

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

Disclaimer

- *Unapproved products or unapproved uses of approved products may be discussed by the faculty; these situations may reflect the approval status in one or more jurisdictions*
- *The presenting faculty have been advised by touchIME and USF Health to ensure that they disclose any such references made to unlabelled or unapproved use*
- *No endorsement by touchIME or USF Health of any unapproved products or unapproved uses is either made or implied by mention of these products or uses in touchIME and USF Health activities*
- *touchIME and USF Health accept no responsibility for errors or omissions*

Radiopharmaceuticals in principle: Mechanism of action and biological effects

Dr Stephen A Graves

Division of Nuclear Medicine
Carver College of Medicine
University of Iowa, Iowa City, USA



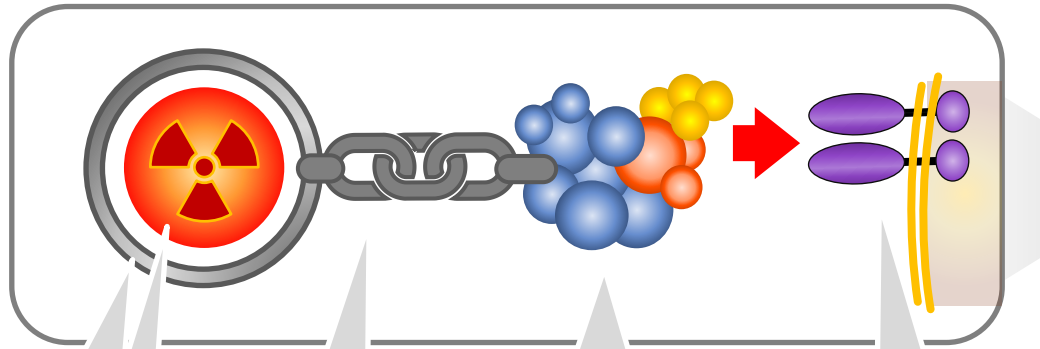


How do we design radiopharmaceuticals for clinical applications?

