



RADIOPHARM THERANOSTICS

NASDAQ: RADX / ASX: RAD

COMPANY PRESENTATION

MARCH 2026



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Company Strategic Positioning

Develop first in class Radiopharmaceutical Therapies

Highly differentiated TARGETS

(all 4 currently in Phase I trials)

PD-L1
HER2
B7H3
KLK3

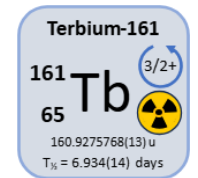
} First in Clinical Development

Protein-based MOLECULES

Half-life-engineered proteins
for optimal biodistribution

Different Size, Fc mutations

Clinically- proven β emitter ISOTOPES



IMAGING BUSINESS

Phase II trial in Brain Mets (close to completion)
- significant commercial potential -

Optionality for continuing development in Phase III or partnering

BUSINESS MODEL

Capital-efficient
Distributed Development
Best-in-class Global Partners

R&D Joint-Venture with



FINANCIALS

CASH runway Q1 2027
ATM available USD 18.9m

LARGEST holder ~15%



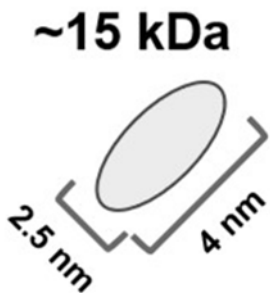
Half-life-engineered proteins for optimal biodistribution

two distinct platforms for our therapeutic molecules

NANOBODIES

BENEFITS OF NANOBODIES

- + Specificity and affinity
- + Binds to different epitopes than approved full-sized antibodies
- + Improved tumor penetration and accumulation (small size)
- + Rapid blood clearance
- + **Non-binding structure can be modified to enhance Human Biodistribution**

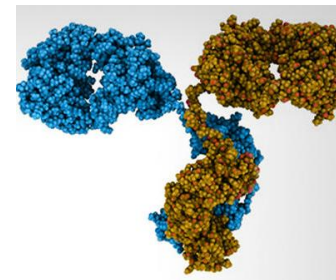


Proprietary platform		
PDL1		CLINICAL
HER2		CLINICAL
PTK7		Preclinical
MDAnderson Cancer Center	Target #1	Preclinical
MDAnderson Cancer Center	Target #2	Preclinical

MONOCLONAL ANTIBODIES

BENEFITS OF MONOCLONAL ANTIBODIES

- + Specificity and affinity
- + Liver Excretion
- + No / limited kidney accumulation
- + Sustained tumor uptake
- + **Structures can be modified in the Fc regions to enhance Human Biodistribution**



mAb platform	
B7H3	CLINICAL
<i>Dual modification at FcRn & FcγR</i>	
KLK3	CLINICAL

Capital-Efficient, Asset-Centric, Distributed Development Model

Experienced, lean team of 17 FTEs leveraging best-in-class global partners



Riccardo Canevari
CEO

- Radiopharm Theranostics CEO since September 2021
- Previously: Chief Commercial Officer of Novartis Company Advanced Accelerator Applications S.A.
- Lead for Lutathera in-market growth strategy & Pluvicto launch strategy
- Senior Vice President & Global Head, Breast Cancer for Novartis Oncology since 2017



Dr. Dimitris Voliotis
CMO

- Radiopharm Theranostics CMO since August 2024
- Previously: SVP Global Development at Convergent Tx and Zentalis Pharma
- Chief Development Officer at CureVac
- Global Head of Clinical Development at Eisai and Bayer



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Clinical Ops



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Manager, CMC



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Clinical Ops



Nicole Villari
Quality Affairs
GMP Manager

Therapeutic Pipeline – Clinical stage and *First-in-Class Radiotherapies*

		PROGRAM	TARGET	INDICATION	ISOTOPE	PRECLINICAL	PHASE I	PHASE II	NOTES
THERAPEUTIC TRIALS	Nanobody Platform	RAD204	PD-L1	PD-L1+ solid tumors	Lu177				Phase I in 4 AUS centers, NCT06305962 Dose Level #3 at 90mCi recruiting
		RAD202	HER2	HER2+ solid tumors	Lu177				Phase I in 5 AUS centers NCT06824155 Dose Level #2 at 75mCi recruiting
	mAb platform	RV01	B7-H3	B7-H3+ solid tumors	Lu177				Phase I in 5 US centers NCT07189871 Dose Level #1 at 35mCi recruiting. First patient dosed.
		RAD402	KLK3	Advanced prostate cancer (>90% express KLK3)	Tb161				Phase I in 6 AUS centers NCT07259213 Dose Level #1 at 30mCi recruiting

	2026			
PROGRAM	Q1	Q2	Q3	Q4
RAD204				Data read out Phase I dose escalation
RAD202				Data read out Phase I dose escalation
RV01		Interim Data first two cohorts (Human biodistribution)		
RAD402		Interim Data first two cohorts (Human biodistribution)		



Molecule: **^{177}Lu -RAD204**

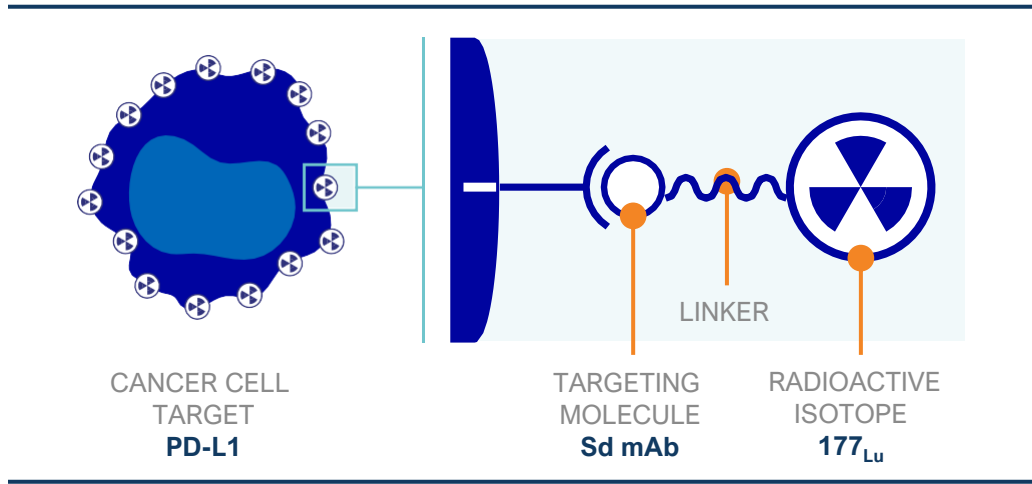
Targeting MoA: **PD-L1**

Therapeutic for: **Multiple Solid Tumors**

COMPETITIVE POSITION

*First in class PDL-1 targeting
radiotherapy in Clinical Development*

RAD 204 utilizes an anti-PD-L1 Nanobody as a targeting moiety



~15 kDa

2.5 nm

4 nm

Anti-PD-L1 Nanobody
High affinity single domain monoclonal antibody

PD-L1 Immune Checkpoint

- Antigen expression mediates evasion of immune responses by cancer cells
- Inhibition leads to antitumor activity

THERAPEUTIC APPLICATION

Expected positioning in the treatment algorithm

Post-IO Non-Small Cell Lung Cancer

Success bar:

docetaxel + ramucirumab with reported ORR ~28.8%, median PFS ~4.1 months.

Phase 1 Trial Design

^{177}Lu -anti-PD-L1 single domain AB in metastatic solid tumors

Primary Objectives

- Safety and tolerability of ^{177}Lu -RAD204
- Recommended ph2 dose of ^{177}Lu -RAD204

Study Design

BOIN for escalation / de-escalation.

Population: History of PD-L1 positive ($\geq 1\%$) metastatic tumors

Imaging Phase 0

Biodistribution, dosimetry and PK with low dose ^{177}Lu -RAD204 in organs of interest and tumor

Therapeutic Phase 1

^{177}Lu -RAD204 dose escalation

	Dose Level	Dose
Phase 0 (Imaging Period with ^{177}Lu -RAD204)	Imaging dose	10 (0.37 GBq)
	Therapeutic DL1	30 mCi (1.1. GBq) ✓
Phase I (Treatment Period with ^{177}Lu -RAD204 _{tr})	DL2	60 mCi (2.2 GBq) ✓
	DL3	90 mCi (3.3 GBq)
	DL4	TBD

Clinical data Phase I

- First (30mCi) and Second Cohort (60mCi) completed, and 6 patient data released.
- Tumor uptake confirmed in all the treated subjects. No tumor reduction above 30% achieved at the first two dose levels
- The safety profile has been very favorable, with few adverse events and no related SAEs observed.
- Currently recruiting Third Cohort (90mCi).



Molecule: **177Lu-RAD202**

Targeting MoA: **HER2**

Therapeutic for: **HER2+ TUMORS**

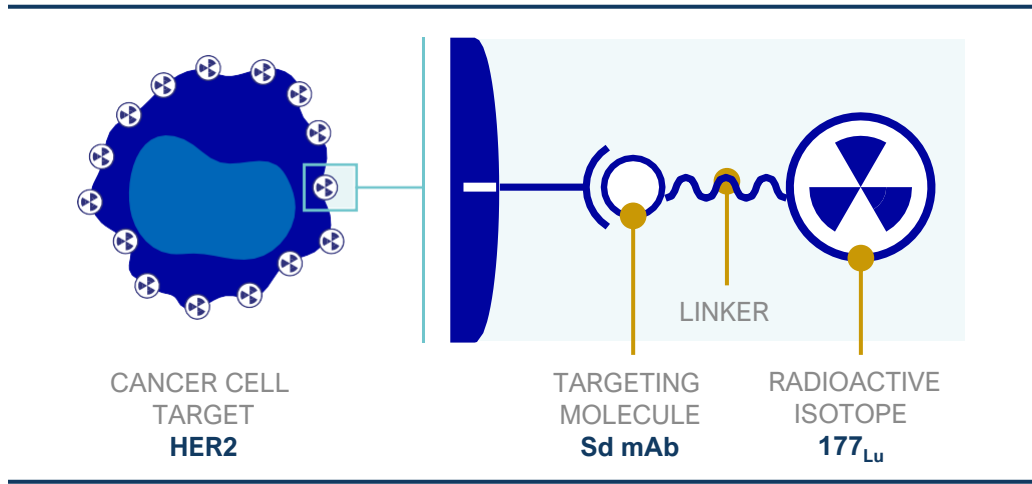
COMPETITIVE POSITION

*Most advanced HER2 radiotherapy in
Clinical Development*

Ahead of:

*Novartis
Clarity*

RAD 202 utilizes an anti-HER2 Nanobody as a targeting moiety



HER2 NANOBODY

High specificity & affinity single-domain antibody

- HER2 pathway well validated
- Overexpression in breast, and gastroesophageal cancers
- Improved tumor penetration, accumulation and rapid blood clearance (small size)

~ 15 kDa

2.5 nm

4 nm

THERAPEUTIC APPLICATION

Expected positioning in the treatment algorithm

Post-Trastuzumab DXd Breast Cancer

Success bar:

tucatinib + trastuzumab + capecitabine (TTC) reported ORR 32.6%, Median PFS 4.7 months

Phase 1 Trial Design

'HEAT' Trial (HER2 Antibody Therapy with Lutetium-177) in subjects with HER2+ advanced solid tumors

Primary Objectives (Phase 1, Treatment):

- Safety and tolerability of ^{177}Lu -RAD202
- Recommended ph2 dose of ^{177}Lu -RAD202

Population:

Her2+ (IHC, ISH) a/m solid tumors

Phase 0 Imaging:

Biodistribution, PK and radiation dosimetry of ^{177}Lu -RAD202_{im} in organs of interest and tumor lesions

Phase I Therapeutic:

^{177}Lu -RAD202_{tr} dose escalation

	Dose Level	Dose
Phase 0 (Imaging Period with ^{177}Lu - RAD202 _{im})	Imaging dose	10 mCi
Phase I (Treatment Period with ^{177}Lu - RAD202 _{tr})	Therapeutic DL1	30 mCi (1.1 GBq) ✓
	DL2	75 mCi (2.7 GBq)
	DL3+	TBD

Clinical Data Phase I

- First Cohort completed (30 mCi), with 3 Patient data released
- Significant tumor uptake observed
- The safety profile very favorable, with few low-grade adverse events and no SAEs observed thus far

- Currently recruiting Cohort #2 at 75mCi

Tumor Uptake | Very High Uptake in the First 3 Patients

PATIENT #1			Absorbed Dose at 30 mCi
Cycle	Lesion	Volume (ml) ²	Dose (Gy), with PVC ^{1,2}
C1	ROI-2	2.26025	3.57
C1	ROI-3	3.634	2.07
C1	ROI-4	10.36125	2.02
C1	ROI-5	16.20966667	0.39

PATIENT #2			Absorbed Dose at 30 mCi
Cycle	Lesion	Volume (ml)	Dose (Gy), with PVC ^{1,2}
C1	ROI-3	5.821	2.732
C1	ROI-6	23.02025	1.581
C1	ROI-7	43.224	1.831
C1	ROI-8	145.1585	1.286
C1	ROI-9	16.796	2.084
C1	ROI-10	20.9355	2.092
C1	ROI-11	20.31075	2.959
C1	ROI-12	20.31075	1.558
C1	ROI-13	30.34125	0.854

PATIENT #3			Absorbed Dose at 30 mCi
Cycle	Lesion (refer to Viedoc for lesion's location for each ROI)	Volume (ml)	Dose (Gy), with PVC ^{1,2}
C1	ROI-2	48.7975	0.848
C1	ROI-3	60.502	0.661
C1	ROI-4	25.8	0.793
C1	ROI-6	26.85	0.964
C1	ROI-11	17.256	1.235

¹Partial Volume Correction applied.

²Density of lesion: soft tissue = 1.0 g/mL. Bone = 1.3 g/mL

³Lesions were contours based on thresholding (40%) method and volume was averaged over all timepoints

*BR = background – shoulder and proximal thigh. T:BR = lesion SUVmax:BR SUVmean

GBq, gigabecquerel; Gy, gray; ml, milliliter; PVC, partial volume correction; ROI, region of interest; SUV, standardized uptake value; SUVmax, maximum standardized uptake value; SUVmean, mean standardized uptake value; TBR, target-to-background ratio.

