THERANOSTICS FOR PERSONALIZED MOLECULAR TARGETED THERAPY OF CANCER

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20th International Symposium on Radiopharmaceutical Sciences (ISRS2013) Jeju International Convention Center Jeju Island, May 12-17, 2013

Lecture Outline

- Definition and principles of THERANOSTICS and Personalized Medicine
- THERANOSTIC radionuclides and Ga-68 generator
- Neuroendocrine tumors (NET) as a paradigm
- Diagnosis of NET by PET/CT (clinical applications)
- Dosimetry (organ & tumor dose calculations)
- Therapy of NET (Peptide Receptor Radiotherapy, PRRT)
- Future perspectives
 - new peptides (antagonists, CXCR4, RGD)
 - **PSMA: THERANOSTICS potential for prostate ca.**

Theranostics

- Theranostics is the combination of a Diag*nostic* Tool that helps to define the right *Thera*peutic Tool for a specific disease.
- The term is not specific to radiopharmaceuticals but in fact was first used there (e.g. as "THERAGNOSTICS" by Suresh Srivastava).
- In NM, THERANOSTICS is easy to apply and to understand, because of an easy switch of the radionuclide from Dx to Rx on the same vector.
- The most prominent and oldest application is radioiodine.
- Used by pharma industry at the beginning of the 90's at the same time the concept of Personalized Medicine appeared.

Personalized Medicine

- The right treatment, for the right patient, at the right time, at the right dose.
 - first time », not anymore targeting the "specific disease" but the "specific tumor of a patient".
- The concept of PM has now been extended to Personalized Health Care that includes all steps relevant for the cure of the patient at an individual level from the first sign of disease up to full recovery, including the physicians, the technologies, the drugs and of course all economic aspects, but also extended to the environment, relatives, nurses...

Molecular Nuclear Medicine and THERANOSTICS within MNM are definitely part of Personalized Health Care.

Recent Results in Cancer Research P.M. Schlag · H.-J. Senn Series Editors

Richard P. Baum · Frank Rösch Editors Theranostics, Gallium-68, and Other Radionuclides

A Pathway to Personalized Diagnosis and Treatment

Indexed in PubMed/Medline





2nd World Congress on Ga-68

(Generators and Novel Radiopharmaceuticals) Molecular Imaging (PET/CT), Targeted Radionuclide Therapy, and Dosimetry (SWC-2013): On the Way to Personalized Medicine

28th February - 2nd March, 2013



Organized by : Department of Nuclear Medicine & PET, PGIMER, Chandigarh, India



Targeted Molecular Imaging and Therapy THERANOSTIC PAIRS The Key-Lock Principle



• ¹⁰⁵Rh, ⁶⁷Cu, ^{186,188}Re

Courtesy Helmut Mäcke (modified)





Paul Ehrlich – Side Chain Theory Amboceptors and formation of antitoxins

Corpora non agunt nisi fixata – "Zauberkugeln" (Magic Bullits) Frankfurt/Main, Germany (1899) Nobel Price 1908 – 105 years ago



RADIOLABELED OCTREOTIDE

From bench to bedside: "Proof of principle"



Courtesy Marion de Jong, Rotterdam

From Trial and Error Medicine to Personalized Medicine



New paradigm: personalized medicine



Breaking the cycle of trial and error medicine

Targeted radionuclide therapy has unique promise for personalized treatment of cancer, because both the targeting vehicle and the radionuclide can be tailored to the individual patient.

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Peptides and Receptors in Image-Guided Therapy: Theranostics for Neuroendocrine Neoplasms

Richard P. Baum, MD, Harshad R. Kulkarni, MD, and Cecilia Carreras, MD



Selected Theragnostic Radionuclides

Radionuclide	T1/2 (d)	Principal γ energy for imaging, KeV (%)	Therapeutic particle(s) (Avg. Energy, KeV, % abundance)	
Scandium-47	3.35	159 (68)	β- (162)	
Copper-67	2.58	185 (49)	β- (141)	
Gallium-67	3.26	93, 184, 296 (40, 24, 22)	15 Auger, 0.04-9.5 KeV, 572% 10 C.E., 82-291 KeV, 30%	
Indium-111	2.80	171, 245 (91, 94)	6 Auger, 0.13-25.6 KeV, 407% 12 C.E., 144-245 KeV, 21%)	
Tin-117m	14.0	159 (86)	8 C.E. (141 KeV avg., 114%)	
Samarium-153	1.94	103 (30)	β- (280)	
Bismuth-213	46 min	441 (26)	β- (425); α (98%, from TI-209 daughter, 2% from Bi-213)	
Actinium-225	10.0	99, 150, 187 (93, 73, 49)	α (7030, 93%)	
lodine-123	13.3h	159 (83)	12 Auger, 23-30.4 KeV,1371% 7 C.E, 0.014-32 keV, 17%	
Astatine-211	7.2 h	79 (21)	α (5867, 42%)	





Finland 112712 Selected Theragnostic PET/Therapy Radiometal Pairs			
Radionuclide Pair Imaging/Therapeutic	T1/2 (days)	Imaging positron, KeV (%)	Therapeutic particle(s) (Avg. Energy, KeV)
Scandium-44m/ Sc-47	2.4 / 3.35	γ ± 511 (94%)	β- (162)
Copper-64/Copper-67	0.53 / 2.6	γ ± 511 (38 %)	β- (141)
Gallium-68/Gallium-67	68 min / 3.26	γ ± 511 (176 %)	15 Auger, 0.04-9.5 KeV, 572% 10 C.E., 82-291 KeV, 30%
Yttrium-86/Yttrium-90	0.61 / 2.7	γ ± 511 (35 %)	β- (935)
lodine-124/lodine-131	4.2/8.0	<u>+</u> 511 (44%)	β ⁻ (181)





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Incidence of Neuroendocrine Tumors

2.5-5.0 per 100,000 Population



Surveillance, Epidemiology and End Results (SEER), US population 1974-2005

Modlin et al., Lancet Oncol. 2008

Cumulative Survival of Foregut-NET According to TNM Stage and Tumor Grade

NET - low incidence, but high prevalence!