Design and construct of radiopharmaceuticals

Radionuclide, vector and target selection¹⁻⁴

Pharmacokinetics, decay profile and toxicity risks²⁻⁴



Chelator

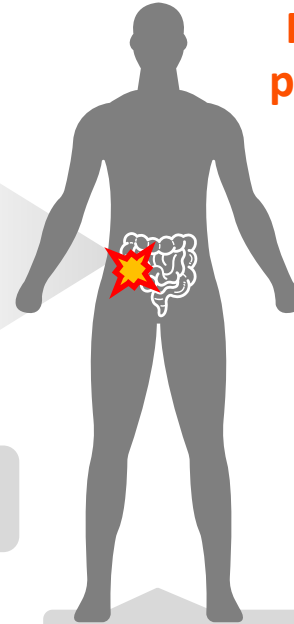
Linker /
spacer

Biomolecule /
vector

Tumour
cell target

Radionuclide

- Antibodies
- Peptides
- Small molecules
- Microspheres
- Nanoconstructs



Required application

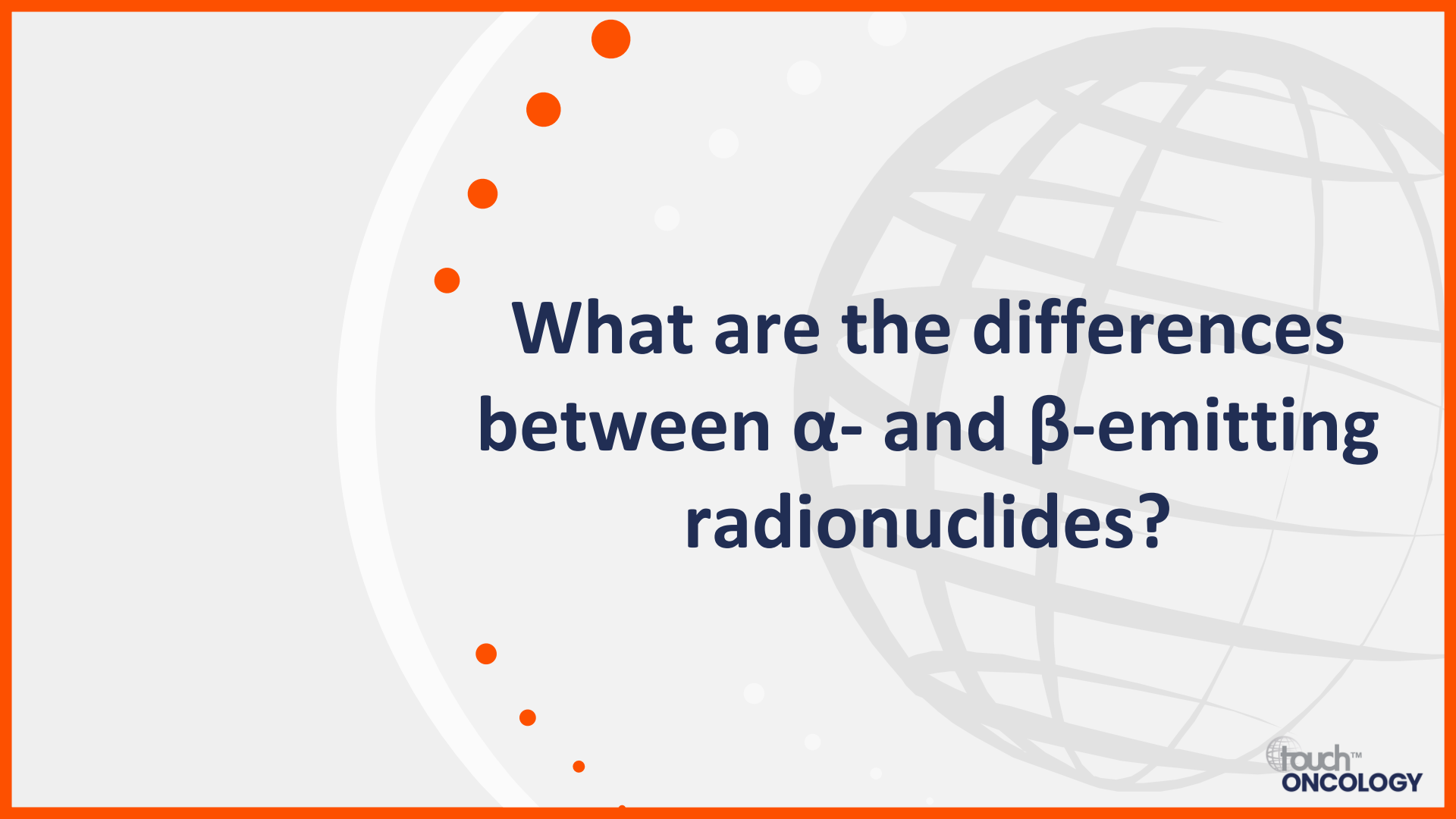
Molecular target
location and type

- Radiation type (α - or β -particles, or γ -rays)^{2,4}
- Half-life^{2,3}
- Daughter products²
- Biological clearance (e.g. renal)⁴

Image adapted from Holik HA, et al. 2022.⁵

1. Pouget JP, et al. *Nucl Med Biol.* 2022;104-5:53-64; 2. Kunos CA, et al. *Semin Radiat Oncol.* 2021;31:3-11; 3. Vermeulen K, et al. *Semin Nucl Med.* 2019;49:339-56;

4. Sgouros G, et al. *Nat Rev Drug Discov.* 2020;19:589-608; 5. Holik HA, et al. *Molecules.* 2022;27:3062.



**What are the differences
between α - and β -emitting
radionuclides?**

DNA damage mediated by α - and β -radiation

Particles with higher LET are more efficient at inducing DSBs

(normalized to tissue energy deposition)