Intellectual Property

THE UNIVERSITY OF TEXAS
MDAnderson
~~Cancer Center~~
Making Cancer History®



RADIOPHARM THERANOSTICS

Molecule: **RV01/BetaBart**

Targeting MoA: **B7H3**

Therapeutic for: **Multiple Solid Tumors**

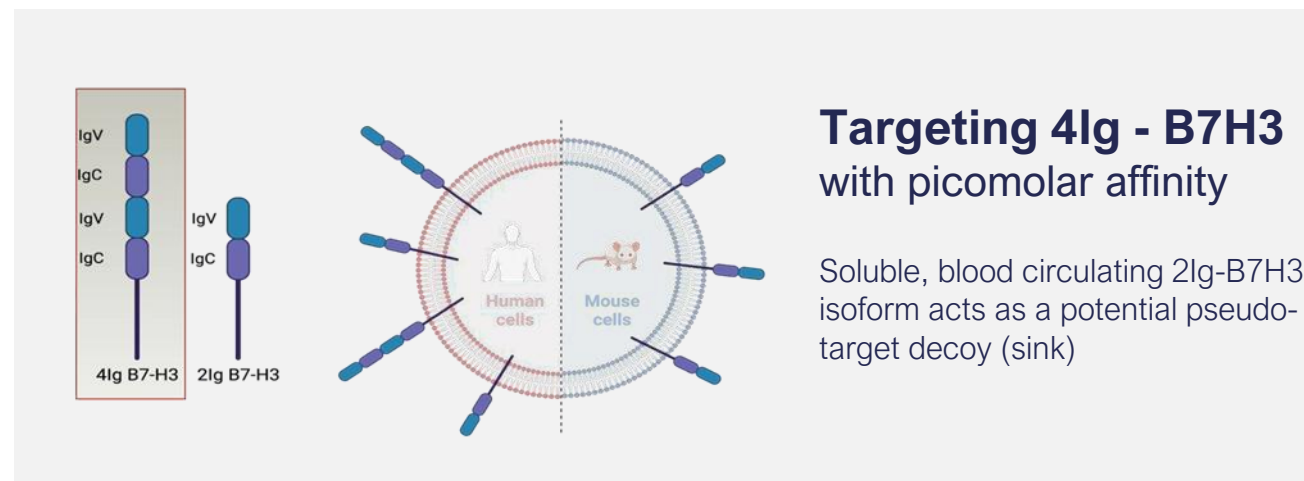
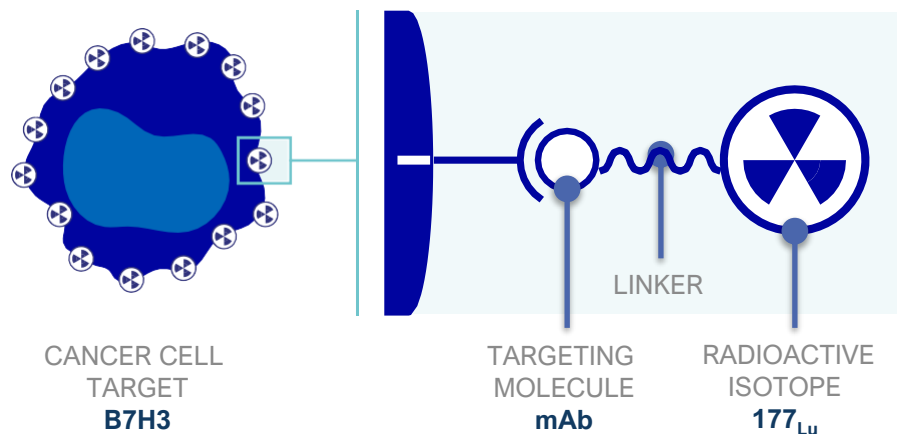
COMPETITIVE POSITION

*Most advanced B7H3 radiotherapy in
Clinical Development*

Ahead of:

*Novartis
Aktis Oncology*

RV 01 (Betabart) first in class selective B7H3 in clinical development



Fc-mutated mAb to improve Biodistribution

- + Reduced affinity for FcRn
 - Faster hepatic excretion (no re-circulation)
- + Reduced affinity for FcγR
 - Reduced bone marrow affinity

Isotope Selection



- + Favorable Therapeutic Index
- + Solid supply Chain
- + Cross-fire effect in small & large lesions

Phase 1 Trial Design

RV 01 (Betabart) basket trial in metastatic solid tumors

Primary Objectives (Phase 1, Treatment):

- Safety and tolerability of ^{177}Lu -RV01
- Recommended ph2 dose of ^{177}Lu -RV01

Population:

All comers, with advanced/metastatic solid tumors:

prostate, lung (NSCLC & SCLC), colorectal, breast(TNBC), H&N, ovarian, endometrial

First patient dosed on Feb 23, 2026

	Dose Level	Dose
Phase I (Treatment Period with ^{177}Lu - RV01)	DL1	35 mCi (1.3 GBq)
	DL2	

B7H3 Radiotherapies: Competitive Situation

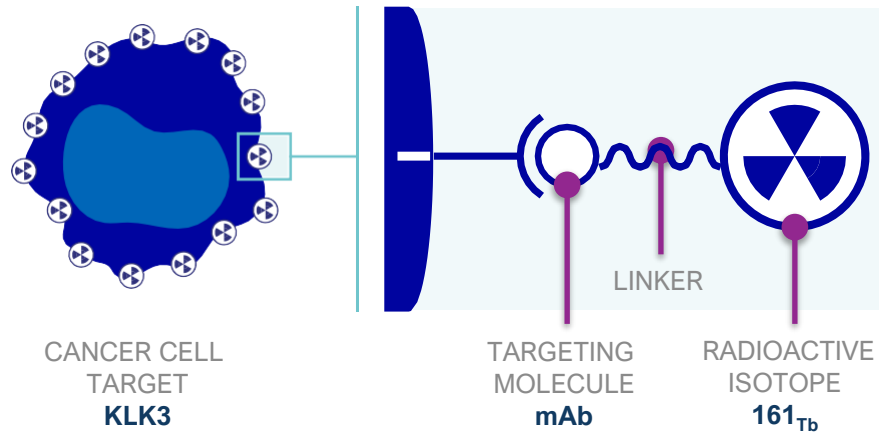
COMPANY	PROGRAM	MOLECULE	INDICATION	ISOTOPE	Pre-IND	IND Granted	Phase I First patient dosed	Phase I first two cohorts completed
RADIOPHARM THERANOSTICS	RV01	mAb with dual Fc modification to reduce blood half-life and bone marrow affinity	Basket trial in multiple solid tumors	Lu177				
AKTIS Oncology	AKY-2519	Mini-protein	Basket trial in multiple solid tumors	Ac 225				
NOVARTIS / MARIANA ONCOLOGY	MC 760	Peptide	Basket trial in multiple solid tumors	?				

Intellectual PropertyMemorial Sloan Kettering
Cancer Center

RADIOPHARM THERANOSTICS

Molecule: RAD 402 – 161Tb**Targeting MoA: KLK3****Therapeutic for: PROSTATE CANCER****COMPETITIVE POSITION***First in class KLK3 radiotherapy in
Clinical Development**First in class Tb161 Company
Sponsored Clinical Trial*

RAD 402 first in class KLK3 targeting mAb



Humanized IgG1

Specifically binding KLK3

High affinity

Internalized by Prostate Cells

UNIQUE BENEFITS IN POST PLUVICTO SETTINGS

- + mAb (liver excretion) avoids excess of kidney uptake after 4-6 cycles of ^{177}Lu -PSMA peptide (kidney excretion)
- + MoA switch to KLK3 after patients become refractory to PSMA targeting treatment
- + Tb161 dual emission (Beta+Auger) has the potential to improve therapeutic efficacy with a favorable safety profile

THERAPEUTIC APPLICATION

Expected positioning in the treatment algorithm

Post-Pluvicto Prostate Cancer

Success bar

SoC post Pluvicto:

Cabazitaxel-containing regimens reported PSA50 at ~28%, rPFS 3-4 months

In Clin Dev:

Ac225-PSMA-617 reported PSA50 at ~65% - still undefined Safety Profile

Phase 1 Trial Design

^{161}Tb -RAD 402 in metastatic Prostate Cancer

Primary Objectives (Phase 1, Treatment):

- Safety and tolerability of ^{161}Tb -RAD 402
- Recommended ph2 dose of ^{161}Tb -RAD 402



Population:

All comers, with advanced/metastatic prostate cancer

	Dose Level	Dose
Phase I (Treatment Period with ^{161}Tb - RAD 402)	DL1	30 mCi (1.1 GBq)
	DL2	

IMAGING

Imaging Pipeline – Clinical Stage and First-in-Class Diagnostics

MOLECULE	PROGRAM	TARGET	INDICATION	ISOTOPE	PHASE I	PHASE II	NOTES
Small Molecule	RAD101	Short Chain Fatty Acid	Brain Mets	F18			Phase 2b in 5 US centers, NCT06777433 12-patient interim analysis released (12/25) Expect to complete enrollment 1Q26
Peptide	RAD301	Integrin AvB6	Pancreatic /NSCLC	Ga68			Phase I enrolling, NCT05799274 8 pts dosed / 9 total

	2026			
PROGRAM	Q1	Q2	Q3	Q4
RAD101		Data read out Phase II primary Endpoint		
RAD 301		Data read out Phase I		



Molecule: **18F-RAD101**

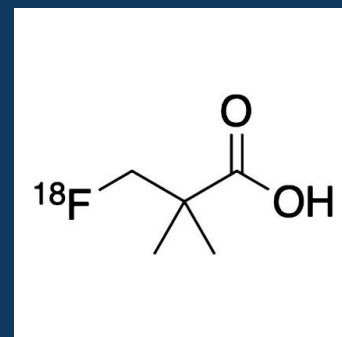
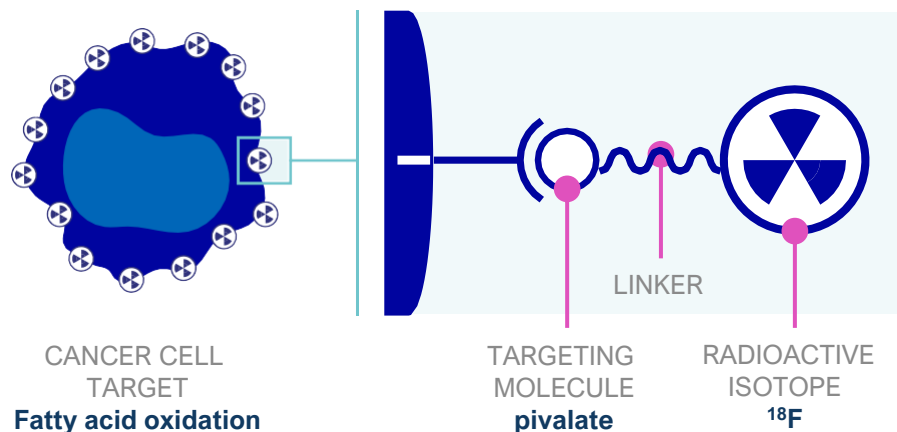
Targeting MoA: **SHORT CHAIN FATTY ACIDS**

Imaging for: **SUSPECTED RECURRENT BRAIN METASTASES**

COMPETITIVE POSITION

*First in class PET agent in Advanced
Clinical Development in Brain Mets*

Imaging for Brain Metastasis



RAD 101 (PIVALATE) SMALL MOLECULE

Selectively targets fatty acid synthase:
overexpressed in tumors but not normal brain cells



FATTY ACID SYNTHASE IS A VIABLE TARGET

- + Upregulation of de novo fatty acid synthesis via Fatty Acid Synthase (FASN) enables cancer cells to grow in lipid-deprived brain microenvironment.
- + Disruption of FASN activity can impair growth of brain metastases, representing a viable therapeutic target.

IMAGING

- + First-in-class Phase IIb imaging study currently recruiting (US).*
- + High unmet need to detect early relapse after Stereotactic Radio Surgery in brain metastases from solid tumors of different origin
- + ~300,000 new subjects diagnosed every year (US only)

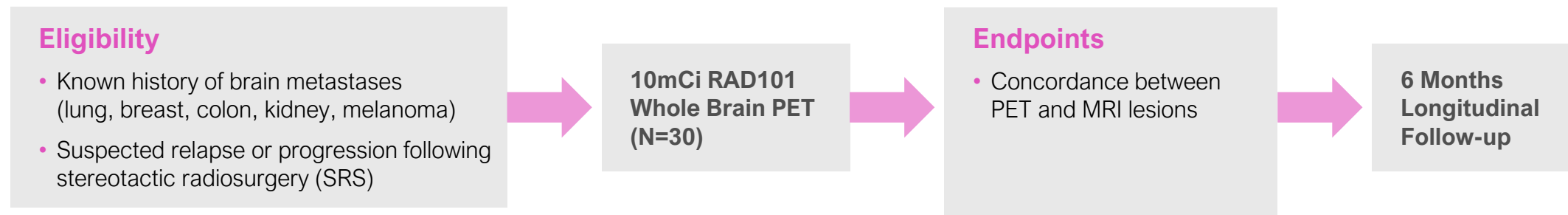
RAD 101 Imaging: Clinical Development

- Phase IIb imaging study currently recruiting in five centers in USA; 50% enrollment achieved.
- No competitor identified; RAD 101 is the only PET agent in clinical development for Brain Mets
- Large Total addressable market: 300,000 new subjects diagnosed every year (US only)

PRECLINICAL	PHASE I	PHASE IIa	PHASE IIb	PHASE III
	UK	UK	USA	
	24 pts	22 pts	30 pts	150 pts

Phase 2b Trial Design

Phase IIb imaging study in participants with suspected recurrent brain metastases from solid tumors



- **Study Design:**

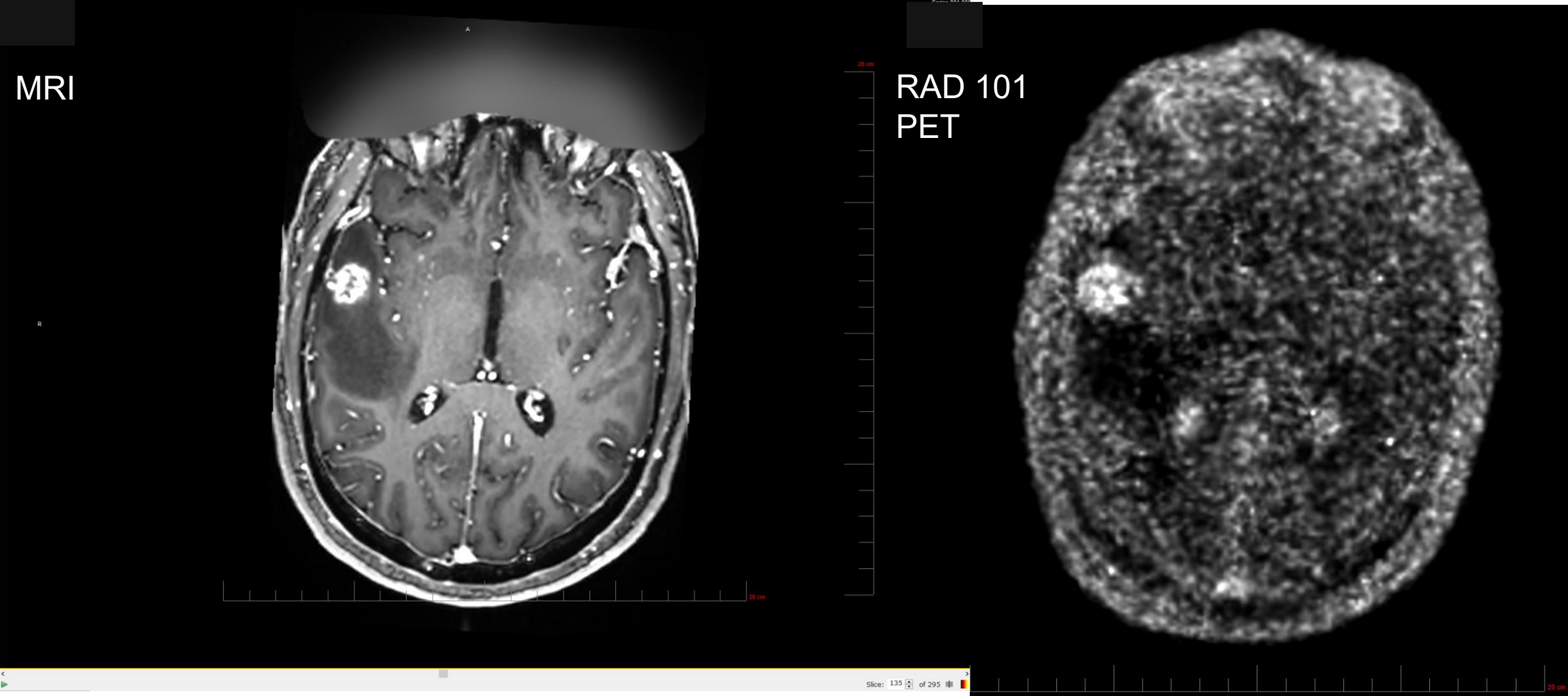
Single dose RAD101, max 370 MBq (10 mCi), administered IV followed by whole brain PET/MRI scan at 60 ± 10 min post-dose. Four-week screening period, 3-day imaging and safety follow-up, longitudinal imaging and data collection up to 6 months. Study size: n=30.

