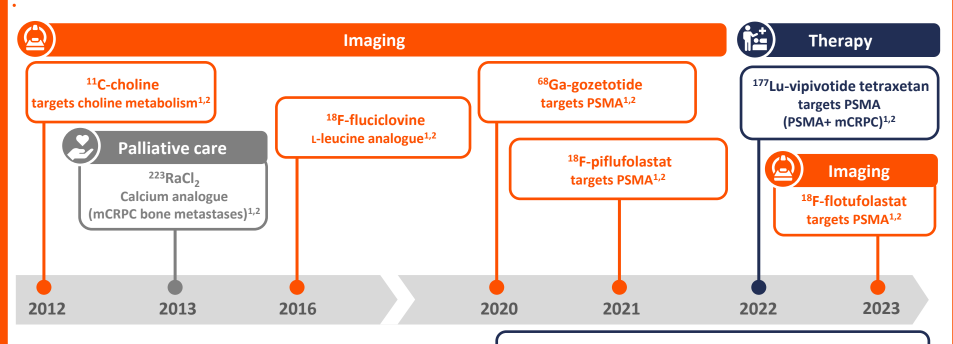
University Hospitals Seidman Cancer Center Case Western Reserve University Cleveland, OH, USA





What is the current status and role of radiopharmaceuticals in the management of prostate cancer?

### FDA-approved radiopharmaceuticals in prostate cancer



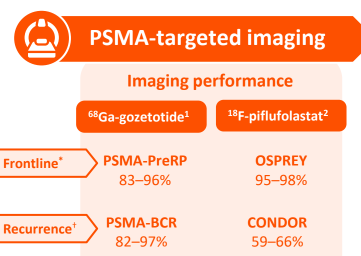
Development of PSMA-targeting radiopharmaceuticals heralds a new era of high-precision theranostics<sup>1</sup>



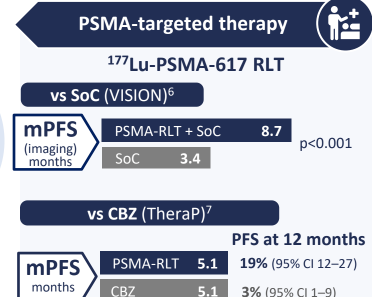
<sup>1.</sup> Jia AY, et al. *Prostate Cancer Prostatic Dis.* 2023;doi:10.1038/s41391-023-00670-6; 2. FDA. Prescribing information searchable by agent. Available at: <a href="https://www.accessdata.fda.gov/scripts/cder/daf/">www.accessdata.fda.gov/scripts/cder/daf/</a> (accessed 19 May 2023).

What have we learned about the value of PSMA-based theranostics in prostate cancer from pivotal trial data?

# Value of radiotheranostics in mCRPC management







- Superior diagnostic accuracy vs conventional imaging (initial staging and recurrent disease)<sup>3,4</sup>
- May guide <sup>177</sup>Lu-PSMA-617 treatment eligibility<sup>4</sup>
- Improved sensitivity at >PSA relative to <PSA<sup>3,4</sup>
- <sup>18</sup>F-PSMA-11 non-inferior vs <sup>68</sup>Ga-PSMA-11<sup>5</sup>

in heavily pre-treated patients with PSMA+ disease<sup>4</sup>

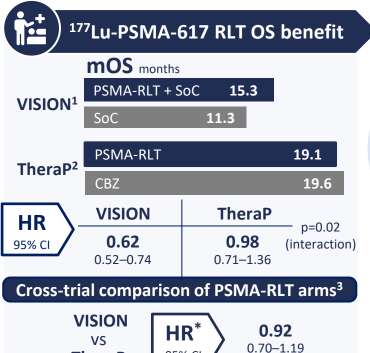
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www.accessdata.fda.gov/drugsatfda\_docs/label/2021/214793s000lbl.pdf (accessed 20 June); 3. Keegan NM, et al. Eur Urol Focus. 2021;7:267–78; 4. Jia AY, et al. Prostate Cancer Prostatic Dis. 2023;doi:10.1038/s41391-023-00670-6; 5. De Man, K et al. Eur Urol. 2022;82:501–9; 6. Sartor O, et al. N Engl J Med. 2021;385:1091–103; 7. Hofman MS, et al. Lancet. 2021;397:797–804.

<sup>\*</sup>Specificity; ¹True positive in ≥1 region(s). CBZ, cabazitaxel; CI, confidence interval; m, median; mCRPC, metastatic castration-resistant prostate cancer; PFS, progression-free survival; PSA, prostate-specific antigen; PSMA, prostate-specific membrane antigen; RLT, radioligand therapy; SoC, standard of care.