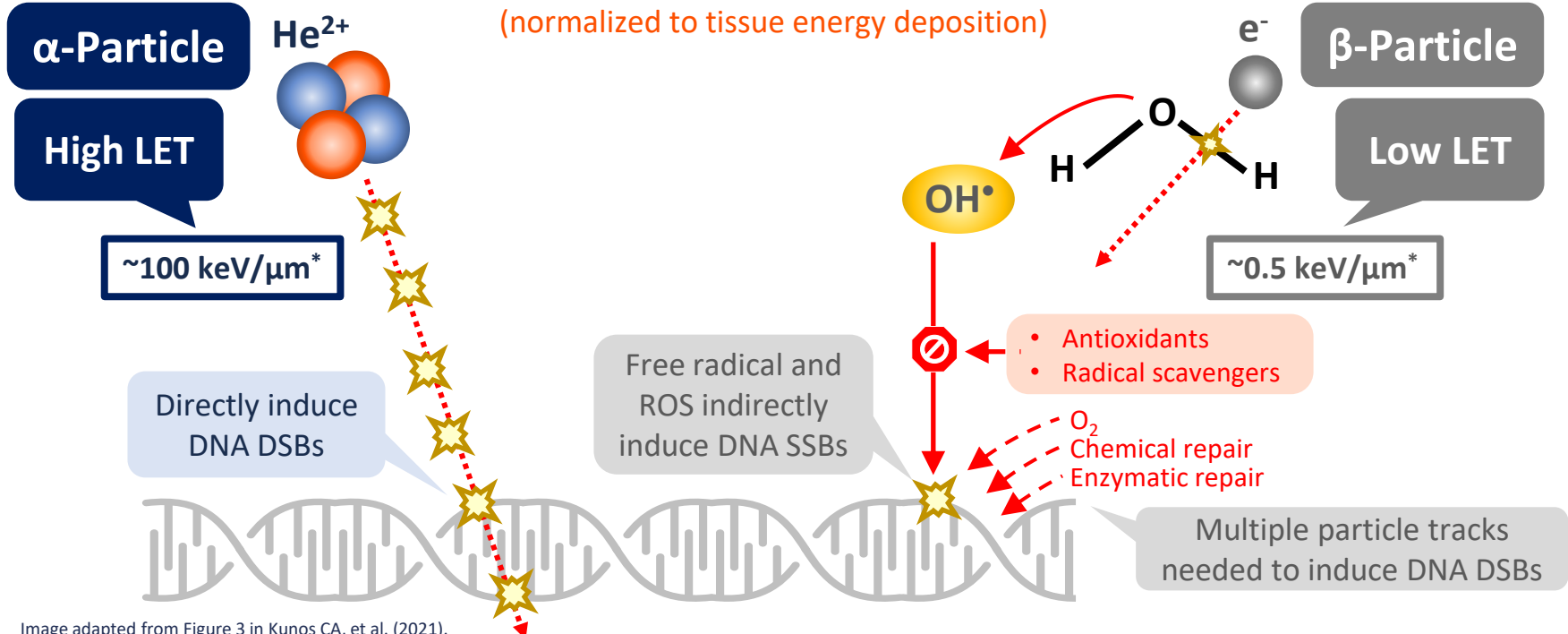


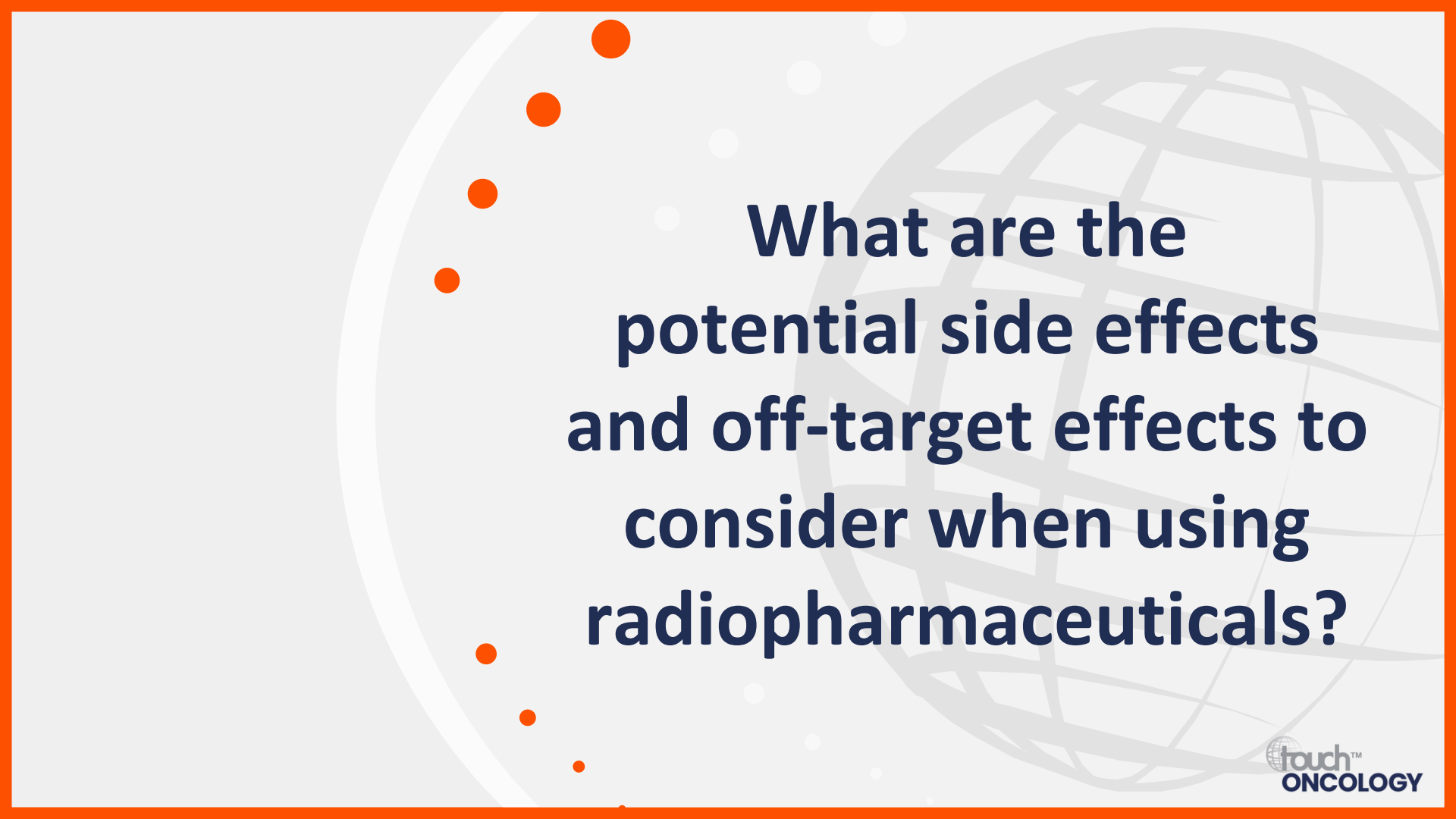
Image adapted from Figure 3 in Kunos CA, et al. (2021).

*Kiloelectronvolts per micrometre ($\text{keV}/\mu\text{m}$) is the standard LET unit of measure.

DSB, double-strand break; e, electron; LET, linear energy transfer; ROS, reactive oxygen species; SSB, single-strand break.

Kunos CA, et al. *Semin Radiat Oncol.* 2021;31:3–11.

- **What are the current approaches to dosimetry when using radiopharmaceuticals?**

The background of the slide features a large, faint globe with a grid of latitude and longitude lines. To the left of the globe, there is a vertical line of seven orange dots of varying sizes. The entire slide is framed by a thick orange border.

**What are the
potential side effects
and off-target effects to
consider when using
radiopharmaceuticals?**

Off-target effects and side effects to consider



Considerations to maximize clinical benefit and minimize off-target and side effects¹



Dosimetry based on absorbed dose to:¹

- Target tumour tissue?
- Non-tumour tissue (at-risk organs)?

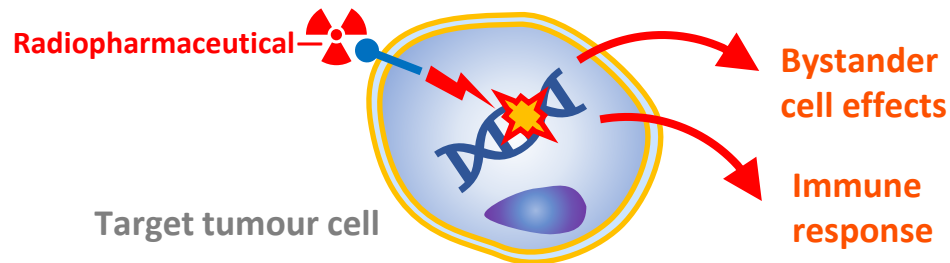
Dose-limiting tissues to consider:²

- Bone marrow
- Kidneys
- Liver
- Lungs
- Salivary glands



More clinical evidence is needed to understand implications of RPTs for dose-limiting tissues, and off-target and side effects²⁻⁴