Clinical Data from Phase IIb

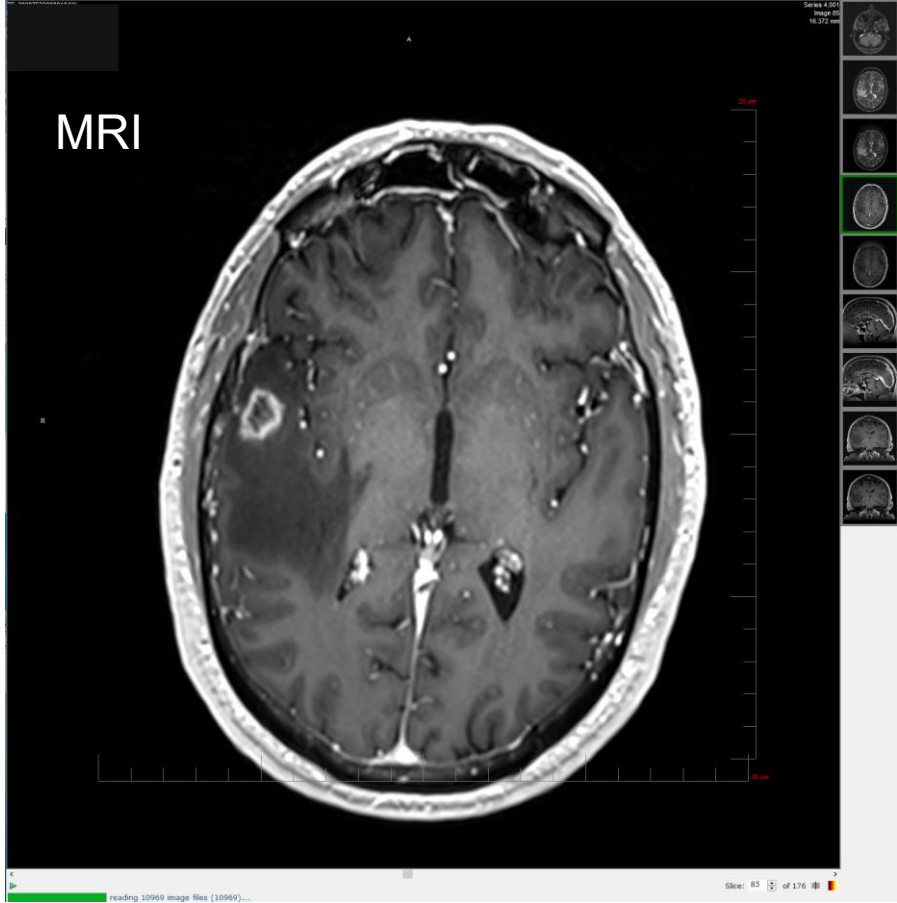
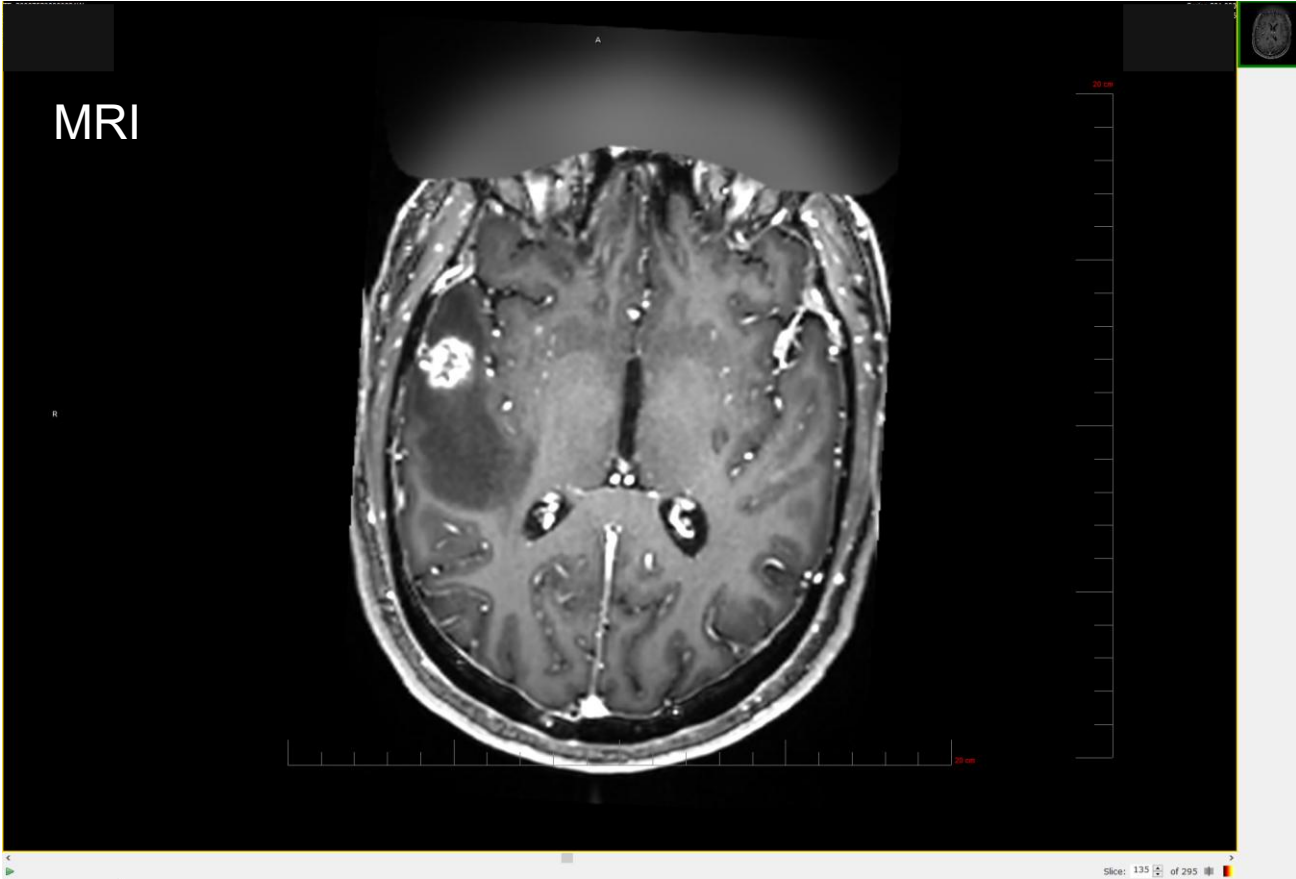
- Interim analysis in 12 patients in the ongoing study released (Dec 2025)
- 11/12 patients (92%) achieved the primary endpoint (Concordance with MRI), with increased metabolic activity in areas with equivocal MRI findings (suspected relapse)
- N=15 subjects dosed as of 11/15/2025
- Images from 6 patients included in this deck, as a representative example of the Interim results

Subject #1 Visit 1 - Oct'25

Concordance between MRI (contrast uptake plus perilesional hypodensity) & PET (tracer uptake)

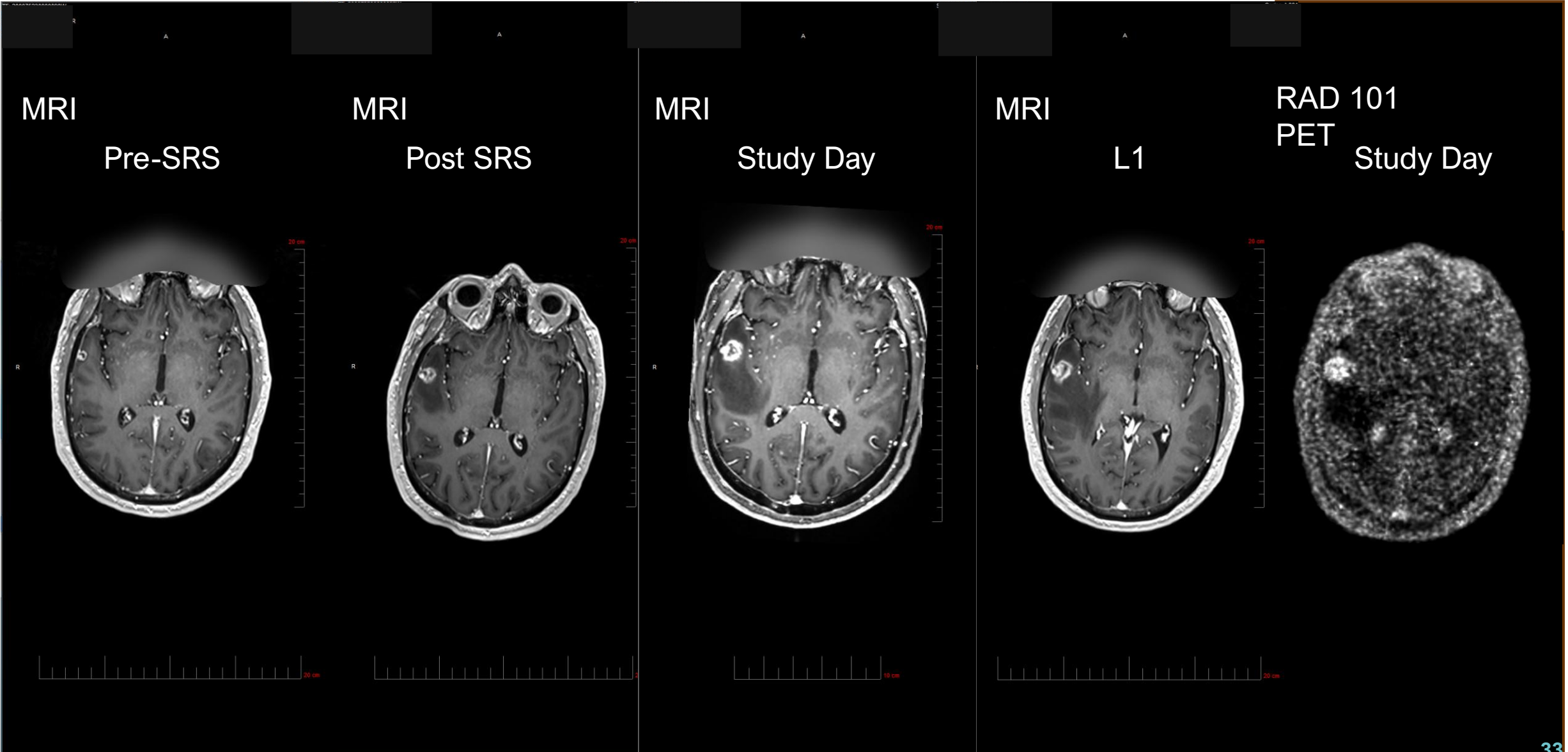


Subject #1 MRI Visit 1 (Oct'25) vs Long 1 (Nov'25)



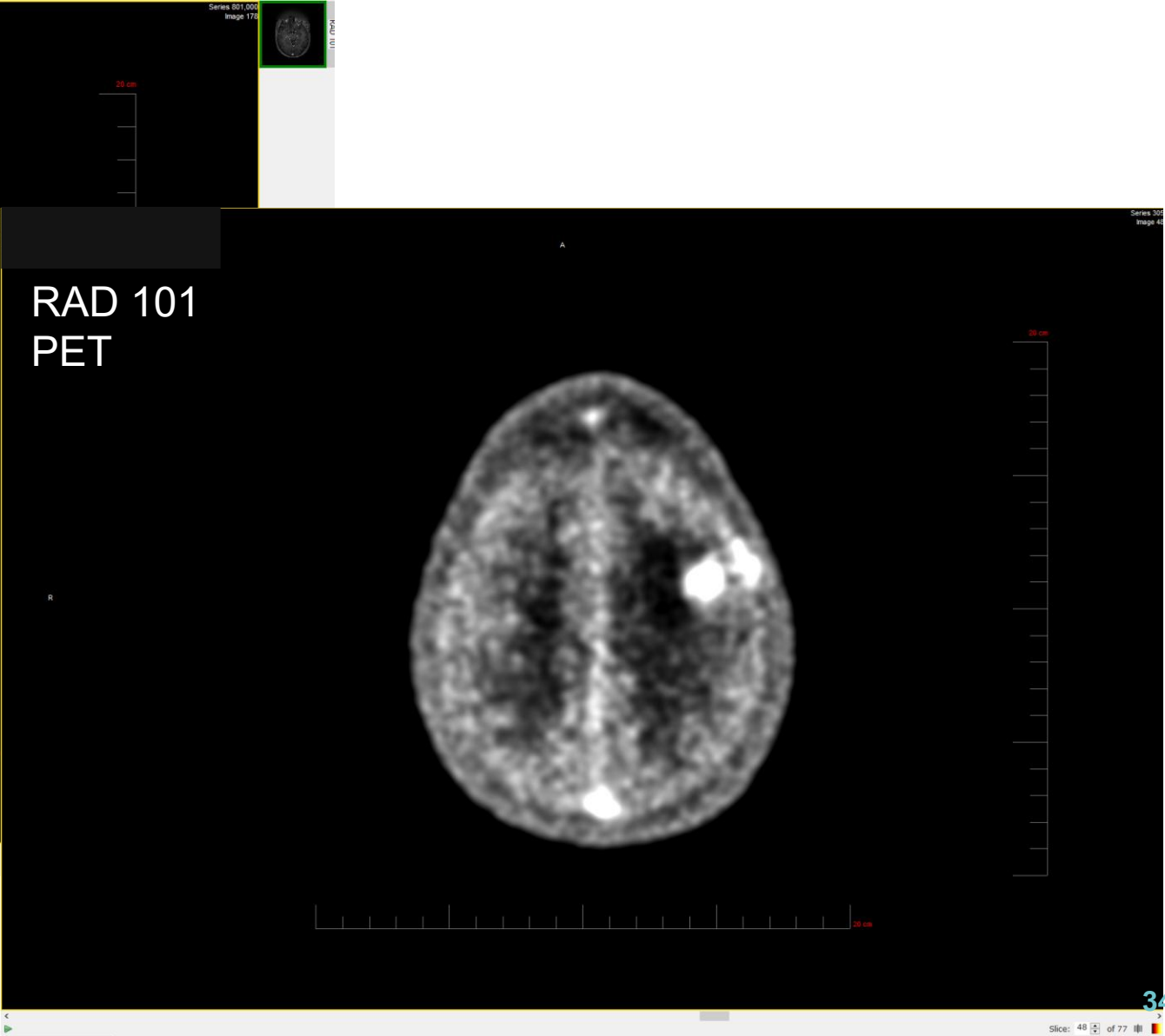
Subject #1 scan dates: 1/25; 9/25;10/25;11/25

Concordance between MRI (contrast uptake plus perilesional hypodensity) & PET (tracer uptake)