Prevalence of neuroendocrine tumors

(cases living with the disease)

= 400 per 1 Million

~20,000 in Korea

~120,000 in US

~ 2,400,000 worldwide

NET - low incidence, but high prevalence!



 Yao JC, Hassan M, Phan A, et al. One hundred years after 'carcinoid': epidemiology of and prognostic factors for neuroendocrine tumors in 35,825 cases in the United States. J Clin Oncol. 2008;26(18):3063-3072.
 National Cancer Institute. Surveillance, Epidemiology, and End Results (SEER) Stat Fact Sheets. http://seer.cancer.gov/csr/1975_2004/results_merged/topic_prevalence.pdf. Accessed September 30, 2009.

NET are the second most common gastrointestinal cancer!

Neuroendocrine Tumors

Tumor/ Syndrome Symptoms Secretion Malignancy **Product** Rate Nonfunctioning unspecific Chromogranin A. 60-80% PP, a/ß-HCG, Calcitonin flush, diarrhoea, Serotonin 100% Carcinoid bronchial obstruction **Syndrome** Insulinoma hypoglycemia Insulin, Proinsulin 5-10% Gastrin 60-80% Gastrinoma / ZES ulcera, diarrhoea Vasoactive 40-75% VIPoma / VMS watery diarrhoea intestinale peptide negrolytic migratory Glucagon Glucagonoma 50-80% erythema, diabetes Somatostatinoma diabetes Somatostatin 50% steatorrhoea, cholelithiasis **GHRHoma** GHRH 100% acromegaly Cushing's syndrome CRH, ACTH **CRH**, **ACTHoma** 90-100%

~ 50%

~ 50%

Typical Carcinoid Facies (Flush)



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Technical advances in PET/CT scanner design



Advances in CT:

- increased number of axial slices
- faster gantry rotation times
- incorporation of dual Straton x-ray tubes
- very fast scan times for cardiac applications
- improved use of the radiation dose (TCM, AEC)
- high quality, low mAs clinical CT scans



75 ms/slice 0.26 s total 0.98 mSv

1998; 75 min scan ECAT EXACT



Advances in PET:

- new faster scintillators (LSO, LYSO)
- higher spatial resolution detectors
- increased sensitivity from extended AFOV
- overall improved count rate performance
- iterative recon, accurate system model
- motion corrected PET images with gating
- improved SNR from Time-of-Flight (TOF)

Courtesy David Townsend, 2010



2010; 5 min scan mCT



Developed in close collaboration between Radiopharmacy PET/CT Center, Zentralklinik Bad Berka and Institute of Nuclear Chemistry Johannes Gutenberg-Universität, Mainz, Germany Zhernosekov K, Filosofov DV, Baum RP.... Rösch F J Nucl Med 2007 (Oct): 48:1741-48



Simultaneous use of several generators



⁶⁸Ga-elution, <u>purificaton</u> and synthesis module

First clinical studies in 2004, up to now over 7,500 studies done at ZKL Bad Berka.



- Gallium chemistry well studied
- Rapid increase of PET centers without cyclotron
- The same peptide can be labeled with ¹⁷⁷ Lu or ⁹⁰Y for radionuclide therapy



Automated cassette-based synthesis for the daily routine production of radiopharmaceuticals, e.g. Ga-68 HA-DOTA-TATE, PSMA, CPCR4, RGD.....

Selected Compounds and their sst-Receptor Affinity

Compound	sst1	sst2	sst3	sst4	sst5
Somatostatin-28ª	5.2 ± 0.3	2.7 ± 0.3	7.7 ± 0.9	5.6 ± 0.4	4.0 ± 0.3
OC	>10.000	2.0 ± 0.7	187 ± 55	>1000	22 ± 6
I-TOC	>10.000	1.3 ± 0.3	128± 22	867 ± 33	50 ± 12
I-TATE	>1.000	0.5 ± 0.2	187± 38	337 ± 57	50 ± 5.8
Ga-DOTA-OC	>10.000	7.3 ± 1.9	120 ± 45	>1.000	60 ± 14
Ga-DOTA-TOC	>10.000	2.5 ± 0.5	613 ± 140	>1.000	73 ± 21
Ga-DOTA-TATE	>10.000	0.20 ± 0.04	>1.000	300 ± 140	377 ± 18
Y-DOTA-OC	>10.000	20 ± 2	27 ± 8	>10.000	57 ± 22
Y-DOTA-TOC	>10.000	11.0 ± 1.7	389 ± 135	>10.000	114 ± 29
Y-DOTA-TATE	>10.000	1.6 ± 0.4	>1.000	523 ± 239	187 ± 50

Improvement of the sst-receptor affinity



sst-Receptor Affinities DOTATATE versus HA-DOTATATE

	sst ₁	sst ₂	sst ₃	sst ₄	sst ₅	lgP
Ga-DOTATATE	> 10.000	0.67 ± 0.25	> 1000	822 ± 327	> 1000	-3.69
Ga-HA-DOTATATE	> 10.000	0.64 ± 0.23	> 1000	> 1000	59.7 ± 15.1	-3.16
Lu-DOTATATE	> 10.000	1.21 ± 0.27	162 ± 16	> 1000	> 1000	-
Lu-HA-DOTATATE	> 10.000	0.73 ± 0.10	93 ± 0.1	> 1000	147 ± 40	-

Ga-68 Growth

a-68 Growth	150	PUBMED Search "Gallium-68 or
o Bombesin		68Ga or Ga-68 "
• RGD	125	
 Biotin 	123	
 Porphyrin-like 		
o Citrate	100	
 Octreotides 	100	2011
 Affibody 		2011
 P-glycoprotein targeted 	75	90+Hits
 Nanoparticles 	70	
 Neurotensin 		
 Glycopeptide 	50	2010
 Proteins 		$60 \pm Hits$
 Nitroimidazoles (hypoxia) 		00 ± 11115
 Many more 	25	
	0	
		\$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$

YEAR

2012 150+ Hits

Courtesy C. Decristoforo

Gallium-68 has the potential to become the Tc-99m for PET/CT! *

In 2013, we have 8 different Ga-68 labeled radiopharmaceuticals in clinical use!