Off-target effects^{3,4}



Modulation of cell DNA repair response to radiation^{3,4}

Deterministic effects

Stochastic effects

Repair

No repair

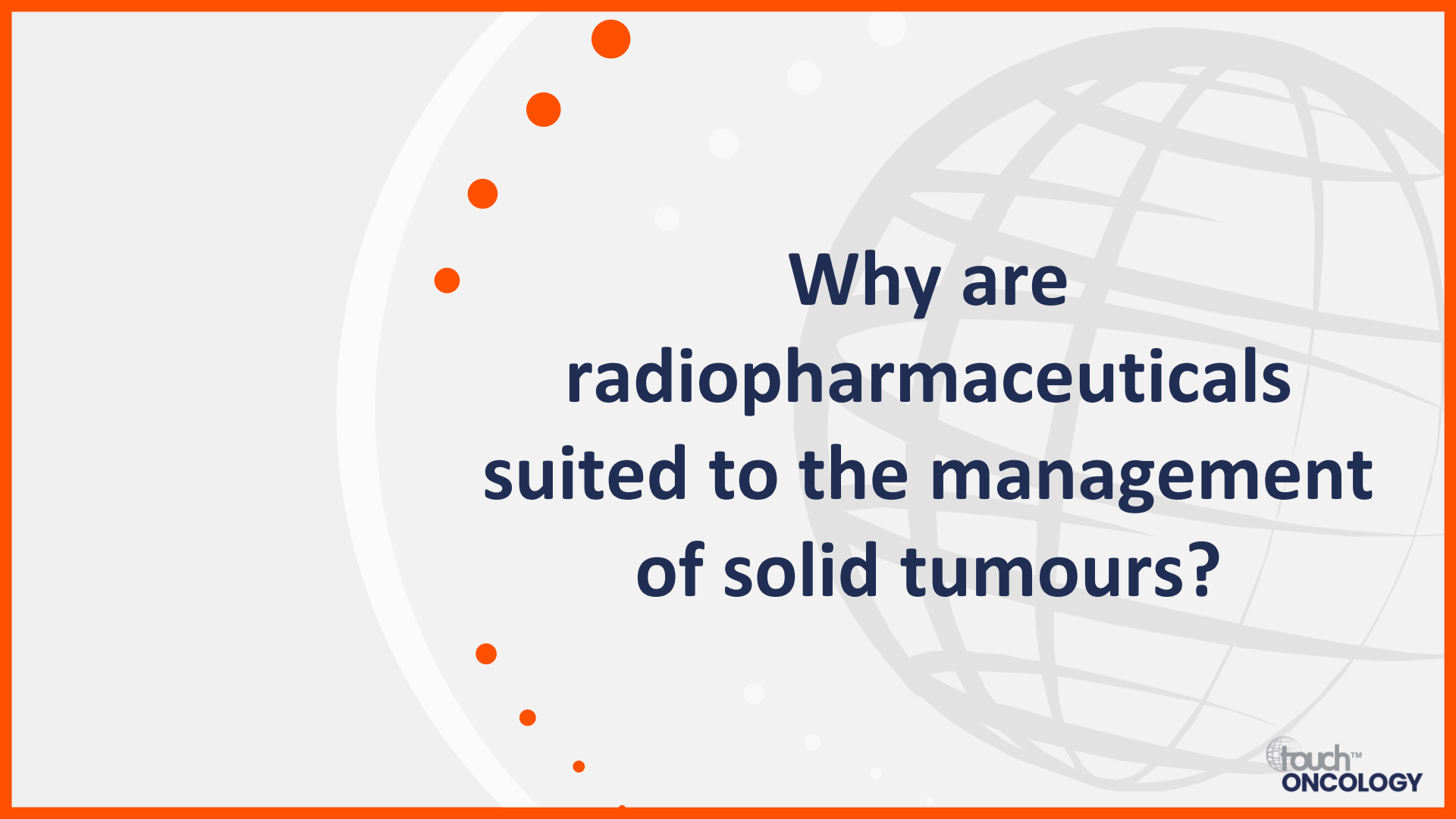
Misrepair

Cell survival

Cell death

- No meaningful change in cell biology
- Risk of carcinogenesis?

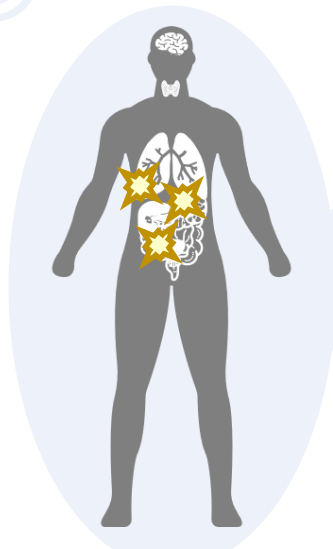
1. Lawhn-Heath C, et al. *Lancet Oncol.* 2022;23:e75-87; 2. Wahl RL, et al. *J Nucl Med.* 2021;62(Suppl. 3):235-355; 3. Pouget JP, et al. *Nuclear Med Biol.* 2022;104-5:53-64; 4. Pouget JP, et al. *Antioxid Redox Signal.* 2018;29:1447-87.



**Why are
radiopharmaceuticals
suited to the management
of solid tumours?**

Addressing unmet needs in solid tumours

Radiopharmaceuticals offer scope for personalized approaches in cancer management¹⁻⁶



Clinical benefit demonstrated in SSTR+ GEP-NETs,^{4,5} mCRPC⁶ and mPPGLs⁷



Systemic therapy able to localize to low volume metastatic disease not amenable to conventional therapy or not visible on radiographic imaging^{8,9}



Biological by-stander effects can induce immune response to systemic disease⁹⁻¹¹

Radiopharmaceuticals is an expanding field, with multiple agents in clinical development¹²

GEP-NET, gastroenteropancreatic neuroendocrine tumour;

mCRPC, metastatic castration-resistant prostate cancer; mPPGLs, metastatic pheochromocytomas and paragangliomas; SSTR, somatostatin receptor.