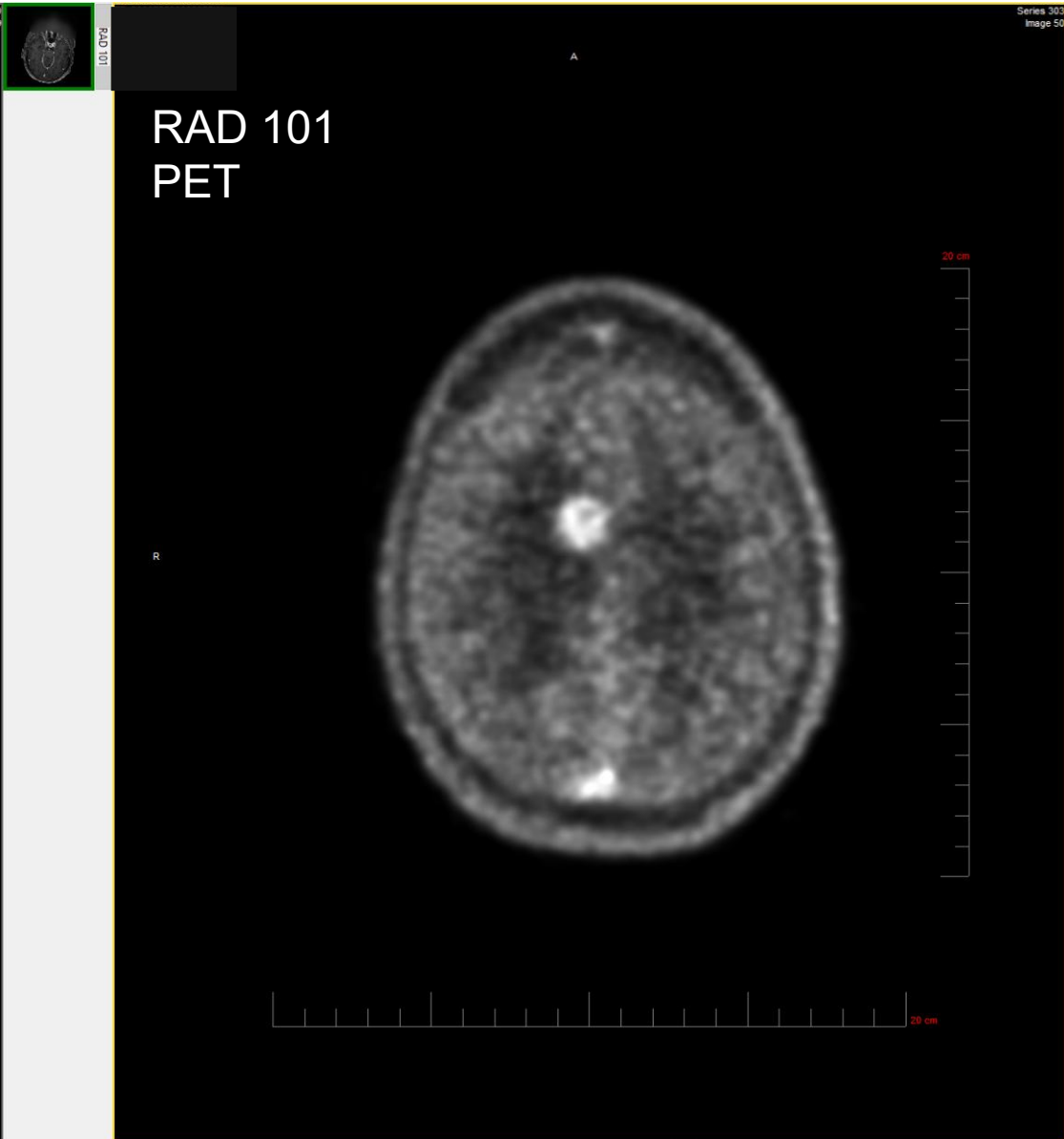
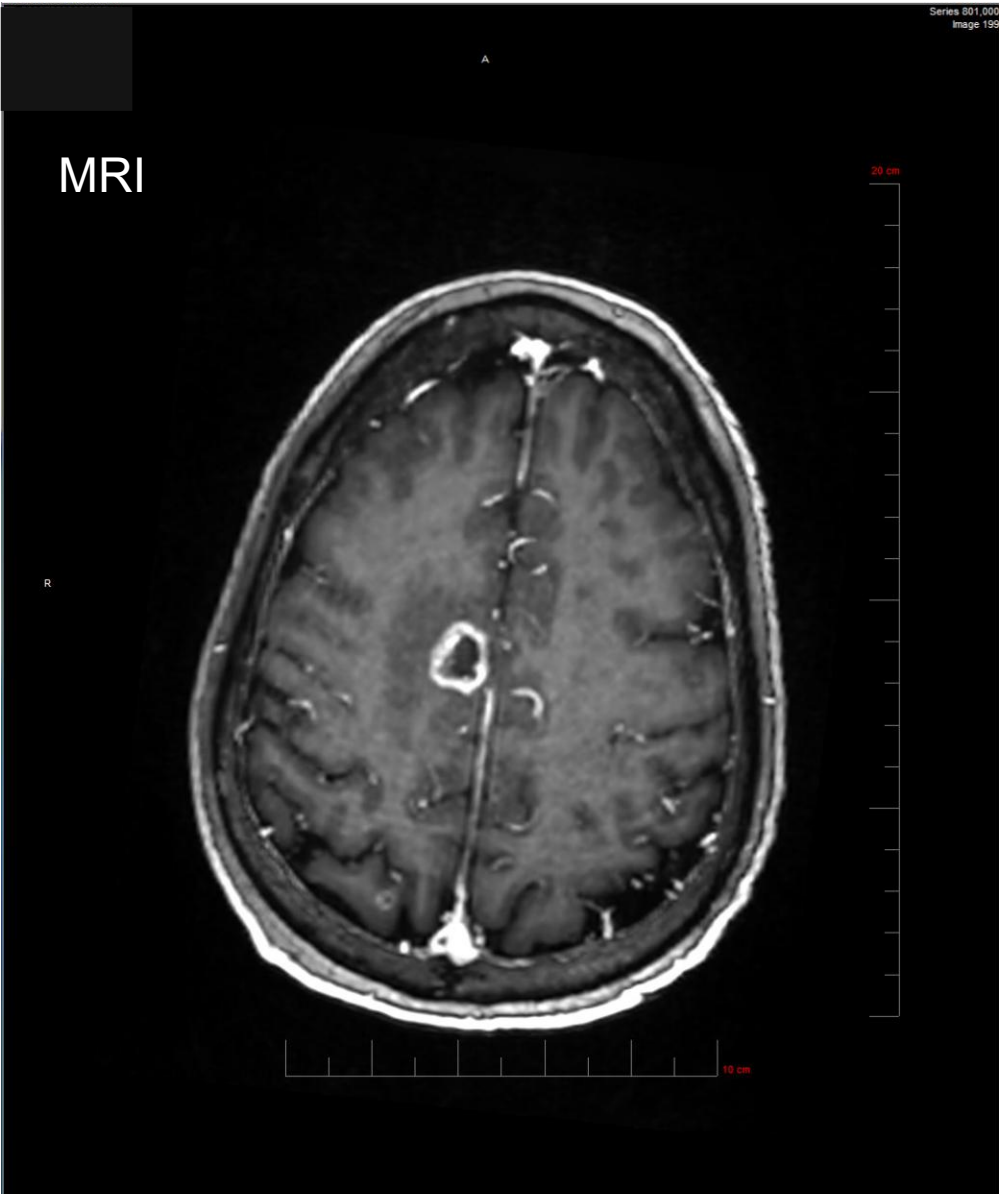
Subject #2

(Concordance between MRI and strong tracer uptake)



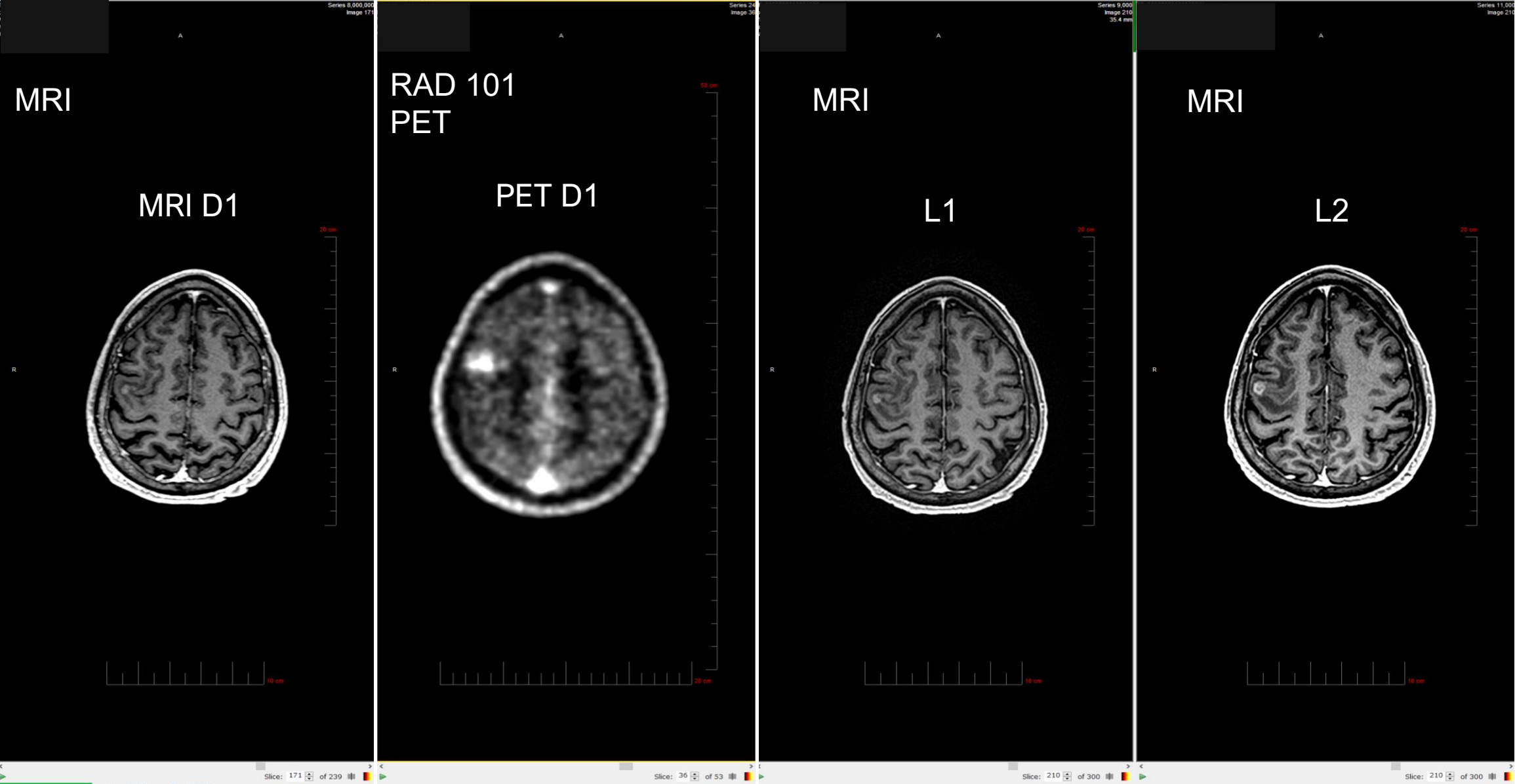
Subject #3 Visit 1 Sep'25 MRI & PET

(Concordance between MRI and strong tracer uptake)



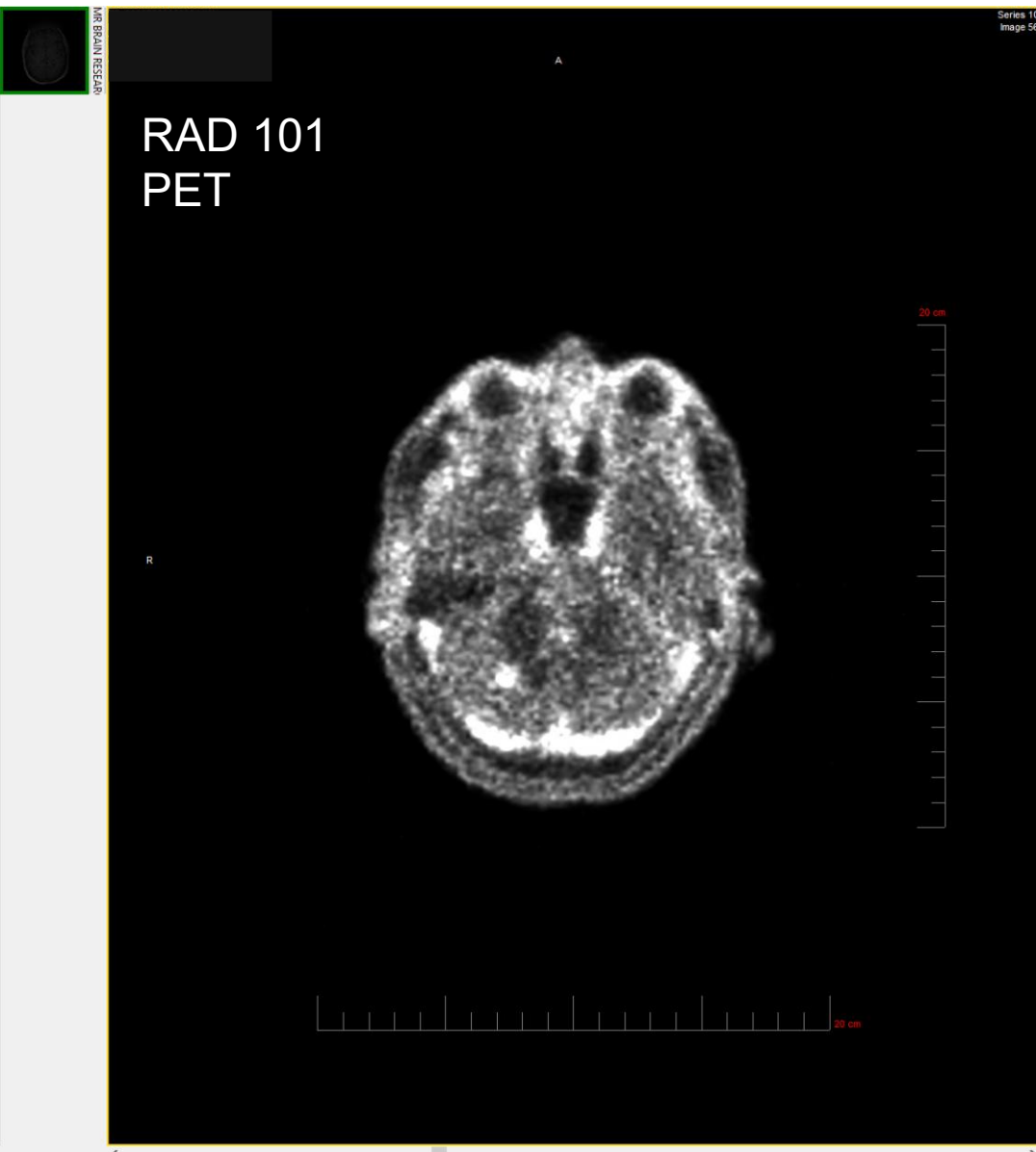
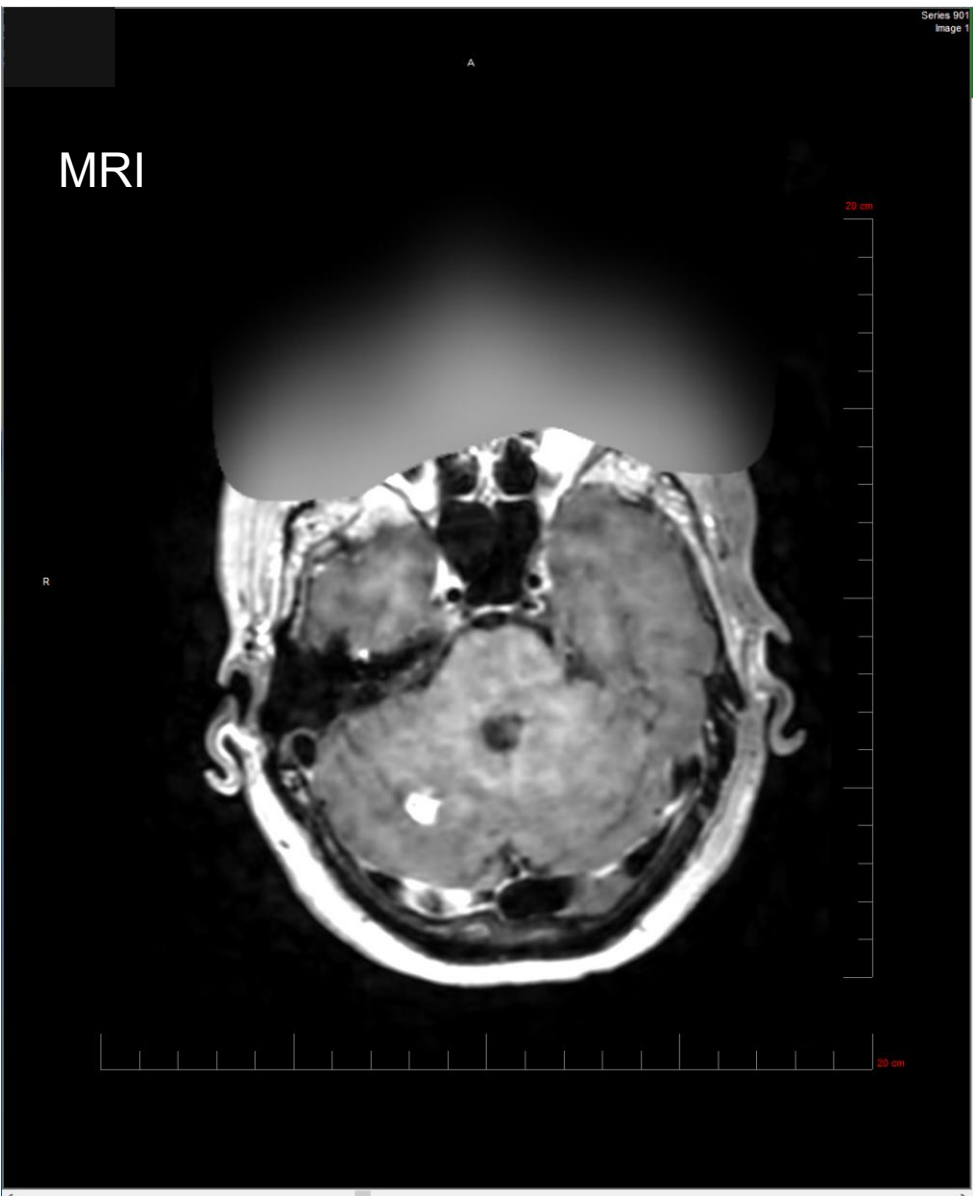
Subject #4 Longitudinal1 (Aug'25) and 2 (Oct'25)