Ga-68 PET/CT Studies Zentralklinik Bad Berka

Ga-68 SMS Receptor PET – Imaging Technique

Images courtesy Heiner Bihl/Gabriele Pöpperl Klinik für Nuklearmedizin •Katharinen-Hospital, Stuttgart



0:20 p.i. 0:40 p.i. 1:00 p.i. 1:20 p.i. 1:40 p.i. Injected activity: 1.5 MBq/kg (100-150 MBq, 3-4 mCi). Start of acquisition: 60-90 min p.i. (30-180 min) Acquisition parameters: 2 min. per bed position Effective radiation dose: 3 mSv for 150 MBq ⁶⁸Ga-DOTATOC (+CT) (Octreoscan® 12 mSv)

Imaging characteristics: fast kinetics, fast renal clearance, high quality images with very low background images with very low background image analysis: visual lesions (3 to 5 mm) already 30 to 60 min. p.i. Image analysis: visual and quantitative (SUV) evaluation

Dr Michael Hofman

Gatate PET/CT vs ¹¹¹In-Octreotide

Staging



Molecular Imaging of NET by SMS-Receptor-PET/CT

 Whole-body diagnosis ("one-stop shop")
 Detection of unknown primary tumors (CUP)
 Evaluation of receptor status before and after PRRT

ZMA-360369/08-04

Whole-body diagnosis ("one stop shop")



Receptor-PET/CT using Ga-68 DOTA-NOC Primary tumor (ileum), liver, lymph node & bone metastases Eur J Nucl Med Mol Imaging DOI 10.1007/s00259-009-1205-y

Eur J Nucl Med Mol Imaging 2010 Jan;37(1):67-77

ORIGINAL ARTICLE

Detection of unknown primary neuroendocrine tumours (CUP-NET) using ⁶⁸Ga-DOTA-NOC receptor PET/CT

Vikas Prasad · Valentina Ambrosini · Merten Hommann · Dieter Hoersch · Stefano Fanti · Richard P. Baum Results In 35 of 59 patients (59%), ⁶⁸Ga-DOTA-NOC PET/

CT localised the site of the primary: ileum/jejunum (14),

Received: 30 November 2008 / Accepted: 12 June 2009

Conclusion Our data indicate that ⁶⁸Ga-DOTA-NOC PET/ CT is highly superior to ¹¹¹In-OctreoSean (39% detection rate for CUP according to the literature) and can play a major role in the management of patients with CUP-NET.

Dept. of Nuclear Medicine/P.E.T. Center, Zentralklinik Bad Berka



Primary tumor (ileum) and synchronous liver metastasis

378.712-M.G.


Superiority of Ga-68 DOTANOC PET/CT over conventional imaging











Osteoblastic metastasis: **CT** positive

CT normal (3 mm lesion on MRI)

Rare localizations detected by PET/CT



Intramuscular metastasis (psoas muscle)

358.775-P.J.

INTRACARDIAC METASTASIS (SEPTUM)







C.E.

1:0,

In Wahl R. (ed.): Principles and Practice of PET and PET/CT. Lippincott Williams & Wilkins, Philadelphia 2008 (p. 411-437).



Principles and Clinical Indications

PATIENT EVALUATION BEFORE PRRT

Receptor density determined by Ga-68 receptor PET/CT:

semiquantitative measurement by

SUV (Standardized Uptake Values)

More than just looking at images..



Treatment decisions based on Ga-68 SMS receptor PET/CT: Bad Berka scoring system is based on SUVs – not on visual analogue scales as previously derived from OctreoScans

Ga-68 DOTA-NOC receptor PET/CT: SUV of primary tumors and metastases

V. Prasad, R.P. Baum Q J Nucl Med Mol Imaging 2010; 54:61-67

SUV	Mean	Range
in primary tumors		
and metastases		
(n = 1,400 studies)		
Primary tumors	19.2	8.2 – 109
Liver mets	20.9	3.3 - 156
Lymph node mets	9.5	4.2 – 152
Bone mets	13.6	3.0 – 20.4
Brain mets	12.3	4.6 – 17.2
Lung mets	2.3	1.6 – 5.6
Abdominal mets	14.8	5.8 – 34.1

ORIGINAL ARTICLE

Molecular imaging with ⁶⁸Ga-SSTR PET/CT and correlation to immunohistochemistry of somatostatin receptors in neuroendocrine tumours

Daniel Kaemmerer • Luisa Peter • Amelie Lupp • Stefan Schulz • Jörg Sänger • Vikas Prasad • Harshad Kulkarni • Sven-Petter Haugvik • Merten Hommann • Richard Paul Baum



From Molecular Imaging to Therapy

lleum NET, size 4 mm



Ileum NET



IHC Scoring for SSTR1-5

Ga-68 DOTA-SMS PET/CT in 34 histologically documented GEP NET patients

44 surgical specimens generated

Only lesions > 1.5 cm on PET/CT were selected to avoid partial volume effect on the semiquantitative parameters





SSTR1: IRS = 2; HER2 = 1+

SSTR2A: IRS = 9; HER2 = 3+

Lymph Node

Does receptor density, as predicted by PET/CT (SUV, MTV, MTD) correlate with immunohistochemistry scores (HER2 & IRS)?



Digitialised Histopathological Classification Definiens XD Image Analysis - Processing Somatostatin receptor imaging using Ga-68 DOTA-NOC PET/CT

results in accurate estimation of the receptor density.

Image Analysis Results SSTR-2	Correlation	Liver Mets SUVmax PET/CT
N1	Correlation Coefficient	-0,733
	P Value	0.02
N2	Correlation Coefficient	-0.750
	P Value umber of Patients:9	0.0158

Results

The correlation coefficients for SUV max, SUVmean, and MTV ranged from 0.83 to 0.99 (p<0.005).

The tumor SUVmax showed a significant correlation with immunohistopathology scores.

A correlation was also found between SSTR1-5 staining and the corresponding pathology grading.

Ga-68 DOTA-SSTR PET/CT provides in vivo histopathology!