1. Kunos CA, et al. *Semin Radiat Oncol.* 2021;31:3-11; 2. Divgi C, et al. *Int J Radiat Oncol Biol Phys.* 2021;109:905-12; 3. Lawhn-Heath C, et al. *Lancet Oncol.* 2022;23:e75-87;

4. Strosberg J, et al. *N Engl J Med.* 2017;376:125-35; 5. Clement D, et al. *Eur J Nucl Med Mol Imaging.* 2022;49:3529-37; 6. Parker C, et al. *N Engl J Med.* 2013;369:213-23;

7. Severi S, et al. *ESMO Open.* 2021;6:10017; 8. Salih S, et al. *Molecules.* 2022;27:5231; 9. Sgouros G, et al. *J Nucl Med.* 2021;62(Suppl. 3):12S-22S;

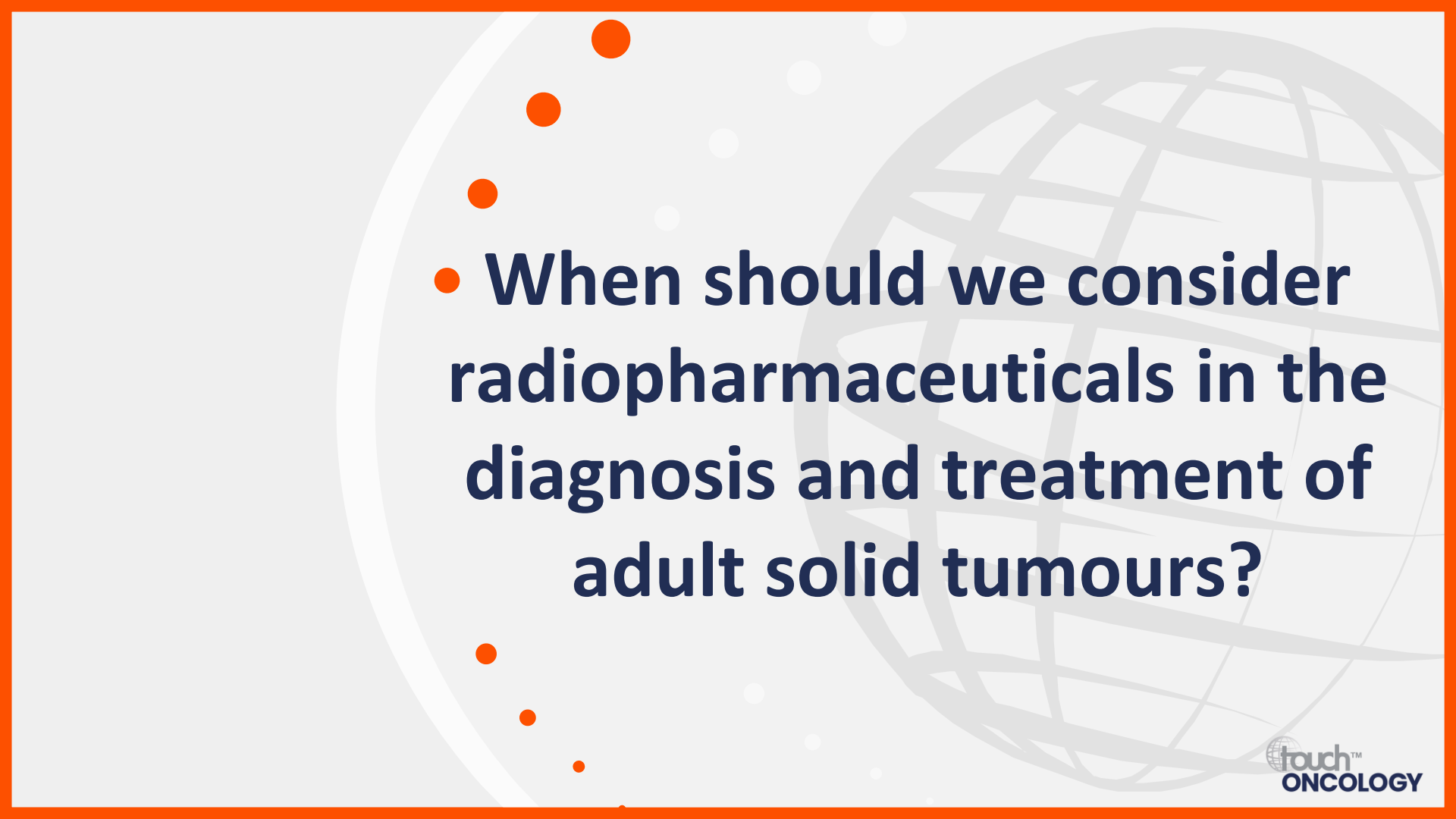
10. Pouget JP, et al. *Nuclear Med Biol.* 2022;104-5:53-64; 11. Pouget JP, et al. *Antioxid Redox Signal.* 2018;29:1447-87; 12. Sgouros G, et al. *Nat Rev Drug Discov.* 2020;19:589-608.