PET positive at Study Date; MRI Progression at Longitudinal 2 (4 months post study)

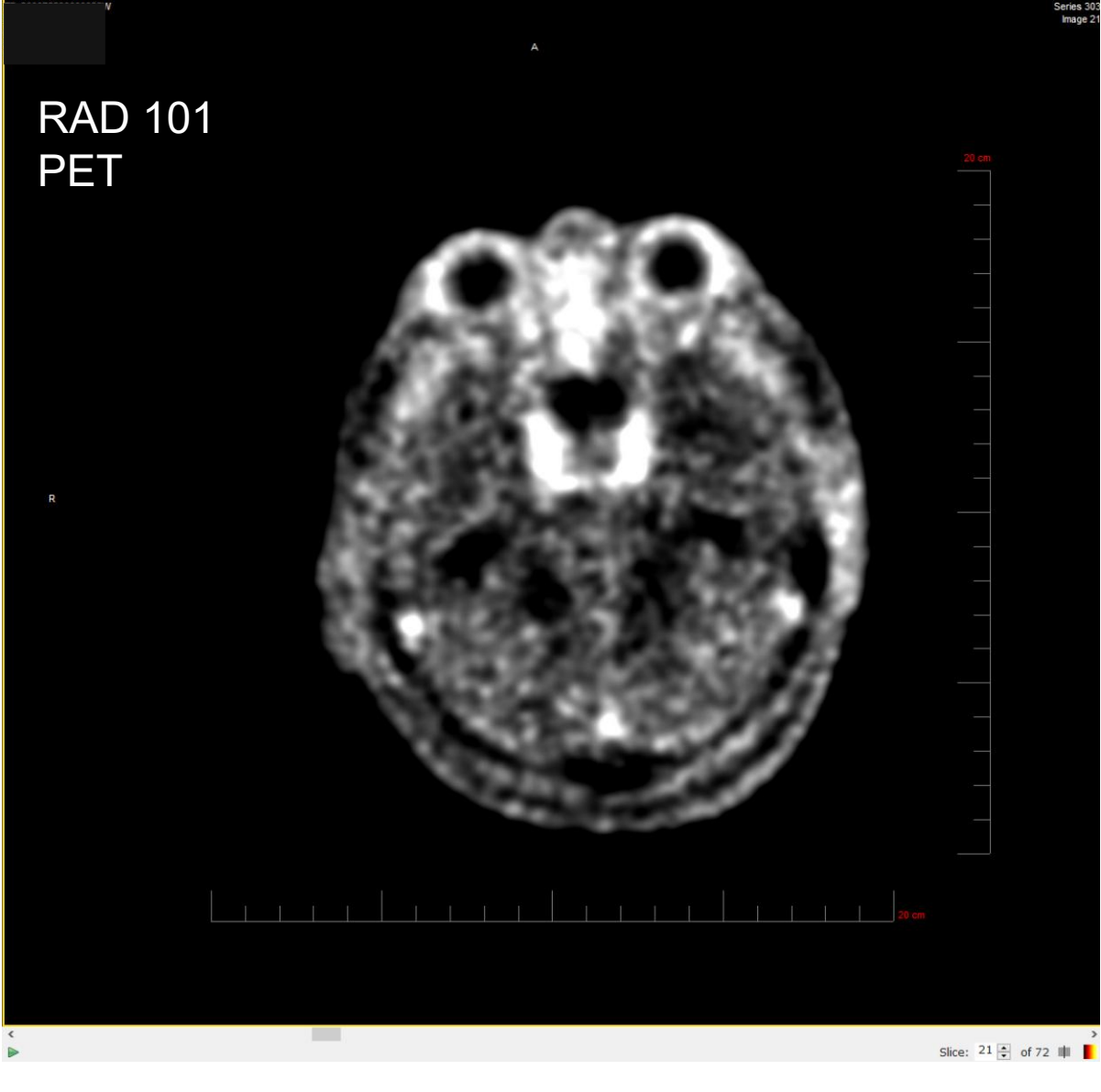
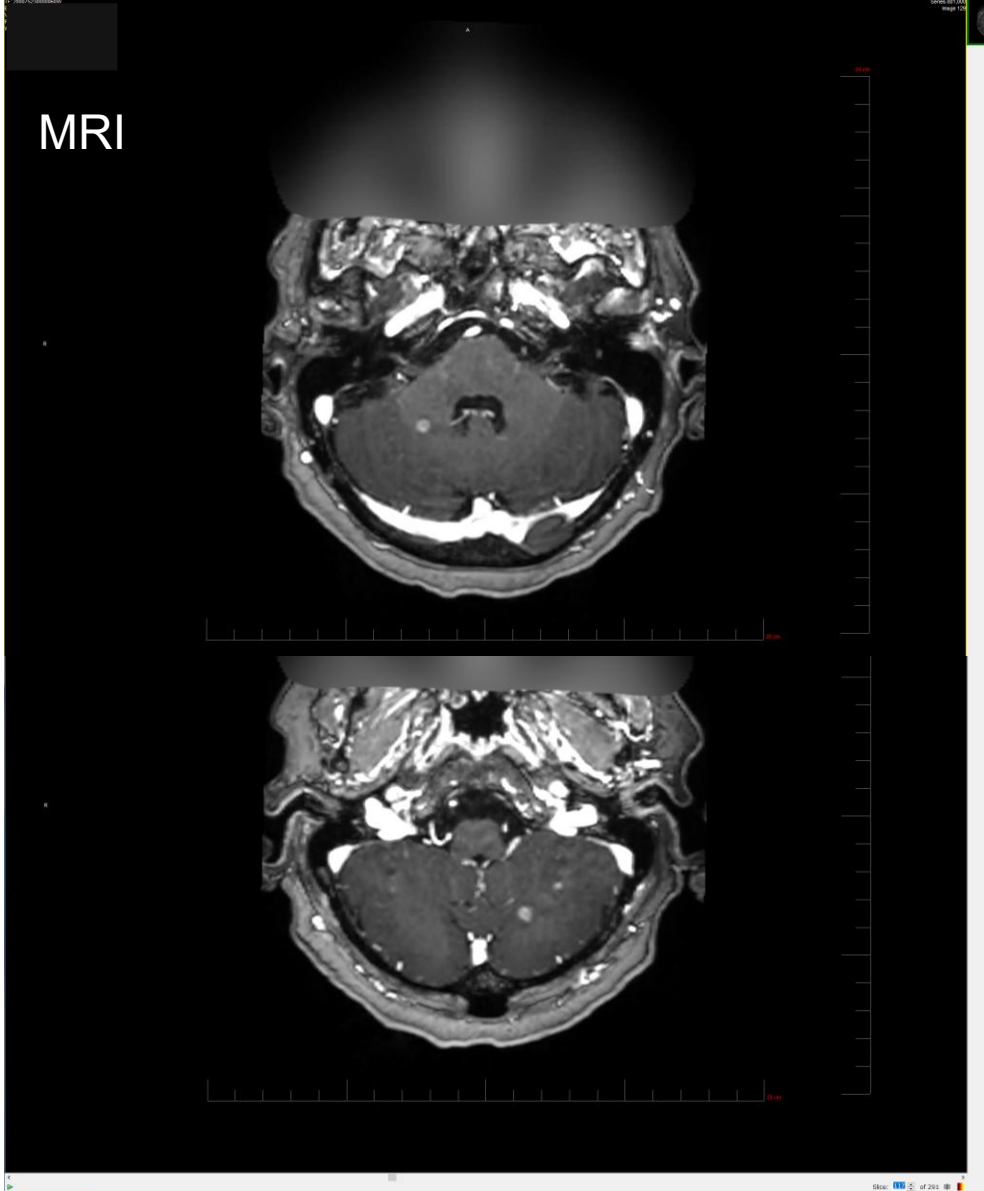


Subject #5 Visit 1 Aug'25 MRI & PET

(Concordance between MRI and strong tracer uptake)



Subject #6 No Active Tumor Detected on PET



Next Steps

Phase IIb

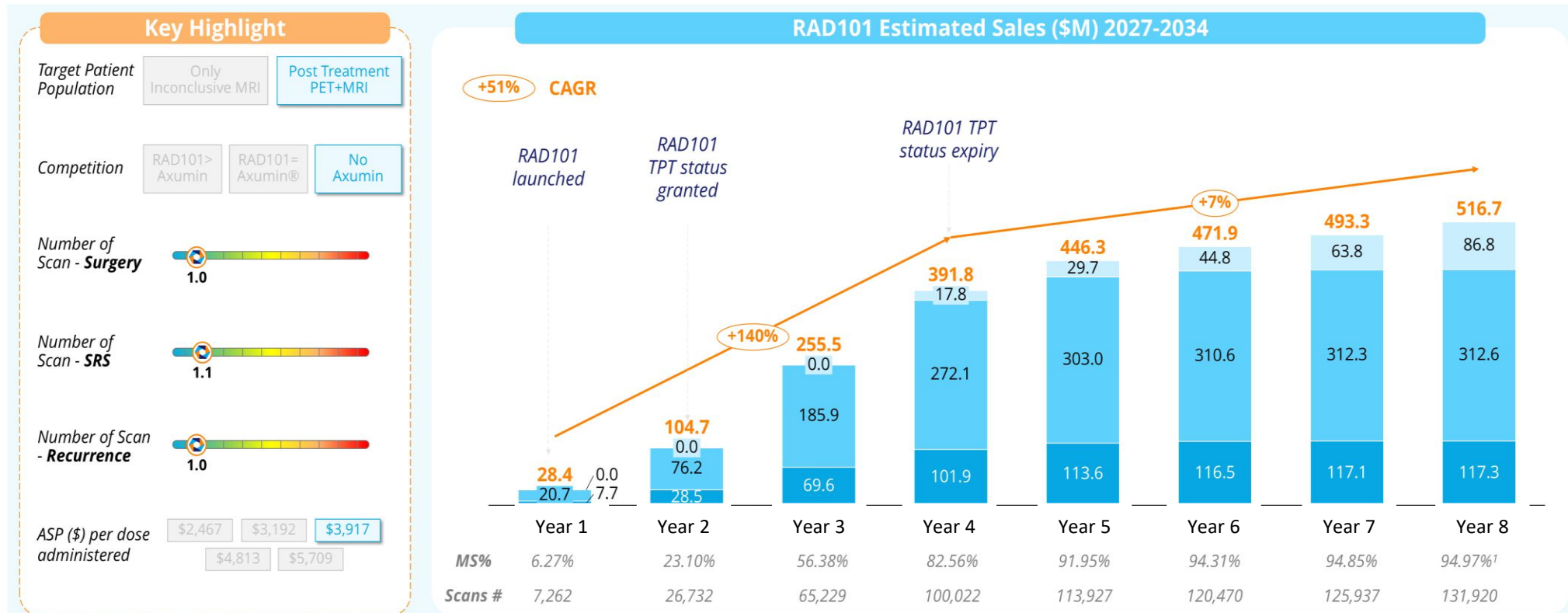
- Trial completion N=30/30 pts by Q1 2026
- Phase 2b primary endpoint readout in the first half of 2026

Phase III

- FDA Meeting to align on Phase III - mid 2026
- Phase III start – Q4 2026

RAD101 COMMERCIAL POTENTIAL: USD\$ >500m yearly sales (USA only)

Third largest imaging molecule after Pilarify (Lantheus) & Illucix (Telix)



Note: ¹In the scenario where RAD101 is the sole product on the market, the peak market share of 100% has been adjusted with a 5% discount factor to account for potential deviations in HCP behavior, such as non-adherence to guidelines or shifts in the SoC. Source: Definitive Healthcare; Primary research; Alira Health analysis.

Legend ■ Assessment of Tx response: Surgery ■ Assessment of Tx response: SRS ■ Assessment of recurrence



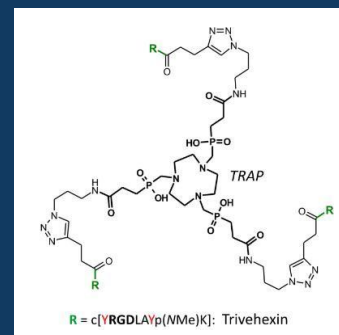
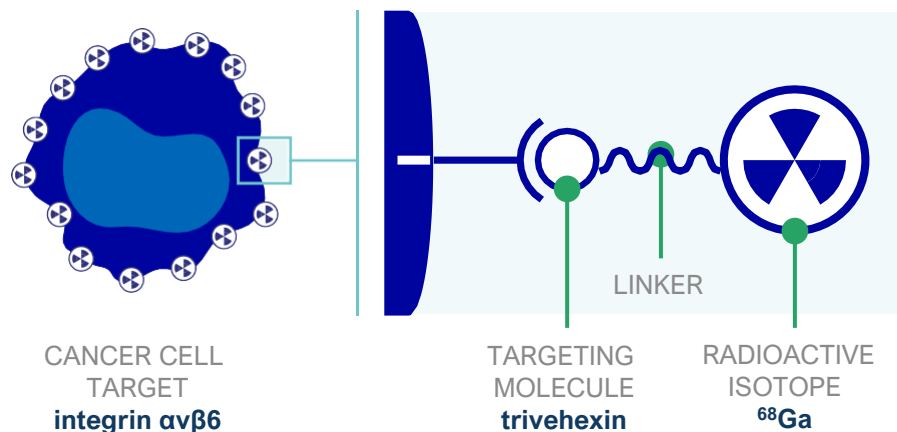
RADIOPHARM THERANOSTICS

Molecule: **68Ga-RAD301**

Targeting MoA: **α V β 6 INTEGRIN**

Imaging for: **PANCREATIC CANCER**

Imaging for Pancreatic Cancer



RAD 301 (Trivehexin) PEPTIDE

- RGD peptide (arginylglycylaspartic acid)
- Integrin $\alpha\beta 6$ receptor antagonist
- Design features include hydrophilicity to reduce non-specific uptake into undesired organs and increase clearance in plasma, trimerization to increase affinity, cyclicity for better selectivity, uptake and tumor retention

INTEGRIN $\alpha\beta 6$

- + Upregulated target often referred to as “cancer integrin” given its role in activation of TGF β ; expression correlates with decreased survival in numerous carcinomas.
- + Pfizer $\alpha\beta 6$ integrin ADC Phase III in NSCLC.




$\alpha\beta 6$ INTEGRIN EXPRESSING TUMORS

- + Pancreatic cancer is first targeted indication (~60% expression).
- + Approx. n=80 subjects already dosed in IIS and under German compassionate use program.
- + Strong peer reviewed presence in several journals and congresses.