On the Way to Personalized Medicine

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Dosimetry: A comparison

External Beam Therapy



Radionuclide Therapy



Courtesy Matthias Blaickner, AIT



Department of Nuclear Medicine / Center for P.E.T.

Zentralklinik Bad Berka, Germany

Patient dosimetry in PRRT using ¹⁷⁷Lu DOTA-NOC, ¹⁷⁷Lu DOTA-TATE and ¹⁷⁷Lu DOTA-TOC



0.03 (0.02-0.08)*

0.6 (0.3-1.6)*

DOTA-TOC

Department of Nuclear Medicine / Center for P.E.T.

*significant

Zentralklinik Bad Berka, Germany

4.9 (0.3-39.7)

0.7 (0.2-2.8)*

Dosimetry - Perspectives

Pre-therapeutic organ and tumor dosimetry using receptor PET/CT and longer lived positron emitters, e.g. Sc-44, Y-86 or Cu-64 and comparison with Ga-68 results.

Selection of the optimal peptide and radionuclide for individual therapy of each patient ("personalized dosimetry") by pretherapeutic measurement of organ and tumor doses.

Personalized dosimetry



Y-86 DOTA-NOC Receptor PET/CT



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Courtesy of Dr. Choi SJ

Therapeutic Radionuclides

Nuclide		Emission	Mean path length	
I-125	60.0d	auger	→ 10nm	Beta
At-211	7.2h	alpha	→ 65nm	Emitter
Lu-177	6.7d	beta/gamma	→ 0.7mm	Physical Properties of Radionuclides Used for PRRT
Cu-67	2.58d	beta/gamma		Remote to de la constant
I-131	8.04d	beta/gamma	→ 0.9mm	rose 11 12
Sm-153	1.95d	beta/gamma	→ 1.2mr	n
Re-186	3.8d	beta/gamma	1.8	3mm
P-32	14.3d	beta		2.9mm
Re-188	17h	beta/gamma		→ 3.5mm
In-114m	n 50d	beta/gamma		→ 3.6mm
Y-90	2.67d	beta		3.9mm

Somatostatin Receptors as Targets for Nuclear Medicine Imaging and Radionuclide Treatment

Helmut R. Maecke¹ and Jean Claude Reubi²

⁴Department of Nuclear Medicine, University Hospital Freiburg, Freiburg, Germany, and ²Division of Cell Biology and Experimental Cancer Research, Institute of Pathology, University of Berne, Berne, Switzerland

THERANOSTIC PAIRS

Personalized medicine is used along with targeted therapy in general. The targeting of somatostatin receptor-positive tumors is an ideal example of this approach. It combines powerful new diagnostics and radiotargeted therapeutics. A diagnostic scan with a γ - or β^+ -emitting nuclear probe is used to identify tumors and metastases that overexpress somatostatin receptors and is therefore predictive of the potential for targeted radionuclide therapy in patients. It also allows the study of dosimetry, thereby estimating

Eur J Nucl Med Mol Imaging DOI 10.1007/s00259-012-2330-6

GUIDELINES

EJNMMI February 2013 ePub ahead of print

The joint IAEA, EANM, and SNMMI practical guidance on peptide receptor radionuclide therapy (PRRNT) in neuroendocrine tumours

John J. Zaknun · L. Bodei · J. Mueller-Brand · M. E. Pavel · R. P. Baum · D. Hörsch · M. S. O'Dorisio · T. M. O'Dorisiol · J. R. Howe · M. Cremonesi · D. J. Kwekkeboom

The pdf of this guidance is available on our website www.prrtinfo.org

ENETS Center of Excellence since 2011 Zentralklinik Bad Berka*

Int. Medicine, Endocrinology, Gastroenterology, Oncology Thoracic, Abdominal/Visceral and Spinal Surgery Interventional Radiology Molecular Radiotherapy & Imaging (PET/CT Center)

including a specialized nuclear medicine ward, medical physcis and GMP radiopharmaceutical facilities/radiopharmacy center "THERANOSTIK"



Radiopeptide Therapy Cycles Zentralklinik Bad Berka 1999 - 2010



FUTURE OF CANCER TREATMENT

Cancers will be classified by molecular phenotypes Organ site \rightarrow secondary classification

Molecular phenotypes will be determined by molecular pathology and by molecular imaging studies (PET, SPECT, MRI, optical) using cancer type specific probes.

Treatment will be targeted specifically against the tumor.

Neuroendocrine tumors are a <u>paradigm</u> for this approach as molecular radiotherapy is applied based on molecular features (i.e. somatostatin receptor expression) of tumors and not based on the organ of origin of the tumor.

Primary tumors of patients with metastatic NETs treated by PRRT n=1100 patients



RADIOPEPTIDE THERAPY (ZKL BAD BERKA)



Neuroendocrine cancer of the right kidney with extensive bilateral liver metastases (size 3.7 cm in S7) and retroperitoneal lymph node (size up to 6.5 cm) and bone metastases.



MOLECULAR RESPONSE BY ⁶⁸GA DOTA-TOC PET/CT IMPROVEMENT OF KIDNEY FUNCTION IN A PATIENT WITH A SINGLE KIDNEY

May 2009 – before PRRT

TER 97 ml/min (35 %)

Sept. 2009 - after 1st PRRT

Jan. 2010 - 4mo after 2nd PRRT

TER 147 ml/min (54 %)

TER 202 ml/min (74 %)

Center for Molecular Radiotherapy / Department of Molecular Imaging (PET/CT) Zentralklinik Bad Berka



MOLECULAR RESPONSE DEMONSTRATED BY ⁶⁸GA DOTATOC PET/CT COMPLETE REMISSION OF LIVER METASTASES

Neoadjuvant sequential PRRT (Y-90 DOTA-TATE) of inoperable progressive pancreatic NET. 32-y-o female, refused chemotherapy.