Understanding radiopharmaceutical therapy: One modality, many entities

Dr Ana P Kiess

Johns Hopkins University School of Medicine
Sidney Kimmel Comprehensive Cancer Center
Baltimore, MD, USA



- 
- **When should we consider radiopharmaceuticals in the diagnosis and treatment of adult solid tumours?**

Using radiopharmaceuticals in solid tumours



Applications and purpose



Imaging¹⁻⁴

- Diagnostic
- Monitoring



Treatment^{2,5,6}

- Curative intent
- Palliative management



Theranostics^{1,2,5}

- Imaging and/or treatment
- Image-guided therapy



Clinical considerations



Molecular targeting^{1,2,7,8}

- Tumour target(s)
- Tissue specificity



Biodistribution^{1-3,5,6}



Clearance and uptake¹⁻³



Absorbed dose^{1-3,6}

- Tumour response
- Potential toxicities

Image provided by corresponding faculty (Dr AP Kiess).

1. Korde A, et al. *EJNMMI Radiopharm Chem.* 2022;7:18; 2. Sgouros G, et al. *Nat Rev Drug Discov.* 2020;19:589–608; 3. Lawhn-Heath C, et al. *Lancet Oncol.* 2022;23:e75–87;

4. Schillaci O. *J Nucl Med.* 2014;55:357–9; 5. Kunos CA, et al. *Semin Radiat Oncol.* 2021;31:3–11; 6. O'Donoghue J, et al. *J Nucl Med.* 2022;63:1467–74;

7. Solnes LB, et al. *J Nucl Med.* 2020;61:311–8; 8. Salih S, et al. *Molecules.* 2022;27:5231.



What radiopharmaceutical modalities are available and/or in development?

Radiopharmaceutical constructs

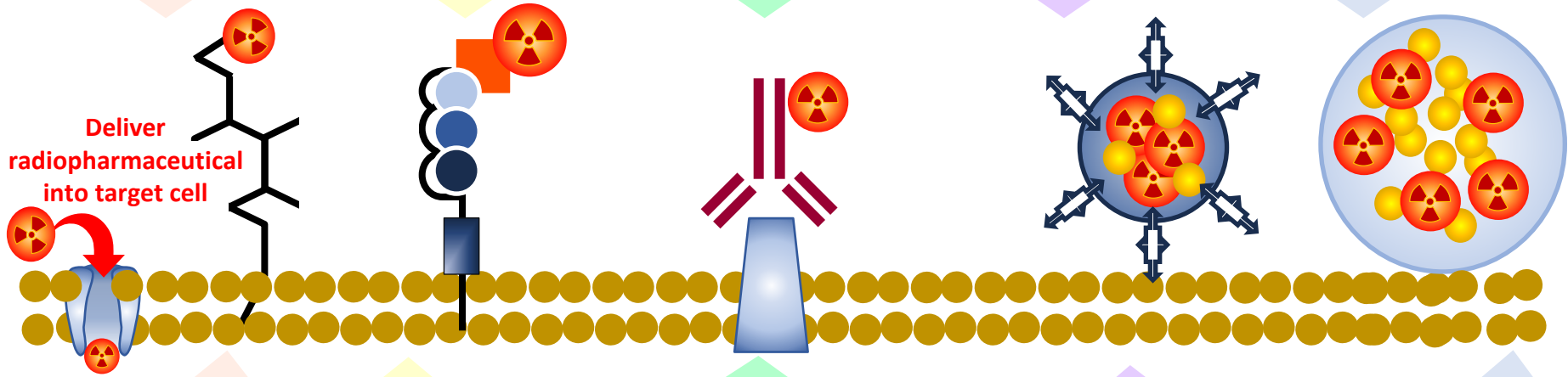
Small molecule

Peptide

Antibody

Nanoconstruct

Microsphere



Deliver radiopharmaceutical into target cell

FR; mIBG;
PL ether analogues
PSMA-targeting

PRRT ^{177}Lu -DOTATATE
Bombesin analogues

Antigen-targeting:
mesothelin; PSMA;
CD22; HER2; B7-H3

In preclinical
development

^{90}Y -microspheres
(glass; resin)

- Small molecules and peptides exhibit rapid targeting and clearance
- Shorter retention in target tissue

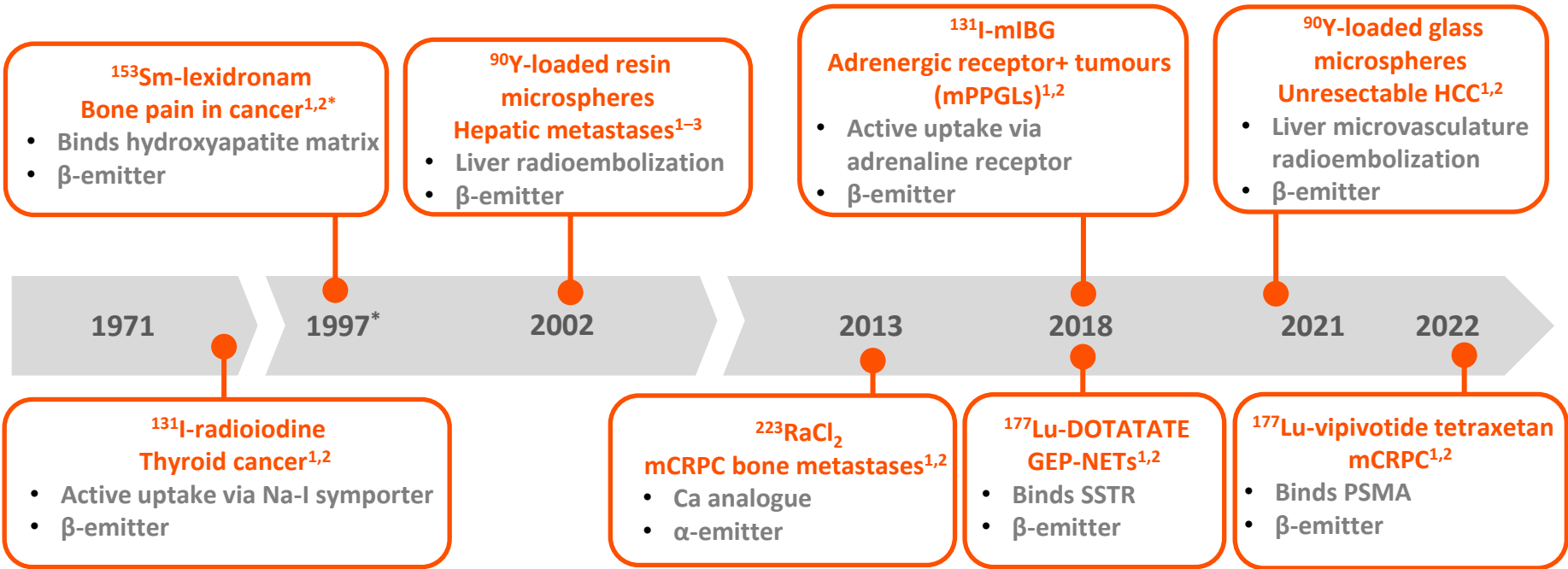
- **Bivalent delivery**
- **Longer retention due to circulating half-life; may lead to off-target toxicities (e.g. haematological)**