80 Subjects Imaged with 68GA-RAD301 to Date

Multi-indication Potential Beyond Pancreatic Cancer

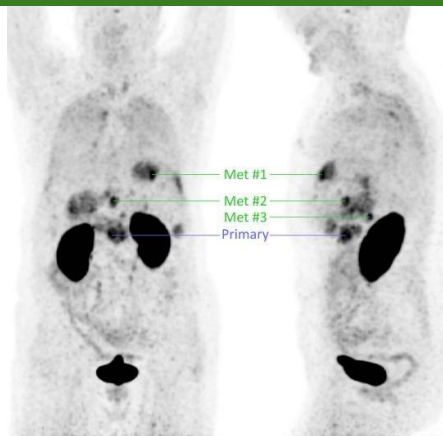
- **44 subjects:** Pancreatic Ductal Adenocarcinoma (PDAC) imaged under 3rd party (Germany) compassionate use*
- **32 subjects:** 12 PDAC, 20 Head & Neck Squamous Cell Carcinoma(HNSCC) imaged in Investigator Initiated Research (IIR)**
- **4 subjects:** single case publications in Non-Small Cell Lung Cancer (NSCLC), Triple Negative Breast Cancer (TNBC), Ovarian, Thyroid Cancer
- **Ongoing Phase I** imaging study in Pancreatic Cancer ongoing at Montefiore, NY and United Theranostic, NJ***
- Phase I is used to confirm Proof-Of Concept in subjects with metastatic pancreatic cancer

3 rd PARTY COMPASSIONATE USE (Germany)*	IIR IN PDAC & HNSCC** + 4 Single Case Publications	PHASE I (USA)***
		
44 pts	32 pts + 4 pts = 36 pts	9 pts
		Ongoing

68Ga-trivehexin PET/MRI Imaging subjects with Pancreatic Tumors

- Detection of $\alpha\beta6$ integrin-expressing tumor lesions in subjects with PDAC
- 66 subjects administered RAD301 (as of 2022)
 - 60 pancreatic cancer and GI tumors
 - 5 with head and neck cancer
 - 1 patient with tumor of unknown origin
- Results indicate that RAD301 can be used to detect and monitor pancreatic cancer
 - Rapid and specific accumulation in many target PDAC primary lesions and metastases
 - Low background accumulation and purely renal elimination

68Ga-TRIVEHEXIN PDAC IMAGING

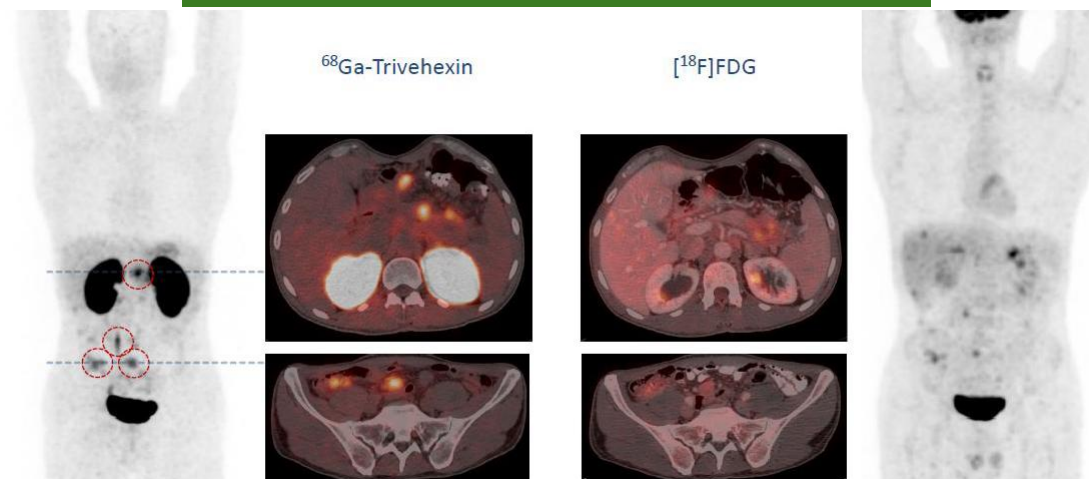


Partnered with TRIMT
Quigley NG Notni J. Eur J Nucl
Med 2021

68Ga-trivehexin PET/CT Imaging vs F18-FDG

- Selective detection of $\alpha\beta6$ integrin-expressing tumor lesions in subjects with PDAC & HNSCC
- 33 subjects administered RAD301
- Results indicate that RAD301 shows incremental value over F18-FDG in PDAC & HNSCC
 - Favorable tumor-to-background contrast vs F18-FDG
 - Sharper images and negligible uptake in the surrounding normal tissue

68Ga-trivehexin PDAC imaging shows superior resolution vs F18-FDG



Partnered with TRIMT
Data presented at World Theragnostic Congress 2022 (Wiesbaden, Germany) & follow up presented at EANM 9/2023 (Vienna)

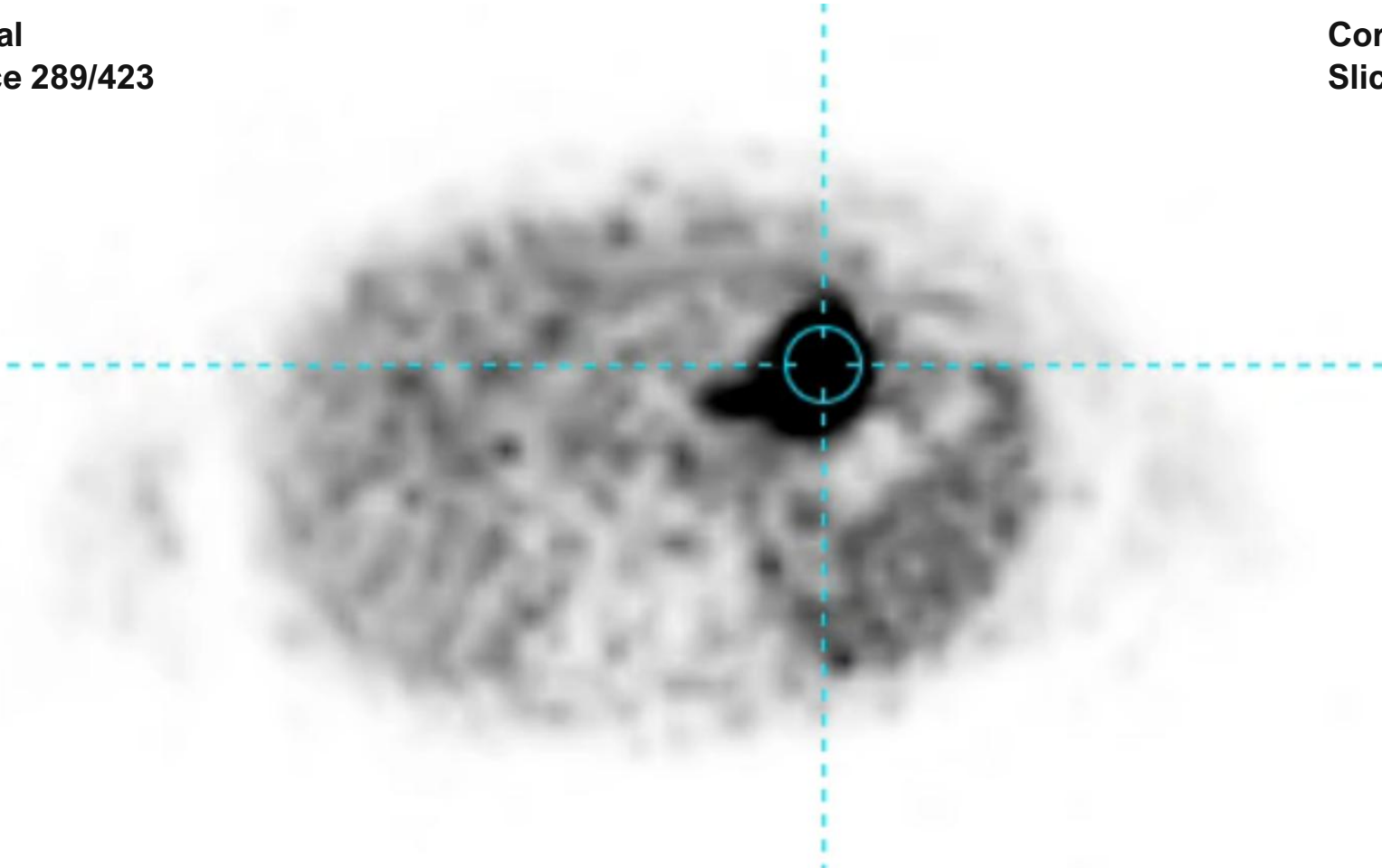
Clinical Data

- Phase 1 company-sponsored study underway in healthy volunteers and pancreatic cancer subjects to characterize biodistribution, image quality and organ/tumor dosimetry
- Preliminary results from n=3 subjects in RADs ongoing study thus far suggest high sensitivity for detection and monitoring of primary tumors and metastatic lesions as small as <1cm
- 8 subjects dosed as 12/17/2025

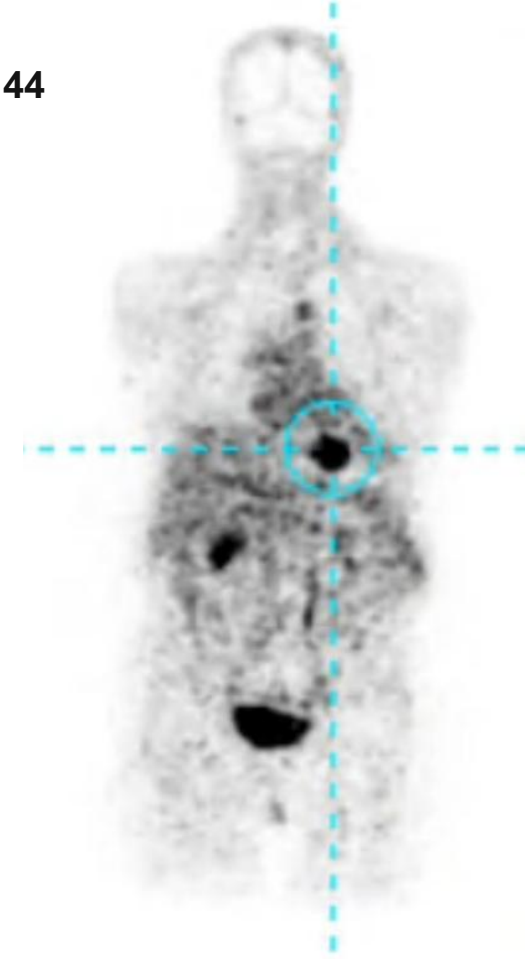
PET/CT Scan | Patient 09

Pancreatic Cancer patient with large pancreatic mass visible in PET

Axial
Slice 289/423



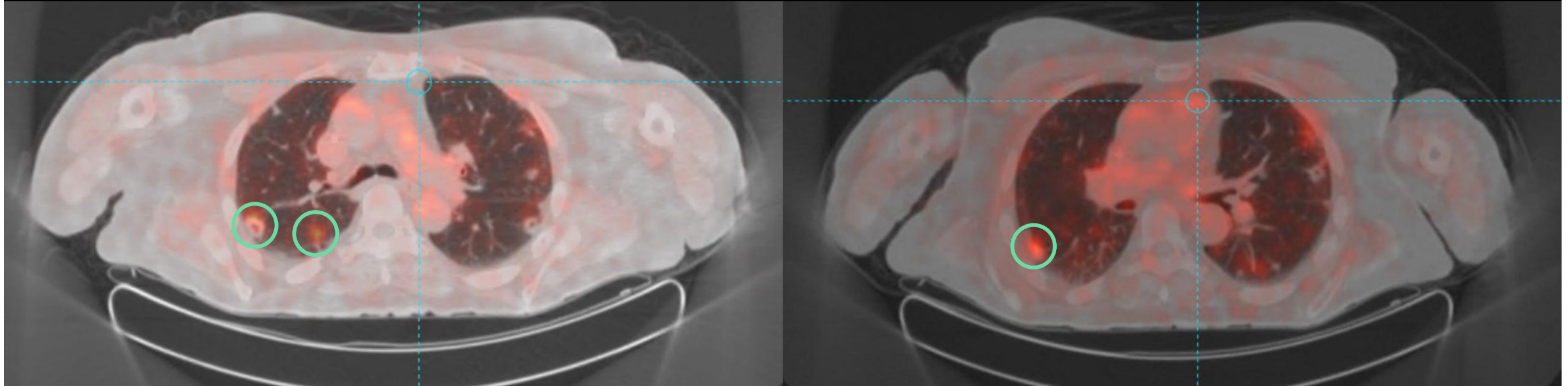
Coronal
Slice 82/144



PET/CT Scan | Patient 15

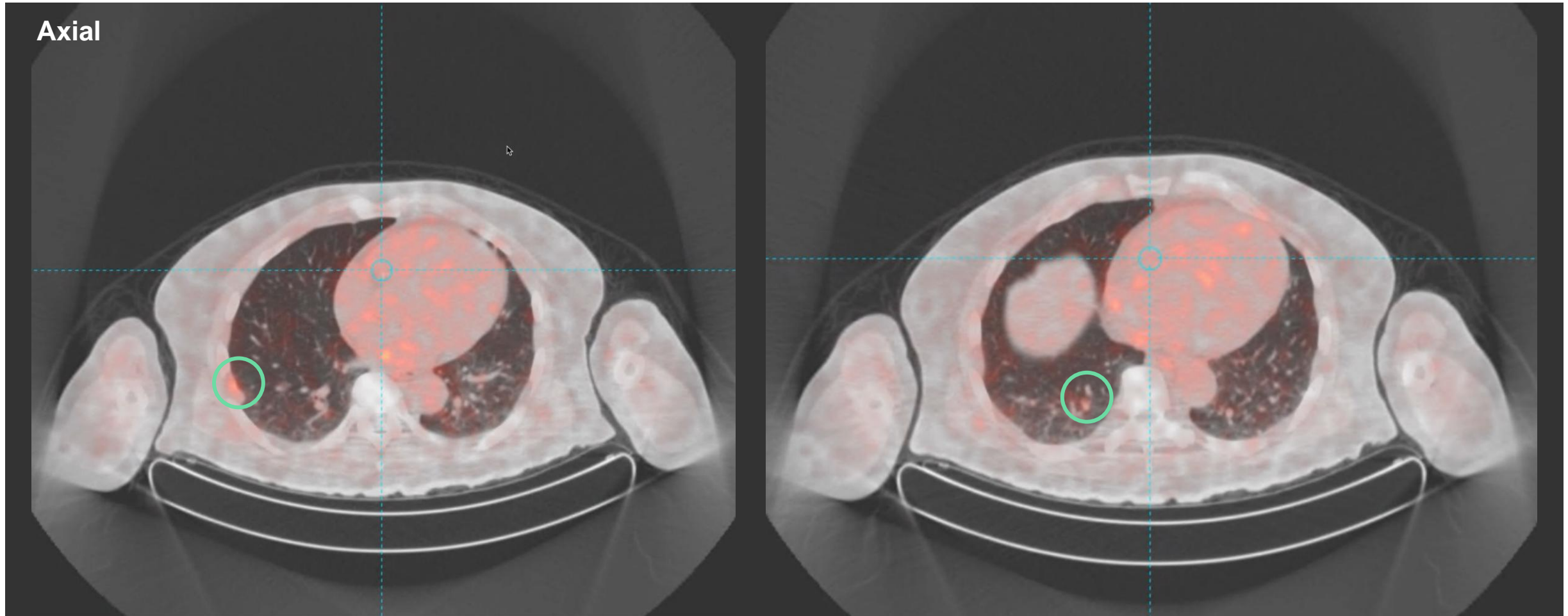
Pancreatic Cancer patient with multiple bilateral metastatic pulmonary nodules ranging in size from 1.3 to 2.2. cm

Axial



PET/CT Scan | Patient 16

Pancreatic cancer patient with multiple metastatic lung nodules <1cm





RADIOPHARM THERANOSTICS

Thank You

www.radiopharmtheranostics.com



Appendix

Chairman and Board



Paul Hopper
Executive Chairman

- Founder of Radiopharm Theranostics Ltd.
- 25 years experience as a life-sciences entrepreneur
- Founder, Chairman, non-executive director or CEO of more than fifteen companies in the US, Australia and Asia
- Previous and current Boards include Imugene, Chimeric Therapeutics, Viralytics, Prescient Therapeutics and Polynoma



Ian Turner



Hester Larkin



Noel Donnelly



Bruce
Goodwin



Riccardo
Canevari





Isotope Selection and Supply Chain

Beta-Emitters: Best proven therapeutic index, secure and reliable sourcing

177-Lutetium






Beta Particles

-  Most used therapeutic isotope
-  Well proven therapeutic index
-  FDA approved for solid tumors
-  Long half-life allows for global distribution

161-Terbium



Beta & Auger Particles

-  Innovative dual atomic particle functionality combining the benefits of Beta cross-fire effect and Auger short-distance high-energy (similar to alpha emission)
-  Potential efficacy in both solid tumors & micrometastases
-  Long half-life allows for global distribution

Tumor Uptake | Up to 3 Gy at 60 mCi

PATIENT #4					Absorbed dose at 60 mCi
Cycle	Lesion	Volume (ml) ²	D1 SUV _{max}	SUV T:BR*	Dose (Gy), with PVC ^{1,2}
C1	Primary	113	2.1	5.1	0.33
C1	Lymph node axillary left (ROI-3)	27	2.7	6.6	0.61
C1	Lymph node supraclavicular left (ROI-4)	27	3	7.3	0.7
C1	Lymph node supraclavicular right (Level V) (ROI-6)	37	1.8	4.3	0.35
C1	Liver Segment VI (ROI-9)	47	6.3	15.4	3.0

PATIENT #5					Absorbed dose at 60 mCi
Cycle	Lesion	Volume (ml) ²	D1 SUV _{max}	SUV T:BR*	Dose (Gy), with PVC ^{1,2}
C1	ROI-4	8.3	2.5	4.7	0.5

PATIENT #6					Absorbed dose at 60 mCi
Cycle	Lesion	Volume (ml) ²	D1 SUV _{max}	SUV T:BR*	Dose (Gy) C1 with PVC ^{1,2}
IM	ROI-7 (Spleen)	16.28	15.2	20.7	2.8

Patients	Average Absorbed dose at 60 mCi
#4	1.0
#5	0.5
#6	2.8
	1.43

Patient 003-009 is not DLT-evaluable (consent withdrawal due to personal reasons)

¹Partial Volume Correction applied.