Before PRRT-1 6 GBq Y-90 SUV 29.4 Before PRRT-2 4.5 GBq Y-90 SUV 25.4 5-mo after PRRT-2 pre Op. SUV 12.5

Laparotomy (PE): Highly differentiated NET, Ki-67 8%, CgA +/ NSE +

Sequential PRRT (Y-90 DOTA-TATE) of Inoperable Pancreatic NET



Jan. 2007

SUV 25.4 **May 2007**

SUV 12.5 **Oct. 2007**

Whipple' Operation – Complete Resection of Pancreatic NET after Neoadjuvant PRRT





Histology revealed nearly total tumor necrosis typical for radiation necrosis

Follow-up at 54 months: Complete Remission

Median overall survivial from start of DUO-PRRNT: 59 months (415 GEP-NET patients)




JOURNAL OF CLINICAL ONCOLOGY

ORIGINAL REPORT

Submitted November 15, 2010; accepted March 14, 2011; published online ahead of print at www.jco.org on May 9, 2011.

Response, Survival, and Long-Term Toxicity After Therapy With the Radiolabeled Somatostatin Analogue [⁹⁰Y-DOTA]-TOC in Metastasized Neuroendocrine Cancers

Anna Indiaf, Philippe Brunner, Navalas Marineek, Matthias Brief, Christian Schnidler, Hohmit Rasen, Helmitt R. Macke, Christoph Boeblitz, Jan Muller Brand, and Martin A. Walter

Screened for eligibility (N = 2,041)	No. of No. of Median Hazard 1.0 - Patients Deaths Survival Ratio P Disease control 621 280 38 years 0.41 y Progress < 001
Not eligible (n = 130) No visible tumor uptake (n = 109) Poor physical condition (n = 19) Impaired bone marrow function (n = 2)	Progress 438 211 1.4 years 0.8 Median: 45.6 months
Eligible for inclusion (n = 1,911)	IENINI 0.4 -
Not treated $(n = 802)$ Transferred to other treatment $(n = 505)$ Refused treatment $(n = 114)$ Loss of transferability $(n = 80)$ Costs not covered $(n = 62)$ Died before first treatment $(n = 41)$	STELE 0.2- 0 2 4 6 8 10 12 14
	Time Since Start of Treatment (years)
Treated with ("yttrium-DOTA)-TOC (ri = 1,109)	No. at risk Total 1,109 381 172 66 27 5 1 0 Disease
Included in Intention-to-treat analysis (n = 1,109) Available for follow-up (n = 1,100) Not available for follow-up (n = 9)	Progress 438 85 28 11 1 0 0 0

JOURNAL OF CLINICAL ONCOLOGY

ORIGINAL REPORT

Adverse events – Basel data

Hematological toxicity 3 or 4: 12.8 % (transient)

Renal toxicity

grade 4 (n = 67 pts) grade 5 (n = 35 pts):

9.2 %

Bad Berka (DUO-PRRT): 1/1002 patients (0.1 %) on dialysis

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New Avenues to Improve PRRT in Future

- **DUO-PRRT** (already routine at our center since 8 years)
- TANDEM-PRRT (concurrent Lu-177/Y-90 PRRT Kunikowska et al.)
- Intra-arterial PRRT (> 100 i.a. treatments up to now)
- **Combined PRRT** (in combination with other treatment modalities)
 - TACE, SIRT, RFA (Hörsch et a. ASCO 2010)
 - chemotherapy (e.g. Capecitabine, Doxorubicin)
 - kinase inhibitors (e.g. Sunitinib, Sorafenib)
- Intra-operative use of probes after PRRT with Lu-177
- Improved dosimetry and radioprotection

Improved peptides (e.g. antagonists) and novel molecules

Ga-68 Labeled Tracers in Clinical Use

- [⁶⁸Ga-DOTA,Tyr³]octreotide (DOTA-TOC)
- [68Ga-DOTA,1-Nal]octreotide (DOTA-NOC)*
- [68Ga-DOTA]-TATE*
- [68Ga-DOTA]-Lanreotide
- [68Ga-DOTA]-Bombesin / AMBA* and DEMOBESIN*
- [⁶⁸Ga-DOTA]-D-Glu-Gastrin (MTC, NET)*
- [⁶⁸Ga-DOTA]-F(ab')₂-herceptin (breast cancer)
- ⁶⁸Ga-Citrate (infection, inflammation)
- ⁶⁸Ga-DOTA-Tyrosin (brain tumors)*
- ⁶⁸Ga-DOTA-HSA Microspheres (lung perfusion)*
- ⁶⁸Ga-NODAGA-RGD (angiogenesis)*
- ⁶⁸Ga-BPAMP (osteoblastic metastases)*
- ⁶⁸Ga-DOTA-α-MSH (melanoma)*
- ⁶⁸Ga-DOTA-SHAL (lymphoma)*
- 68Ga-PSMA (prostate cancer
- ⁶⁸Ga-CXCR4 (acdenocarcinomas)
 ...and many more to come!

*first use in Bad Berka







2012; 2(5):437-447. doi: 10.7150/thno.3645

Review

THERANOSTICS: From Molecular Imaging Using Ga-68 Labeled Tracers and PET/CT to Personalized Radionuclide Therapy — The Bad Berka Experience

Richard P. Baum[™], Harshad R. Kulkarni

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Received: 2011.10.16; Accepted: 2011.12.02; Published: 2012.05.07

Antagonist labels more sst₂ sites than agonist in human cancer tissues



79



SMS-Agonist Ga-68 DOTA-TOC PET/CT

SMS-Antagonist: Ga-68 DOTA-JR10 PET/CT

Comparison of ¹⁷⁷Lu-DOTATATE and ¹⁷⁷Lu-DOTA-JR11 dosimetry

Patient with NEC (G3) of the bladder with lymphnode and uterus metastases, shows progression after surgery and treatment with Somatostatin analogues



Imaging of Neoangiogenesis



A new approach: [68Ga]-RGD

Dept. of Nuclear Medicine/P.E.T. Center, Zentralklinik Bad Berka

Ga-68 RGD PET/CT in adenocarcinoma of lung Molecular imaging of angiogenesis



Center for Molecular Radiotherapy / Department of Molecular Imaging (PET/CT) Zentralklinik Bad Berka

Breast cancer with extensive bone metastases (first in human study). In total, 25 metastases were RGD positive and 12 lesions were FDG-avid.