B7-H3, B7 homolog 3 protein; CD, cluster of differentiation; FR, folate receptor; HER2, human epidermal growth factor receptor-2; Lu, lutetium; mIBG, meta-iodobenzylguanidine; PL, phospholipid; PRRT, peptide receptor radionuclide therapy; PSMA, prostate membrane-specific antigen; Y, yttrium.
Sgouros G, et al. *Nat Rev Drug Discov.* 2020;19:589–608.

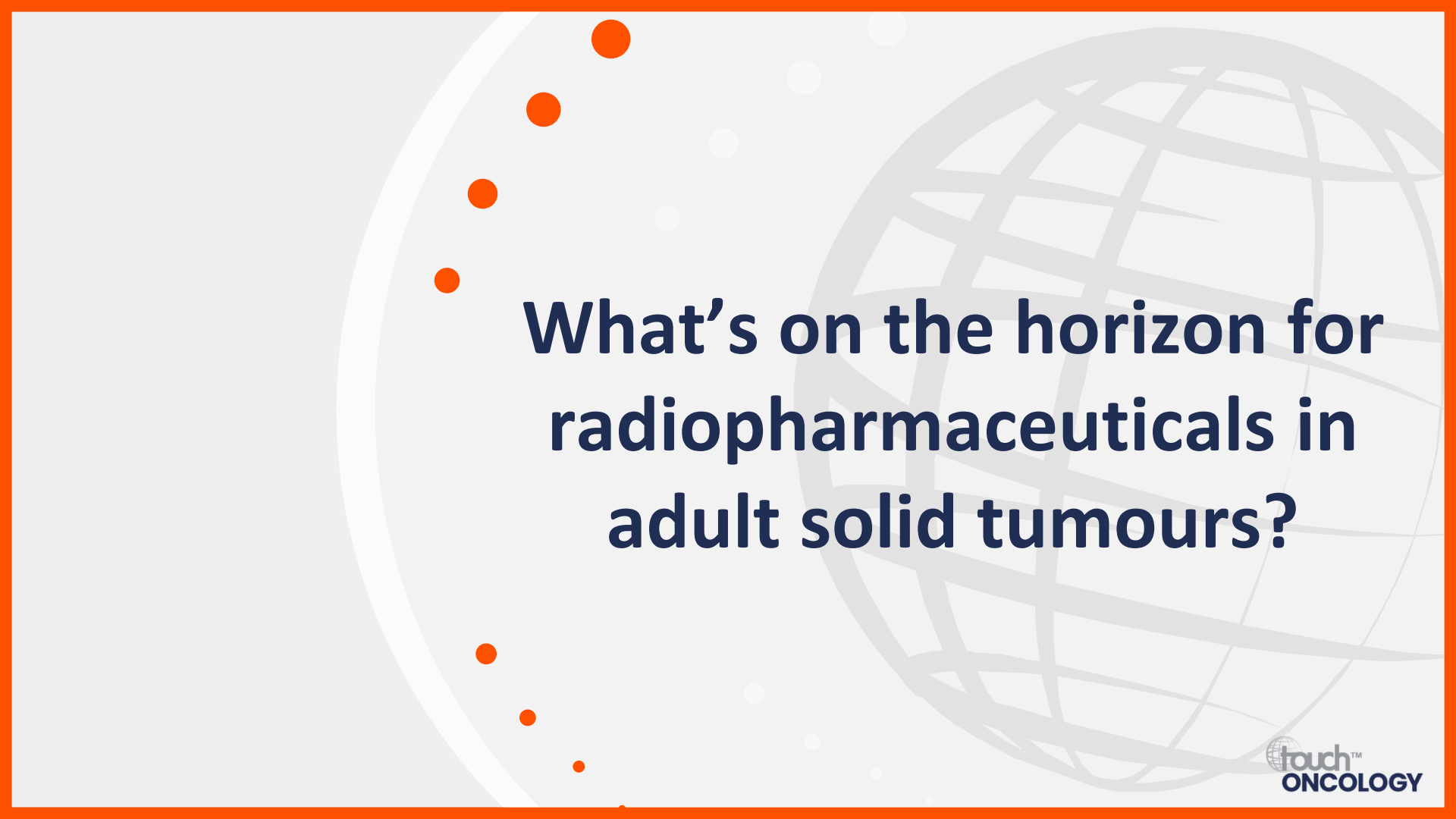


**What radiopharmaceuticals
are currently approved in
adult oncology indications?**

FDA-approved radiopharmaceuticals

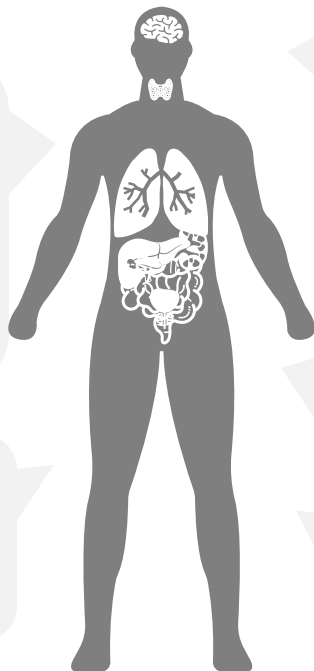


*¹⁵³Sm-lexidronam has been discontinued (production terminated by manufacturer). Ca, calcium; GEP, gastroenteropancreatic; HCC, hepatocellular carcinoma; I, iodine; Lu, lutetium; mCRPC, metastatic castration-resistant prostate cancer; mIBG, meta-iodobenzylguanidine; mPPGLs, metastatic pheochromocytomas and paragangliomas; Na, sodium; NET, neuroendocrine tumour; PSMA, prostate-specific membrane antigen; RaCl₂, radium chloride; SSTR, somatostatin receptor; Y, yttrium.
1. Sgouros G, et al. *Nat Rev Drug Discov.* 2020;19:589–608; 2. FDA prescribing information available and searchable by agent at <https://www.accessdata.fda.gov/scripts/cder/daf/> (accessed 23 March 2023); 3. Stubbs RS, Wickremesekera SK. *HPB (Oxford).* 2004;6:133–9.



**What's on the horizon for
radiopharmaceuticals in
adult solid tumours?**

Future of radiopharmaceuticals in solid tumours



Prostate cancer/tumour neovasculature

- ^{177}Lu -PNT2002 (PSMA-targeting)^{1,2}
- ^{227}Th -PSMA-TTC (PSMA-targeting)^{1,3}
- ^{225}Ac -PSMA-617 (PSMA-targeting)⁴
- ^{225}Ac -J591 (PSMA-targeting)⁵
- ^{225}Ac -DOTA-h11B6 (HK-2-targeting)⁶

GRPR+ advanced solid tumours (e.g. breast, prostate and GISTs)

- ^{177}Lu -NeoBOMB1 (GRPR-targeting)^{1,7}

Brain and CNS/DSRCT and other solid peritoneal tumours

- ^{131}I -omburtamab (B7-H3-targeting)^{1,8,9}

NETs


- ^{177}Lu -satoreotide tetraxetan (SSTR-targeting)^{1,10}
- ^{68}Ga -DOTA-JR11 (SSTR-targeting)^{1,10}
- ^{68}Ga -satoreotide trizoxetan (SSTR-targeting)^{1,11}
- ^{212}Pb -DOTAMTATE (SSTR-targeting)^{1,12}

Advanced stage solid tumours/adenocarcinomas

- $^{177}\text{Lu}/^{90}\text{Y}$ -FAP-46 (FAP-targeting)^{13–15}