1. FDA. <sup>68</sup>Ga-gozetotide PI. Available at: www.accessdata.fda.gov/drugsatfda docs/label/2022/212642s002lbl.pdf (accessed 20 June 2023); 2. FDA. <sup>18</sup>F-piflufolastat PI. Available at:

### Refining understanding with additional analyses



95% C

**TheraP** 



### **PSMA-PET predictive value**



3 /11	110	B 1/I
VIS	SIO)	M4
V 15	$\sim$	

Whole-body SUV<sub>mean</sub>

	<b>Highest quart</b>	tile L	owest quartile
rPFS months	14.1	VS	5.8
mOS months	21.4	VS	14.5

#### TheraP<sup>5</sup>

**PSA response (PSMA-RLT vs CBZ)** 



**SUV**<sub>mean</sub> **≥10 12.19** 3.42–58.76

**SUV**<sub>mean</sub> <**10 2.22** 1.11-4.51 p=0.039<sup>1</sup>



<sup>\*</sup>Unadjusted HR,3 †adjusted p-value (p<sub>adj</sub>) for treatment-by-SUV<sub>mean</sub> interaction.5 CBZ, cabazitaxel; CI, confidence interval; HR, hazard ratio; m, median; OS, overall survival; OR, odds ratio; PET, positron emission tomography; PSA, prostate-specific antigen; PSMA, prostate-specific membrane antigen; RLT, radioligand therapy; rPFS, median radiographic progression-free survival; SoC, standard of care; SUV, standardized uptake value. 1. Sartor O, et al. *N Engl J Med*. 2021;385:1091–103; 2. Hofman MS, et al. *J Clin Oncol*. 2022;40(Suppl. 16):5000; 3. Soon YY, et al. *J Clin Oncol*. 2023;41(Suppl. 16):5045; 4. Kuo P, et al. *J Clin Oncol*. 2022;40(Suppl. 16):5002: 5. Buteau P, et al. *Lancet Oncol*. 2022;23:1389–97.

What are the key safety considerations when integrating 177Lu-PSMA-RLT into the management of patients with prostate cancer?

### Safety considerations with <sup>177</sup>Lu-PSMA-RLT



### Dosimetry

 Informed by <sup>177</sup>Lu-DOTATATE safety data and EBRT absorbed dose constraints on bone marrow and kidney<sup>1,2</sup>

### **Key differences EBRT vs RLT<sup>1</sup>**

- Prescribing protocols
- Treatment schedules
- Dose rates
- Tissue uptake



Frequent adverse reactions (≥20%)<sup>3,4</sup>



Common laboratory abnormalities (≥30%)<sup>4</sup>

- Fatigue
- Dry mouth
- Nausea
- Anaemia
- ↓ Appetite
- Constipation
- Arthralgia
- Back pain

- Lymphopenia
- Leukopenia
- Thrombocytopenia
- ↓ Calcium
- ↓ Haemoglobin
- **J** Sodium
- Consider long-term toxicity in risk/benefit assessments for radiolabelled agents<sup>5</sup>
- Prevention is key as some end-organ toxicities may be irreversible<sup>5</sup>
- Consider baseline patient characteristics e.g., pre-treatment haemoglobin,<sup>6</sup> cytopenias<sup>7</sup>

EBRT, external beam radiation therapy; PSMA, prostate-specific membrane antigen; RLT, radioligand therapy; TAT, targeted alpha therapy.

- 1. Jia AY, et al. Prostate Cancer Prostatic Dis. 2023;doi:10.1038/s41391-023-00670-6; 2. Hofman MS, et al. Lancet Oncol. 2018;19:825–33;
- 3. Sartor O, et al. N Engl J Med. 2021;385:1091–103; 4. FDA. 177 Lu vipivotide tetraxetan PI. Searchable at: <a href="https://www.accessdata.fda.gov/scripts/cder/daf/">www.accessdata.fda.gov/scripts/cder/daf/</a> (accessed 23 May 2023);
- 5. FDA. 2020. Available at: <a href="https://www.fda.gov/media/144843/download">www.fda.gov/media/144843/download</a> (accessed 23 May 2023); 6. Nelson AA, et al. J Clin Oncol. 2023;41(Suppl. 16):e17065;
- 7. Abdelrazek AS, et al. J Clin Oncol. 2023;41(Suppl. 16):e17057.



What key clinical questions remain, and how are trials aiming to address these?