MOLECULAR IMAGING OF TUMOR NEOANGIONESIS BY THERANOST ⁶⁸GA NODAGA RGD vs. Metabolic FDG Imaging

TITI (0 (\$) (🔍

Cyclen-based tetraphosphinate chelator

for preparation of radiolabeled tetrameric bioconjugates with ultrahigh affinity



	Biomolecule
WUU Magi	
Bomolecule	
	Biomolecule

Compound	IС ₅₀ [рм]	relative activity
DOTPI(RGD) ₄	109 ± 7	12.2
Cu-DOTPI(RGD) ₄	73 ± 5	18.2
Lu-DOTPI(RGD) ₄	72 ± 4	18.5
TRAP(RGD) ₃	249 ± 7	5.3
Ga-TRAP(RGD) ₃	193 ± 5	6.9
c(RGDfK)	1330 ± 90	1

 177 Lu- and 64 Cu-DOTPI(RGD)4 are fully stable in DTPA / EDTA and plasma over 7 d / 12 h.

III \bigcirc \bigcirc \bigcirc Cu-64 labelled triazacyclononane-triphosphinate chelators



 $\alpha = 3$.



Simecek et al. Dalton Transactions 2012

Ga-68 TRAP(RGD)₃ PET/CT



Center for Molecular Imaging and Molecular Radiotherapy, Zentralklinik Bad Berka, Germany in collaboration with H.J. Wester (labeling performed using SCINTOMICS module)



Bone infiltration (L3) of plasmacytoma with invasion of the spinal cord and of the psoas muscle. There is also involvement of T-9 vertebra (marrow) and left scapula. Periarthritis right shoulder.

CXCR4 is highly expressed in various tumors and metastases

CXCR4

- Chemokine-Receptor (GPCR)
- Coreceptor for HIVinfection
- CXCL12 is the only endogenous ligand

SDF-1α (CXCL12)

Stromal Cell derived Factor

- chemotactic cytokine
- 68 amino acids

CXCR4 and SDF-1a mediate

- Physiological conditions: Homing of lymphocytes and hematopoietic stem and progenitor cells (HSPC)
- Pathological conditions: tumorigenesis tumor progression metastasis



Liotta LA Nature (2001)410:24

IIII O V P Functional roles of the chemokine receptor CXCR4

Co-receptor for HIV-entry



Angiogenesis

Tumor progression and metastasis



Healing and tissue remodelling



skin wounds, genetically diabetic mice, treated with 6mg/kg AMD3100 in saline or saline alone, and then examined 0, 7, and 14 days later



Theranostics 2013, Vol. 3, Issue 1





2013; 3(1):1-2. doi: 10.7150/thno.5760

Editorial

CXCR4 Chemokine Receptor Overview: Biology, Pathology and Applications in Imaging and Therapy

Orit Jacobson¹, Ido Dov Weiss²

- 1. Hadassah Hebrew University Hospital, Cyclotron radiochemistry unit, Jerusalem, Israel.
- 2. Hadassah Hebrew University Hospital, the Goldyne Savad Gene Therapy Institute, Jerusalem, Israel.

TIT O Ý 🛛

	cxxr4 Search Clear Fields >				
XCR4	CANCER TISSUE	100			
•	Overall cancer tissue statistics				
	0% 25% 50% 75%	100%			
OTEIN					
Y/ANTIGEN	arder Histological Alphabetical				
CIW .	Antibody CABDE 1447				
UNRADCA FLOR	Tissue Antibody staining	Level of antibody			
ISBUE	Breast cancer	staining			
	Carcinoid	Stream			
	8 Cervical cancer				
	S Colorectal cancer	Hoderate			
	Carl Endometrial cancer	Weak			
	Sioma	Negative			
	I Head and neck cancer				
	Sector Cancer				
	6 Lung cancer	Dictionary			
	lymphoma la	TSENT			
	Melanoma	(N) Dictionary			
	Ovarian cancer				
	Pancreatic cancer				
	Prostate cancer				
	Renal cancer				
	🗇 Skin cancer				
	Stomach cancer				
	Testis cancer				
	I Thyroid cancer				
	4 Urothelial cancer				



Structure of ⁶⁸Ga-CPCR4-2



Gourni E. et al. J Nucl Med. 2012; O.Demmer et al. Chem. Med. Chem. 2011

Imaging CXCR4 Receptor Expression of Lung Metastases



ex-vivo µ-Autoradiography: lung of a mouse 1h p.i. of n.c.a. ¹²⁴I-CPCR4 with OH-1 tumor on the shoulder



Quantitative PET Imaging Study

in OH-1 SCLC Tumor Bearing Mice



Gourni E. et al. J Nucl Med. 2012



[⁶⁸Ga]CPCR4-2 PET

[⁶⁸Ga]CPCR4-2 PET

[¹⁸F]FDG PET/CT



[⁶⁸Ga]CPCR4-2 PET/CT

Wester HJ, Schwaiger M, Beer A, Keller U et al. (unpublished)

68Ga CPCR4-2 PET/CT



Pancreatic head Right breast tumor LN 29 year-old female patient with poorly differentiated neuroendocrine carcinoma of unknown primary (CUP-NEC, first appearance in the left breast) with extensive lymph node metastases. Ga-68 CXCR-4 PET/CT shows intense CXCR-4 expression in the previously SMS-R positive metastases, most pronounced in the cervical and mediastinal lymph nodes as well as in the right breast (relatively mild to moderate in the other breast lesions). In the pancreatic head, a CXCR-4 positive, SMS-R negative lesion is detected (most probably corresponding to the primary tumor). Uptake is also noted in metastases in both humeral heads.

Center for Molecular Imaging and Molecular Radiotherapy, Zentralklinik Bad Berka, Germany in collaboration with H.J. Wester (labeling performed using SCINTOMICS module)

Lecture Outline

- Definition and principles of THERANOSTICS and Personalized Medicine
- THERANOSTIC radionuclides and Ga-68 generator
- Neuroendocrine tumors (NET) as a paradigm
- Diagnosis of NET by PET/CT (clinical applications)
- Dosimetry (organ & tumor dose calculations)
- Therapy of NET (Peptide Receptor Radiotherapy, PRRT)
- Future perspectives
 - new peptides (antagonists, CXCR4, RGD)
 - **PSMA: THERANOSTICS** potential for prostate ca.