²Density of lesion: soft tissue = 1.0 g/mL. Bone = 1.3 g/mL

³Lesions were contours based on thresholding (40%) method and volume was averaged over all timepoints

*BR = background – shoulder and proximal thigh. T:BR = lesion SUV_{max}:BR SUV_{mean}

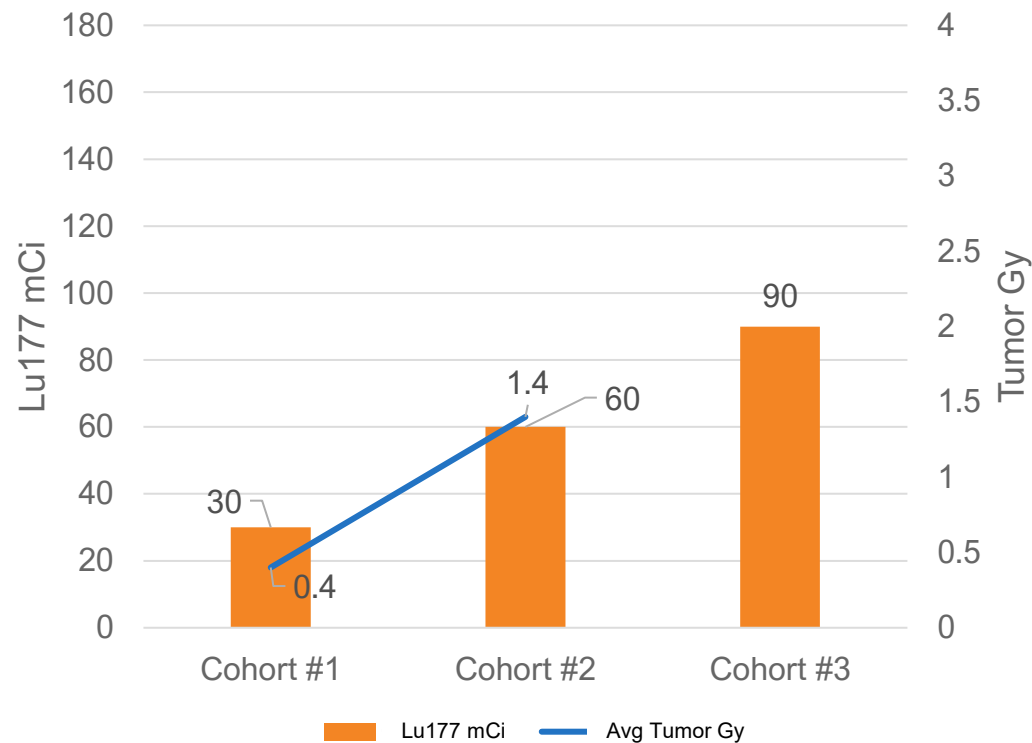
Tumor Uptake | Significant increase at DL2 vs DL1

COHORT#1

Average Absorbed dose at 30 mCi	
Patients	Dose (Gy), with PVC ¹
#1	0.56
#2	0.45
#3	0.21
	0.41

COHORT#2

Average Absorbed dose at 60 mCi	
Patients	Dose (Gy), with PVC ¹
#4	1.0
#5	0.5
#6	2.8
	1.43



RADIOPHARM + MD ANDERSON JOINT VENTURE CREATED IN 2022

Mandate: Develop novel radiopharmaceutical therapies



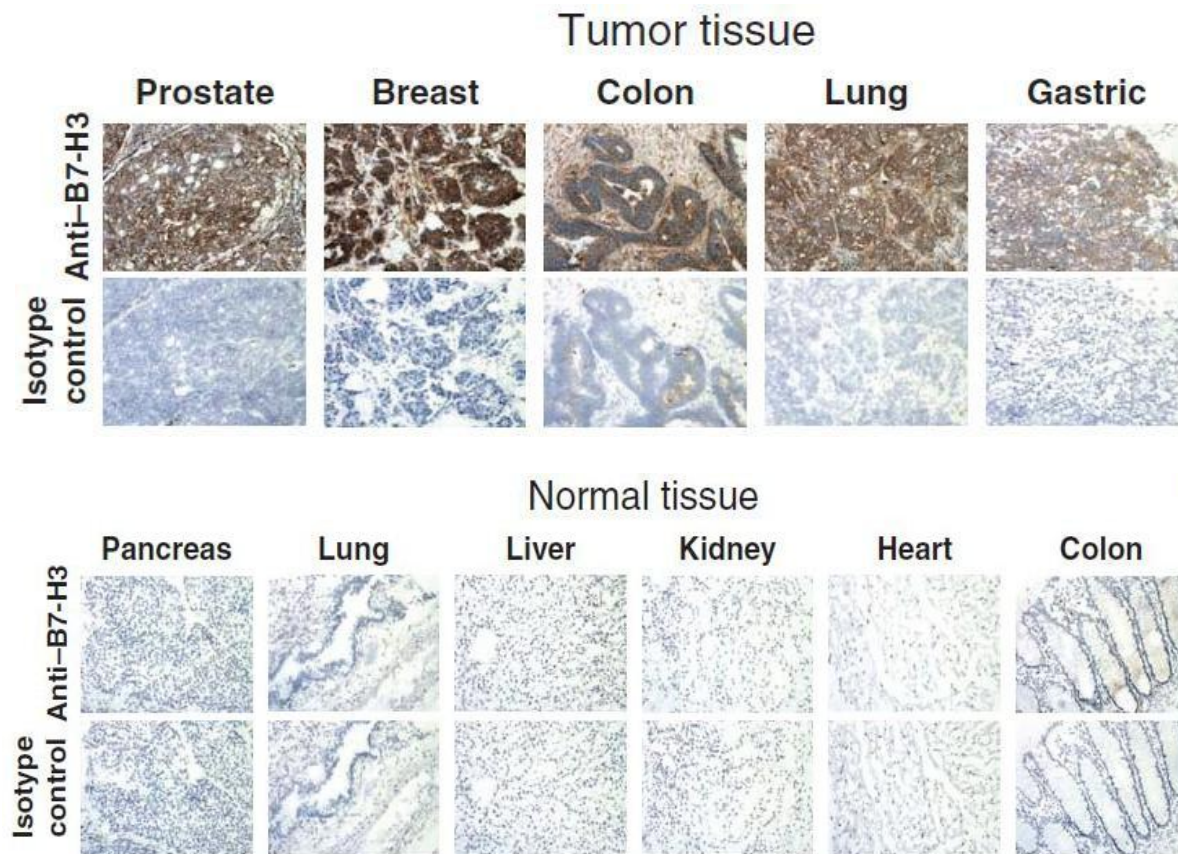
Main Achievements

	2023	2024	2025
B7H3 mAb (RV01)	Double Fc-mutation introduced	IND enabling studies completed	IND clearance received for B7H3 & First patient dose expected
Undisclosed (RV02)	Protein-based vector selected	Preclinical studies	Final candidate selection ongoing
Undisclosed (RV03)	Protein-based vector selected	Preclinical studies	Final candidate selection ongoing
Undisclosed (RV04)	Peptide-based vector selected	Peptide screening ongoing	Preclinical studies planned

Joint Venture share distribution

	2022	2023	2024	2026
RAD	51%	51%	75%	87.5%
MD Anderson Cancer Center	49%	49%	25%	12.5%

B7H3 - Highly Attractive Pan-Tumor Target

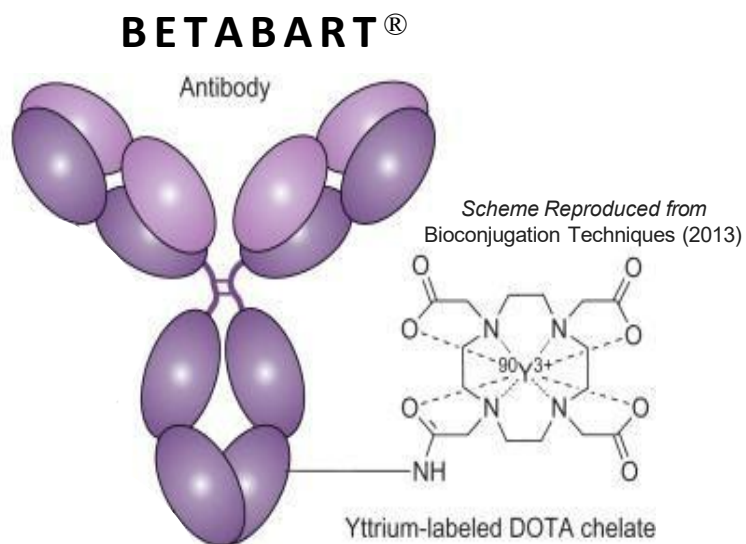


High B7-H3 Expression Levels in Solid Tumors

Potential Indications	B7-H3 Positive*		2+ or Above	
Head and Neck Cancer	19/19	100%	19/19	100%
Kidney Cancer	77/78	99%	75/78	96%
Glioblastoma	65/66	98%	63/66	95%
Thyroid Cancer	34/35	97%	33/35	94%
Mesothelioma	41/44	93%	39/44	89%
Melanoma	132/146	90%	94/146	64%
Prostate Cancer	88/99	89%	51/99	52%
Pancreas Cancer	69/78	88%	45/78	58%
Bladder Cancer	134/156	86%	123/156	79%
Lung Cancer	324/379	85%	300/379	79%
Breast Cancer	189/249	76%	156/249	63%
Ovarian Cancer	59/79	75%	36/79	46%

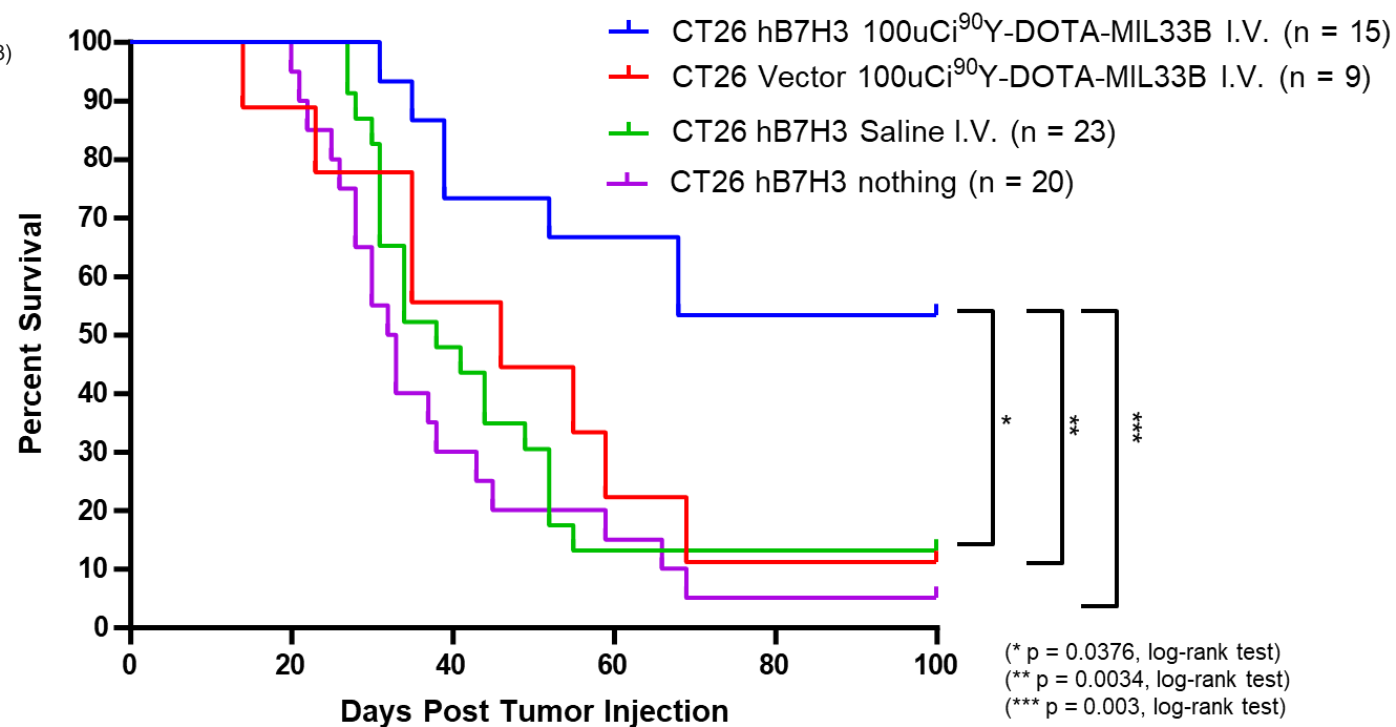
*B7-H3 positivity reflects any grade staining (1-3+) via FFPE tumor microarray (cytoplasmic, membrane, and vasculature staining); B7-H3 is expressed on tumor as well as tumor vasculature.

Betabart: Fc-mutated mAb With Strong Preclinical Efficacy



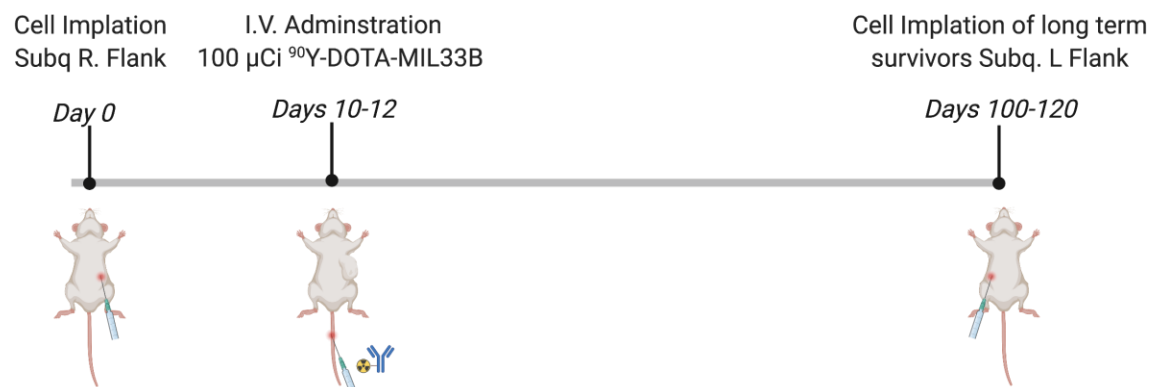
- + Reduced affinity for FcRn
Faster hepatic excretion (no re-circulation)
- + Reduced affinity for FcγR
Reduced bone marrow affinity