### ASCO 2023 insights: New approaches in mCRPC

#### **New combinations**



LuPARP<sup>1</sup> (NCT03874884)

#### Phase I: <sup>177</sup>Lu-PSMA-617 plus PARPi (olaparib)

9-cohort dose escalation study





- <sup>177</sup>Lu-PSMA-617</sup> 7.4 GBq 6 weekly, 6 cycles
- Olaparib 50–300 mg BD (3+3 escalation)\*



- PSA50\* 66% (n=21/32)
- PSA90\* 44% (n=14/32)
- ORR 78% (n=7/9)



- No DLTs across doses
- No grade 4 TRAEs

Anaemia (7%)
Thrombocytopenia (3%)
Neutropenia (7%)

RP2D: 7.4 GBq <sup>177</sup>Lu-PSMA-617 plus 300 mg BD olaparib days -4–18 of each 6-weekly cycle

**New agents** 



 $TAT + RLT^2$  (NCT04886986)

Phase I: <sup>225</sup>Ac-J591 plus <sup>177</sup>Lu-PSMA-I&T (PNT2002)

Dose escalation study





- <sup>225</sup>Ac-J**591** 30, 35 or 40 KBq/kg
- 177Lu-PNT2002 6.8 GBq



94% experienced

- PSA50<sup>†</sup> 61% (n=11/18)
- Day 8 SPECT/CT confirmed accurate tumour targeting



Dual PSMA targeting is feasible and tolerable:

- DLTs at 40 KBq/kg only
- No grade 4 TRAEs

Anaemia (17%)
Thrombocytopenia (11%)
Pain (5%)

RP2D: 35 KBq/kg <sup>225</sup>Ac-J591 plus 6.8 GBq <sup>177</sup>Lu-PNT2002

\*Day: 2–15 or 4–12 or 4–18; \*PSA response defined as the proportion of patients achieving either a reduction of 50% (PSA50) or 90% (PSA90) from baseline.

ASCO, American Society of Clinical Oncology; BD, twice daily; Bq, Becquerel; CT, computerized tomography; DLT, dose-limiting toxicity; G, giga; k, kilo; mCRPC, metastatic castration resistant prostate cancer; ORR, overal response rate; PARPi, poly (ADP-ribose) polymerase inhibitor; PSA, prostate-specific antigen; PSMA, prostate-specific membrane antigen; RLT, radioligand therapy; RP2D, recommended phase 2 dose; SPECT, single-photon emission computed tomography; TAT, targeted alpha therapy; TRAE, treatment-related adverse event.

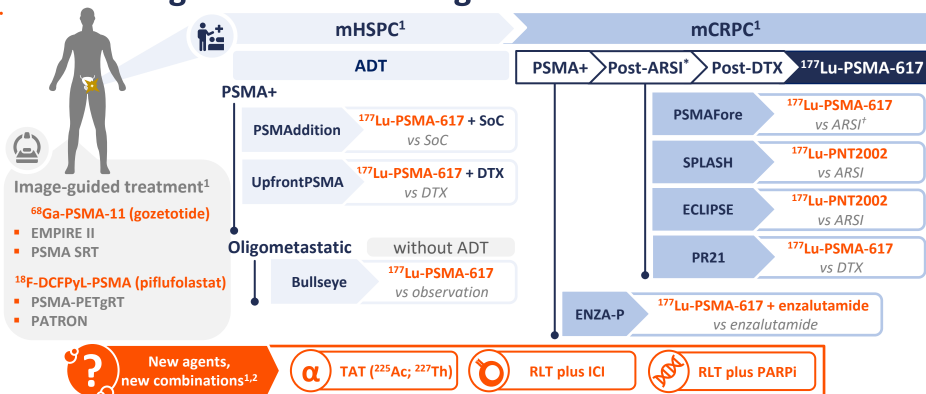
1. Sandhu S, et al. J Clin Oncol. 2023;41(Suppl. 16):5005; 2. Tagawa S, et al. J Clin Oncol. 2023;41(Suppl. 16):5018.

G3



How might theranostics impact the future management of prostate cancer, now and in the future?

### **Evolving role of PSMA-targeted radiotheranostics**



<sup>\*</sup>Progression on prior ARSI: †ARSI not previously used.

1. Jia AY, et al. Prostate Cancer Prostatic Dis. 2023;doi:10.1038/s41391-023-00670-6; 2. Jang A, et al. Ther Adv Med Oncol. 2023;15:1-12.



ADT, androgen deprivation therapy; ARSI, androgen-receptor signalling inhibitor; ChT, chemotherapy; DTX, docetaxel; ICI, immune checkpoint inhibitor; mCRPC, metastatic castration-resistant prostate cancer; mHSPC, metastatic hormone-sensitive prostate cancer; PARPi, poly (ADP-ribose) polymerase inhibitor; PSMA, prostate-specific membrane antigen; RLT, radioligand therapy; SoC, standard of care; TAT, targeted alpha therapy.

# Radiopharmaceuticals for adult solid tumours: Challenges and opportunities for implementation

### **Dr Erik Mittra**

Oregon Health & Science University Portland, OR, USA





- How can nuclear medicine
- specialists and oncologists work effectively together to successfully implement radiopharmaceuticals into
  - oncology practice?