Prostate Cancer 2013

- Prostate cancer is the most common cancer diagnosis in Western countries
 - > 220,000 new cases annually in the U.S
 - > 400,000 new cases in Europe
 - > 900,000 worldwide incidence
- Second most common cause of cancer-related death in Western men
 - Estimated 32,000 deaths in 2011 in U.S
 - Estimated 89,000 deaths in Europe
 - Estimated 258,000 deaths worldwide
- >20% reduction in deaths since 1990 in U.S
 - Screening
 - Improved treatment
 - Statin use
 - But increasing population of at-risk men

Molecular Imaging of Prostate Cancer by PET



Bombesin Receptors in Human Prostate Cancer

Histology Autoradiography



Bombesin receptors are expressed in PIN (prostate intraepithelial neoplasia) and prostate cancer, but not in normal prostate or hyperplasia (N=36)

Markwalder & Reubi. Cancer Res (1999) 59:1152-1159

Clinical Work with Bombesin-based Agonists

- ➢ ¹⁷⁷Lu-AMBA
- ➢ ⁶⁸Ga-AMBA
- ➢ ⁶⁸Ga-BZH3
- ➢ ^{99m}Tc-BOM

- L. Bodei et al, R.P. Baum et al
- R.P. Baum et al
- L. Strauss et al
- C. Van de Wiele et al
- F. Scopinaro et al



AMBA, a powerful bombesin-based agonist

Lantry et al. J Nucl Med 2007

R.P. Baum et al.J Nucl Med 2007 (abstr.)

L. Bodei et al. EJNMMI 2007 (abstr.)

1st World Congress on Ga-68 and Peptide Receptor Radionuclide Therapy (PRRNT)

THERANOSTICS - on the Way to Personalized Medicine





Asp296 Glu294 Ser298 Val295 Thr29 Ser293 Met299 vr292 Leu300 Val303 His291 Phe302 Arg288 Tyr290 Thr304 Ser305

Jensen RT et al, *Pharmacol Rev,* **2008**, 60, 1-42 Tokita K et al. *J Biol Chem,* **2001**, 276, 36652-36663







Patient with prostate cancer recurrence Increasing PSA, CI normal. Ga-68 GRPR antagonist PET/CT 60 Min p.i.

Ga-68 DEMOBESIN

Displacement of [¹²⁵I-Tyr⁴]bombesin (31,200 cpm; 39,000 dpm) from PC-3 membranes (50 µg/batch 19/07/2005) by increasing amounts of DOTA-DAla¹¹-Demobesin-1-mimic 11-11-2008



Lymph node mets



Ga-68 DOTA-Demobesin



F-18 Cholin



Ga-68 DOTA-Demobesin PET/CT in a 71 year-old patient with poorly differentiated carcinoma of the prostate demonstrates bombesin receptor positive mediastinal, axillary and abdominal lymph node metastases.

Center for Molecular Imaging and Molecular Radiotherapy, Zentralklinik Bad Berka, Germany



Post-therapy scans (anterior view) with Lu-177 DOTA-Demobesin in the same patient exhibit rapid wash-out of the tracer from the metastases with time.

IIII 🖉 🌾 😢 Urea-based ⁶⁸Ga-PSMA Inhibitor (GCP-II) for PET Imaging



Evaluation of PSMA-avid small molecules



SPECT/CT scans at 4 h p.i.



370 MBq, planar WB scans (left) and SPECT/CT, 4h p.i.

Barret JA et al. J Nucl Med 2013; 54:380–387

PSMA as a target for radiolabeled small molecules.



Fig. 1 a Maximum intensity projection image of a PET scan performed on day 5 after administration of ¹²⁴I-MIP-1095 shows multiple lesions in bones and lymph nodes, and also accumulation in the salivary and lacrimal glands. b Whole-body scan 7 days after administration of 5 GBq ¹³¹I-MIP-1466 in the same patient



Fig. 2 PET images in the same patient using the ⁶⁸Ga-labelled PSMA inhibitor Glu-NH-CO-NH-Lys(Ahx)-HBED-CC (a) and ¹⁸F-fluoroethylcholine (b). The scan with the PSMA ligand shows significantly more lesions than the fluoroethylcholine scan in which only one metastasis is seen

⁶⁸Ga-PSMA PET/CT



68Ga-PSMA PET/CT

Patient representative for disseminated lymph node and bone metastases of prostate cancer.

68Ga-PSMA PET/CT

Patient with low PSA level (0.01 ng/ml) and lymph node metastases.

Minimal PSA elevation despite visible tumor lesions suggests dedifferentiation of prostate cancer metastases.

At PSA levels < 2.2 ng/ml, lesions suspicious for cancer were observed in 60 % of the patients. At PSA levels > 2.2 ng/ml, lesions were detected in all patients.


New GCP-II Ligands for THERANOSTICS of Prostate Ca





Internalization, LNCaP cells (37°C, DMEM + 5% BSA)

Activity accumulation of EuK-Sub-KFF-[68Ga]DOTA

Activity accumulation of 0.5 nM EuK-Sub-KFF[⁶⁸Ga]DOTA





Internalization, 125.000 LNCaP cells/well; 37°C, DMEM + 5% BSA



New GCP-II Ligands for Imaging and Therapy



Weineisen M, Schottelius M, Simecek J et al. (unpublished)

Indication: rising PSA (8 ng/ml); intense PSMA-avid tumor in right lower quadrant of the prostate with an SUV of 9.3, high probability of malignancy



Ga-68 PSMA PET/CT – Detection of Primary Tumor

Benign prostatic hyperplasia: Enlarged prostate (6.2 x 5.6 cm), PSA 50 ng/ml. No evidence of a PSMA-avid primary tumor or metastases



Ga-68 PSMA PET/CT seems to be highly specific!