56% SURVIVAL WITH SINGLE INJECTION AFTER 100 DAYS

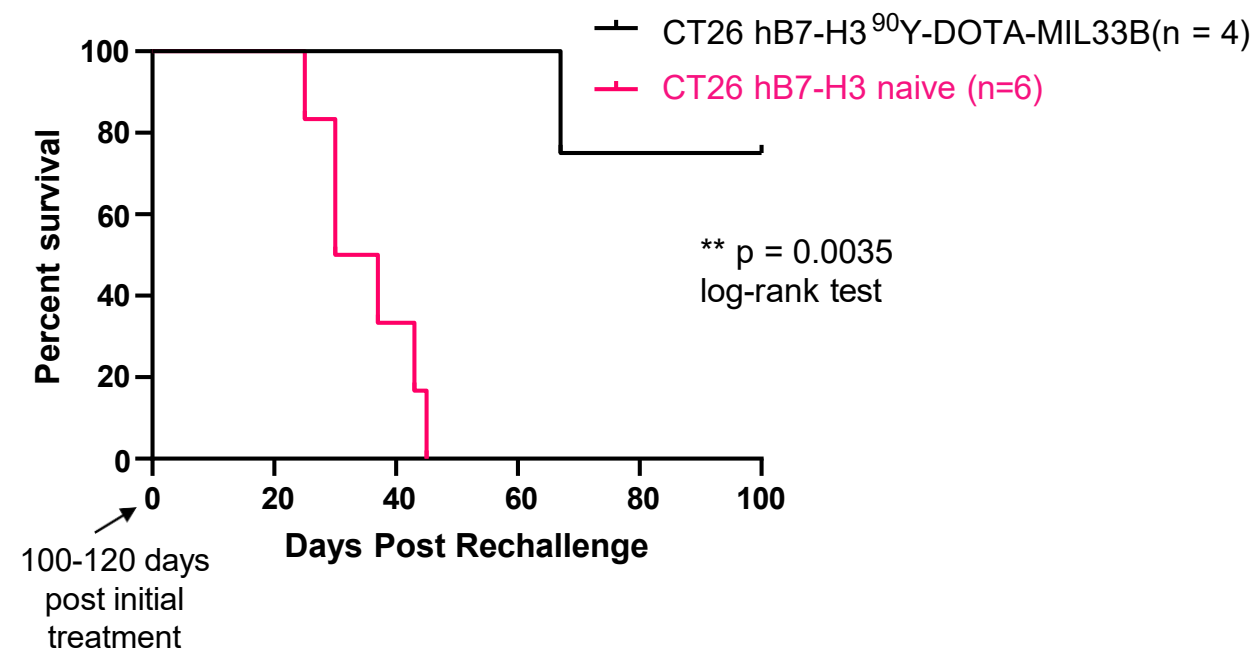


Preclinical Efficacy – Immunological Memory

LONG-TERM SURVIVORS WITH ^{90}Y -DOTA-MIL33B RECHALLENGED WITH CT26 TUMOR CELLS

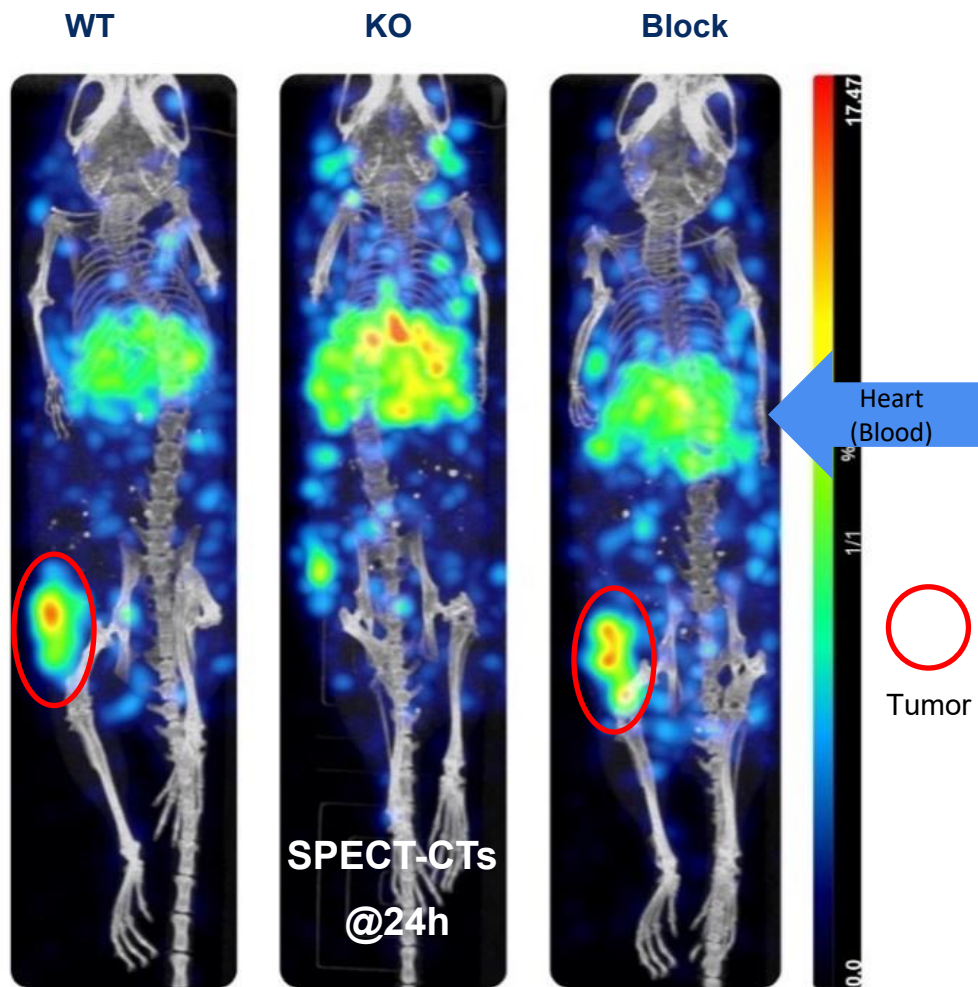


RECHALLENGE SURVIVAL: PRE-TREATED SURVIVORS VERSUS TREATMENT-NAIVE MICE



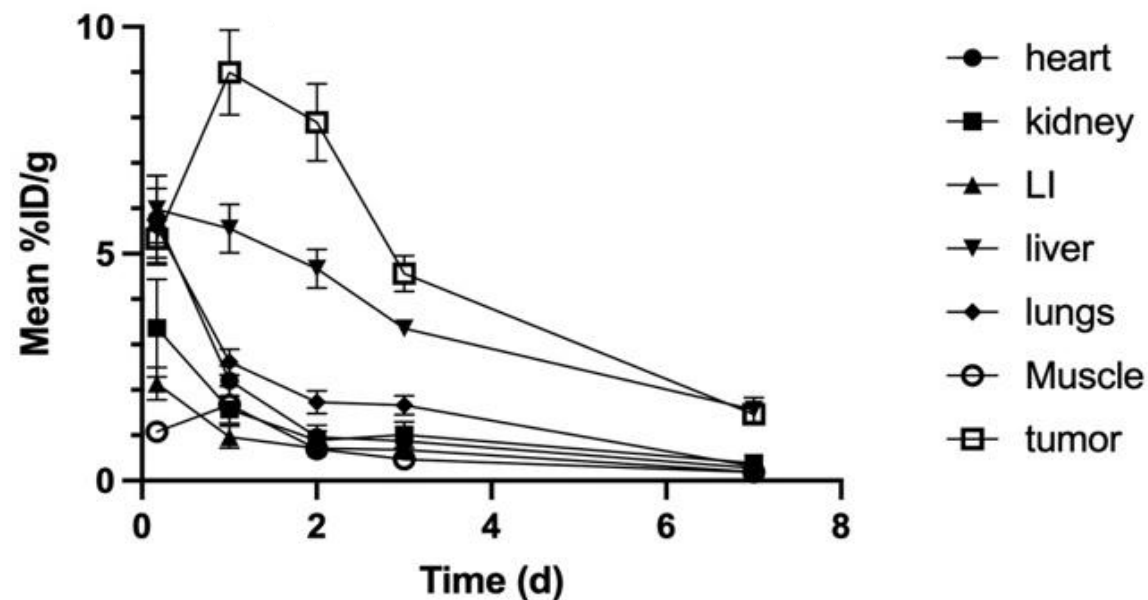
In-Vivo Biodistribution of ¹⁷⁷Lu-BetaBart

Specific Tumor Targeting

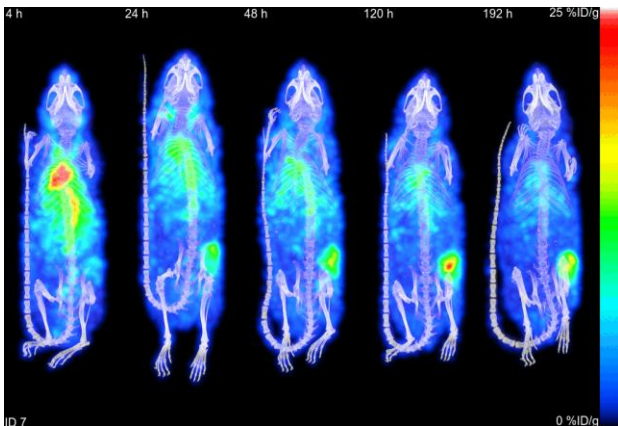


Biodistribution By Design

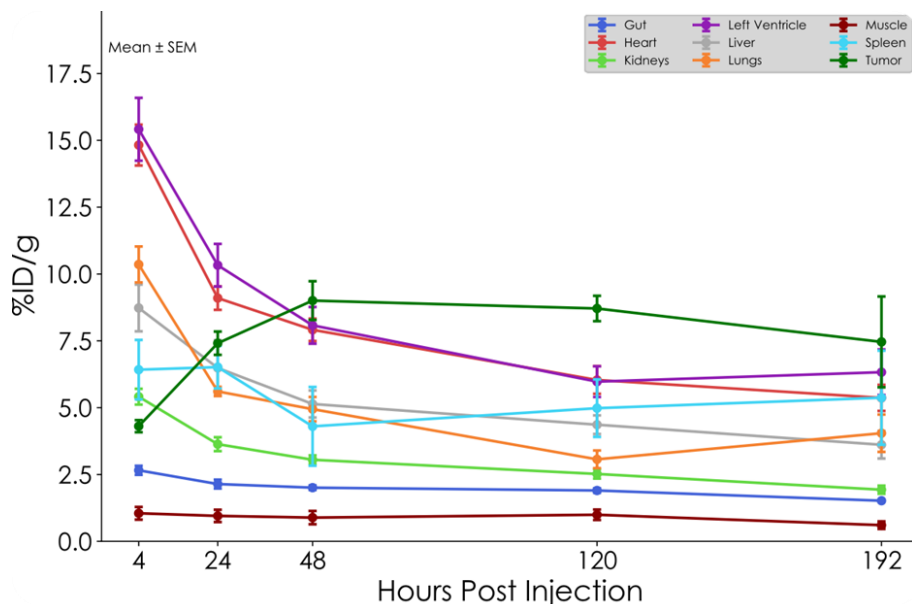
- ✓ High tumor uptake
- ✓ Liver excretion
- ✓ Very low kidney uptake



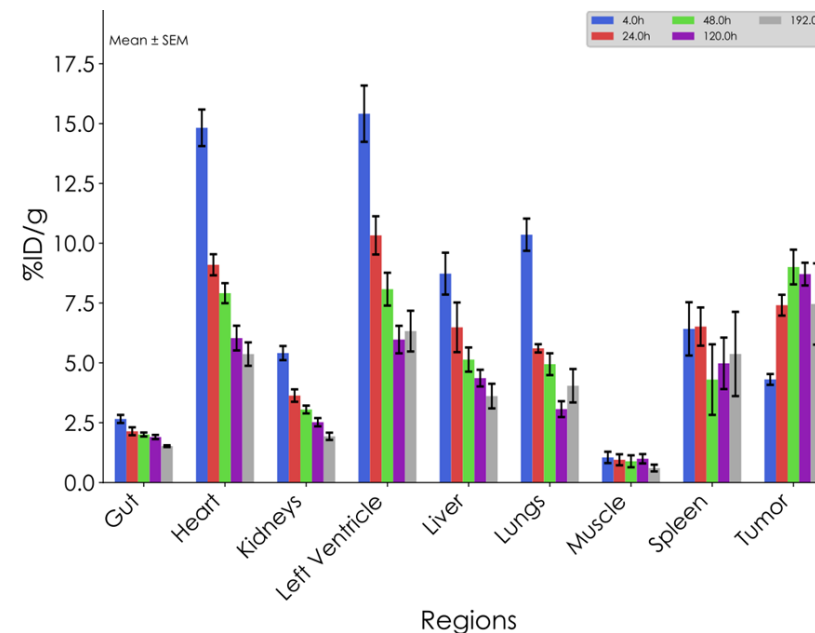
RAD 402 - In Vivo Findings (SPECT BioD in Mice)



- SPECT imaging showed RAD 402 concentration within the heart and lungs at the first time point (4 h) and greatest signal retention in the tumor at the later time points (D2 to D8)
- ¹⁶¹Tb-RAD400 showed good tumor uptake and retention up to 8 d post-injection

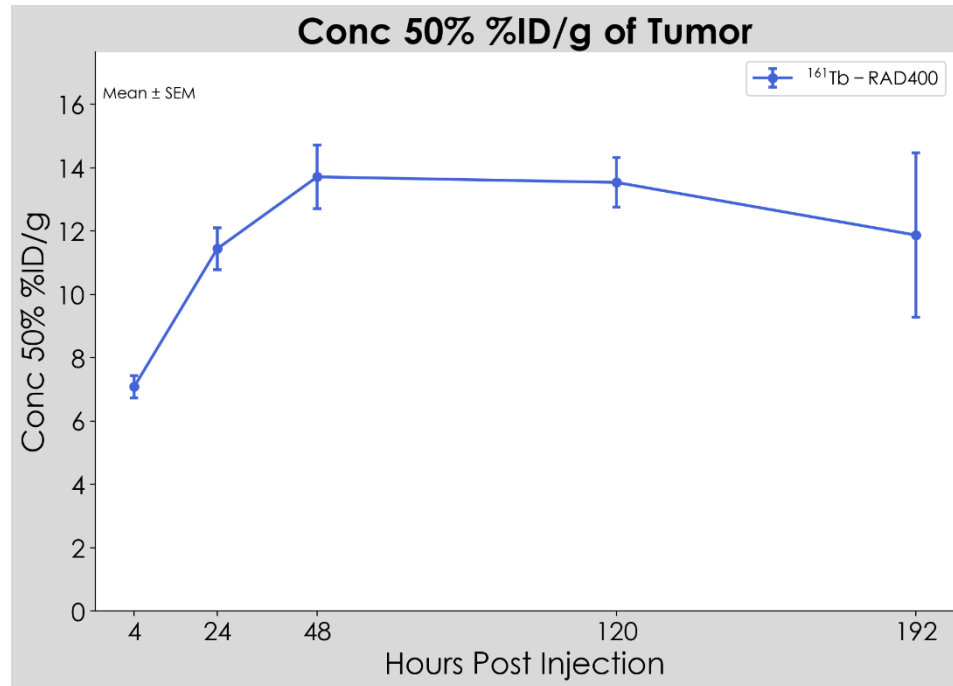


161Tb-RAD400 activity concentration in all ROIs between 4 h and 192 h p.i.

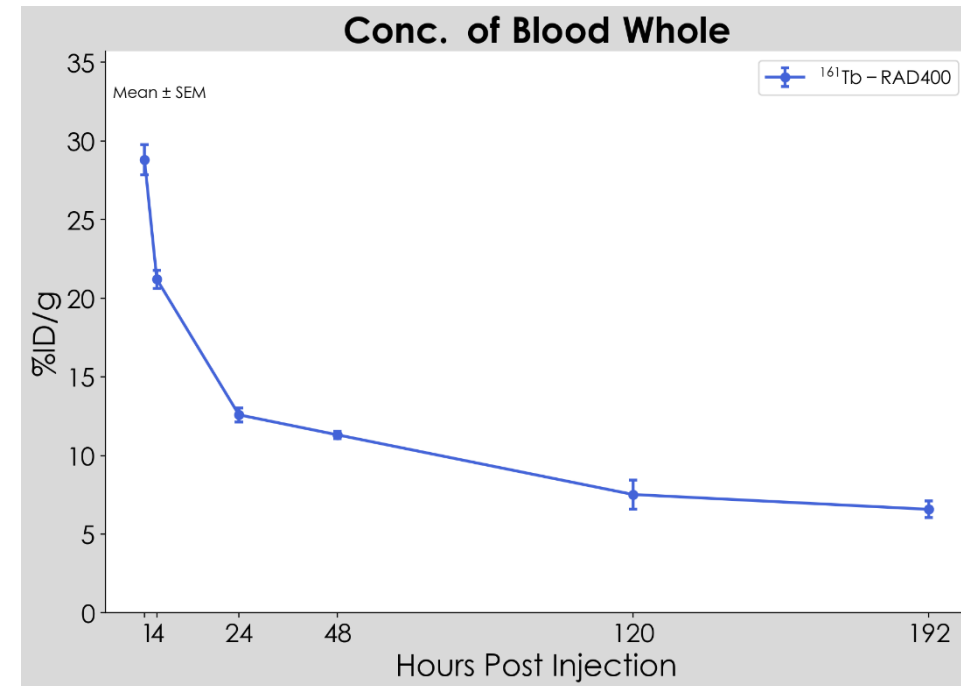


Activity concentration at 4h (blue), 24h (red), 48h (green), 120h (purple), and 192h (grey) p.i.

RAD 402 - In Vivo Findings (SPECT BioD in Mice)



SPECT-derived tumor activity concentration



Activity uptake in blood between 4 h and 192 h p.i.