# Preparation, communication and collaboration are key







# Defining roles and responsibilities

- Referring and treating physicians, and AUs administering radiopharmaceuticals, may differ
- Collaboration between oncologic workflows and theranostic centre

# Active presence and participation of AUs

- AUs (nuclear medicine specialists and radiation oncologists) are key for awareness, acceptance and consideration of radiopharmaceutical options
- Communication with clinicians managing cancer patients is essential

# Coordinating interdisciplinary HCP involvement

- MDT expertise is needed
   e.g. AUs, nurses, RSOs, medical
   physicists, radiochemists/pharmacists
- Co-ordinating patient follow-up and care beyond specialist centres

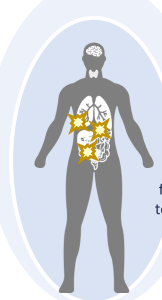


Effective collaboration within and across specialties throughout the patient journey is key to support integration of radiopharmaceuticals into oncology practice



What is the value of radiopharmaceutical theranostics in achieving high-precision medicine in oncology?

# Personalizing the care continuum with theranostics



Personalized management

from drug discovery to diagnosis, through to treatment and monitoring



### Refined patient selection<sup>1-3</sup>

- Actionable molecular target(s)
- Treatment eligibility evaluation



### Molecular imaging<sup>1-3</sup>

- Refined whole-body approach to management
- Prediction and prognostication

Theranostic 'pairs'



### Targeted therapy<sup>1-4</sup>

- Ongoing drug and radionuclide development
- Tailoring individualized dosimetry

Radiopharmaceuticals in nuclear medicine are leading the way in theranostics development, offering the potential for high-precision oncology management in adult solid tumours<sup>1–3</sup>





What are the barriers to the adoption of radiopharmaceuticals as a gold standard treatment in adult oncology?

# Identifying needs to support broader adoption



Many practitioners
would like to utilize
radiopharmaceuticals
more actively, but
barriers to wider
implementation remain<sup>1</sup>

56%\* of radiation oncologists surveyed in the US wanted to actively prescribe more RPT<sup>1</sup>



Referral pathways and MDT collaboration<sup>1,2</sup>

- Ill-defined referral pathways
- Not enough individuals for full MDT availability



Workforce and training<sup>1-3</sup>

- Lack of trained HCPs
- Need for RPT expertise and interdisciplinary collaboration



Treatment infrastructure<sup>1-3</sup>

 Variation in approaches between specialist vs community clinics



Logistics and supply chains<sup>1,2</sup>

 Limitations in availability and delivery of RPT to various institutions



Governance and regulation<sup>1,2</sup>

 Clarity needed on licensing requirements and clinical guidelines

HCP, healthcare professional; MDT, multidisciplinary team; RPT, radiopharmaceutical therapy.





<sup>\*</sup>n/N=74/131.

What strategies do you suggest to help overcome the barriers associated with adopting radiopharmaceuticals in oncology?

# Implementing broader adoption



Referral pathways and MDT collaboration<sup>1,2</sup>

- Minimum expected caseload
- Partner with referring oncologist/physician(s)
- 'Nuclear medicine champions'



Workforce and training<sup>1-3</sup>

- Steering committees and MDT implementation
- Nursing capacity and pharmacy support
- Nuclear medicine technologist input



Treatment infrastructure<sup>1-3</sup>

- Type of treatment and institution/centre size
- Appropriate clinical space(s) and protocols
- Outpatient vs inpatient management



Logistics and supply chains<sup>1,2</sup>

- Defined care implementation and coordination
- Radionuclide generation and handling
- Decontamination and waste disposal protocols



Governance and regulation<sup>1–3</sup>

- RAM licensing
- Safety requirements
- Designated RSOs and AUs



Patient support considerations<sup>1–3</sup>

- Educational materials for patients
- Counselling and support
- Transition from trial to real-world settings

and embedding within oncology workflows is needed1

**Effective integration** 

Touch™ ENDOCRINOLOGY Where do you see radiopharmaceuticals within oncology in the next 5 years?

# Radiopharmaceuticals: Towards a new standard of care?





generation to guide practice<sup>1,2</sup>

- Optimize dosimetry
- Innovate trial design
- Long-term follow-up data



Expanding options with new agents and/or indications<sup>2</sup>

- New targets and applications
- New combination regimens
- New radionuclides and constructs
- Use earlier in oncology pathways



Scaling of infrastructure and expertise to support delivery<sup>1,2</sup>

- Update training programs
- Networks of expertise
- Unlock development pipelines

Radiopharmaceuticals and theranostics offer scope for a new standard of personalized management in oncology<sup>1</sup>