Recurrent prostate cancer with multiple metastases (pulmonary, bone and lymph nodes)



Center for Molecular Imaging and Molecular Radiotherapy, Zentralklinik Bad Berka, Germany in collaboration with H.J. Wester (labeling performed using SCINTOMICS module)

MULTIPLE METASTASES OF PROSTATE CANCER



Ga-68 PSMA PET/CT

Center for Molecular Imaging and Molecular Radiotherapy, Zentralklinik Bad Berka, Germany in collaboration with H.J. Wester (labeling performed using SCINTOMICS module)





Activity in intestines and kidneys







Left pedicle of T-8

Lu-177 PSMA SPECT/CT 24h post-therapy

Left bronchohilar LNM

<u>First Results</u> 4200 MBq EuK-Sub-KFF-[¹⁷⁷Lu]DOTAGA

	MBq		
Time	GK	Kidney left	Kidney right
0,6	4200,00	133,69	104,40
3,6	2104,04	60,66	48,50
20,7	246,66	23,38	23,17
46,0	154,57	14,13	13,42
69,1	104,26	8,72	8,04
	%IA		
Time	GK	Kidney left	Kidney right
0,6	100,0%	3,2%	2,5%
3,6	50,1%	1,4%	1,2%
20,7	5,9%	0,6%	0,6%
46,0	3,7%	0,3%	0,3%
69,1	2,5%	0,2%	0,2%



Half-life:

Whole Body = 45h Kidneys = 34h

First in human treatment with a PSMA targeting probe - EuK-Sub-KFF-[¹⁷⁷Lu]DOTAGA for Lu-177 labeling



Center for Molecular Imaging and Molecular Radiotherapy, Zentralklinik Bad Berka, Germany in collaboration with H.J. Wester (labeling performed using SCINTOMICS module)



F-18 Fluoride PET/CT

Target lesion in left ischium



Ga-68 PSMA PET/CT shows additionally liver metastasis and infiltration of the glans penis (histologically proven)

Skeletal metastases





CT positive

CT negative (marrow metastases)

Ga-68 PSMA PET/CT

Eur J Nucl Med Mol Imaging (2010) 37:834 DOI 10.1007/s00259-009-1355-y

IMAGE OF THE MONTH

PET/CT imaging of osteoblastic bone metastases with ⁶⁸Ga-bisphosphonates: first human study

Marco Fellner • Richard P. Baum • Vojtěch Kubiček • Petr Hermann • Ivan Lukeš • Vikas Prasad • Frank Rösch



Acknowledgments Support by the EC through COST actions D38 and BM0607 is gratefully acknowledged. M. Fellner · F. Rösch (Ed) Institute of Nuclear Chemistry, Johannes Gutenberg University, Fritz-Strassmann-Weg 2, 55128 Mainz, Germany e-mail: frank.rocsch@uni-mainz.dc

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V. Kubiček · P. Hermann · L Lukeš Department of Inorganic Chemistry, Faculty of Science, Charles University in Prague, Prague, Czech Republic

Dept. of Nuclear Medicine/P.E.T. Center, Zentralklinik Bad Berka

BPAMD

- labeling in almost quantitative yield with ¹⁷⁷Lu
- perfect for routine: dilute 15 GBq batch to 20 mL in syringe with stabilizing gentisic acid (1-2 mg total)
- stability of >95 % in 24 h

Combine diagnosis AND therapy with the same ligand!

=THERANOSTICS



Lu-177 BPAMD post-therapy planar images (posterior views) in a patient with metastatic prostate cancer, showing rapid renal clearance, high uptake in skeletal metastases and long effective half life.



Baum and Kulkarni. Theranostics 2012, 2(5)

72 year-old male patient with poorly differentiated adenocarcinoma of the prostate, Gleason score 7 (4 +3), first diagnosed 6 years before, s.p. radical prostatectomy, pelvic lymphadenectomy, EBRT (66.6 Gy) of the prostate, seminal vesicles and iliac lymph nodes developed painful bone metastases treated by androgen blockade and biphosphonates, followed by palliative treatment with Sm-153 EDTMP.

For progressive metastases with intense skeletal pain, he underwent 4 cycles of Lu-177 BPAMD treatment with a cumulative administered activity of 19.9 GBq Lu-177. F-18 (fluoride) PET/CT was performed pre- and post-therapy. Compared to the previous examination, there was a mixed pattern, i.e., on the one hand, clear regression with significant reduction in the intensive osteometabolic activity (uptake on F-18 PET/CT) of the metastases in pelvis and sacrum, but also evidence of new osseous metastases in the axial skeleton.



A: F-18 PET/CT MIP image pre-therapy;
F: F-18 PET/CT MIP image after 4 cycles of Lu-177 BPAMD treatment;
B, C, D, E: Lu-177 BPAMD whole-body planar images 45 hours after injection (first, second, third and fourth cycles respectively).



Macrocyclic Bisphosphonates



DOTA derivatives







NOTA derivatives





M. Fellner, R. P. Baum, V. Kubíček, P. Hermann, I. Lukeš, V. Prasat, F. Rösch *Eur. J. Nucl. Med. Mol. Imaging* 2010, *37*, 834.



ex vivo Biodistribution



male Wistar rats, 60 min p.i.



* Data: Fellner M, Bergmann R, In Vivo Comparison of DOTA Based Ga-68 Bisphosphonates for Bone Imaging 2013

Ga-68 NO2A-BP - a new and improved biphosphonate



77-year-old male with Ca prostate with multiple skeletal metastases, s.p. palliative TUR-P, lymphadenectomy, radiotherapy to prostate and multiple vertebrae, palliative chemotherapy and Strontium-89 therapy

Translational Research: Crossing the Valley of Death

National Institutes of Health (NIH):

- "Clinical and basic scientists don't really communicate"
- Excellent basic research, but lack of translation
- Where do we go from here?



Nature 453, 840-842, 2008



11111 Zentralklinik Bad Berka IN THE REPART

Thanks to the team...

and to our patients!

International Symposium Notecular Diagnosis and Therepy of Cancer in Nuclear Medicine The Future is now May 2-3, 2008

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Memorizing.. You remember 10 % - reading 20% - listening 30 % - seeing 50 % - seeing & hearing 70 % - talking about 90 % - what you're doing