- ^{177}Lu -FAP-2286 (FAP-targeting)^{13,16}

Ac, actinium; B7-H3, B7 homolog 3 protein ;CNS, central nervous system; DSRCT, desmoplastic small round cell tumour; FAP, fibroblast activation protein; Ga, gallium; GIST, gastrointestinal stromal tumour; GRPR, gastrin-resistant peptide receptor; HK-2, human kallikrein-2; I, iodine; Lu, lutetium; NET, neuroendocrine tumour; Pb, lead; PSMA, prostate-specific membrane antigen; SSTR, somatostatin receptor; TCC, targeted thorium conjugate; Th, thorium; Y, yttrium.

1. Sgouros G, et al. *Nat Rev Drug Discov.* 2020;19:589–608; 2. NCT04647526; 3. NCT03724747; 4. NCT04597411; 5. NCT03276572; 6. NCT04644770; 7. NCT03872778; 8. NCT05064306; 9. NCT04022213; 10. NCT02609737; 11. NCT03220217; 12. NCT03466216; 13. Calais J. *J Nucl Med.* 2020;61:163–5; 14. Liu Y, et al. *Eur J Nucl Med Mol Imaging.* 2022;49:871–80; 15. Ferdinandus J, et al. *J Nucl Med.* 2022;63:727–34; 16. Baum RP, et al. *J Nucl Med.* 2022;63:415–23. All trial information available at: <https://clinicaltrials.gov/> (accessed 22 March 2023).

The background of the slide features a large, light gray globe with a grid of latitude and longitude lines. To the left of the globe, there is a vertical line of seven orange dots of varying sizes, arranged in a slightly curved pattern. The entire slide is framed by a thick orange border.

**What more is needed
to support integration
of radiopharmaceuticals
into clinical pathways in
adult oncology?**

Radiopharmaceuticals: An interdisciplinary endeavour

Expanding knowledge and multidisciplinary team involvement¹

Steps to realizing the potential of radiopharmaceuticals



Wider access to education and clinical training to expand access to expertise (e.g. training in radionuclide dosimetry)^{2,3}



Frameworks for multidisciplinary collaboration^{1,3}



High-quality evidence to support use³



Addressing healthcare infrastructure needs (e.g. staff and additional imaging costs)³



Optimizing patient communication²⁻⁴

1. Sgouros G, et al. *Nat Rev Drug Discov.* 2020;19:589–608; 2. Divgi C, et al. *Int J Radiat Oncol Biol Phys.* 2021;109:905–12; 3. Lawhn-Heath C, et al. *Lancet Oncol.* 2022;23:e75–87; 4. Kohl P, et al. *Front Nucl Med.* 2023;3:1127692.