



Highlights of the 30th Annual Congress of the EANM, Vienna 2017: “Yes we can – make nuclear medicine great again”

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Abstract

The 30th Annual Congress of the European Association of Nuclear Medicine (EANM) was held in Vienna, Austria, from 21 to 25 October 2017 under the chairmanship of Professor Francesco Giammarile. As always, the Congress was a great success: more than 6,379 participants came from 90 countries from all continents. Participants were presented with an excellent programme consisting of symposia, and scientific and featured sessions, CME sessions, and plenary lectures. These lectures were devoted to nuclear medicine imaging and therapy, including hybrid imaging and molecular life sciences. Additionally, the latest technology and innovations in the field were presented, and added to the success of the Congress. This review summarizes the major scientific contributions which were selected from more than 1,900 submitted abstracts, and presented in the closing highlights session. They cover the diverse areas of nuclear medicine, with particular focus on oncology, cardiovascular science, neurology, technological innovation and novel tracers, and also other clinical sciences. A particular focus of the Congress was on targeted radionuclide-based therapies, which all show promising and great innovations. The Congress was a unique opportunity to be thoroughly updated on this research. This Highlights Lecture could only be a brief summary of the large amount of data presented and discussed during the meeting, which can be found in much greater detail in the Congress proceedings book, published as volume 44, supplement 2 of the *European Journal of Nuclear Medicine and Molecular Imaging* in October 2017.

Keywords EANM 2017 · Annual congress · Abstracts · Highlights · Nuclear medicine · Radionuclide therapy and dosimetry · Radiopharmaceuticals · PET · SPECT

Introduction

From 21 to 25 October 2017 the 30th Annual Congress of the European Association of Nuclear Medicine (EANM 2017) took place in Vienna, Austria. With more than 6,379 participants on site and 1,571 online this congress is by far the most important European event in nuclear medicine, bringing together a multidisciplinary community involved in the different fields of nuclear medicine. Participants from 90 different countries attended, with more than 1,300 attendees from non-European countries, indicating the worldwide importance

of the meeting. The Scientific Programme Committee chaired by Francesco Giammarile organized the programme that covered all topics relevant to nuclear medicine comprising 529 oral and 1,021 poster presentations in more than 100 sessions during the meeting.

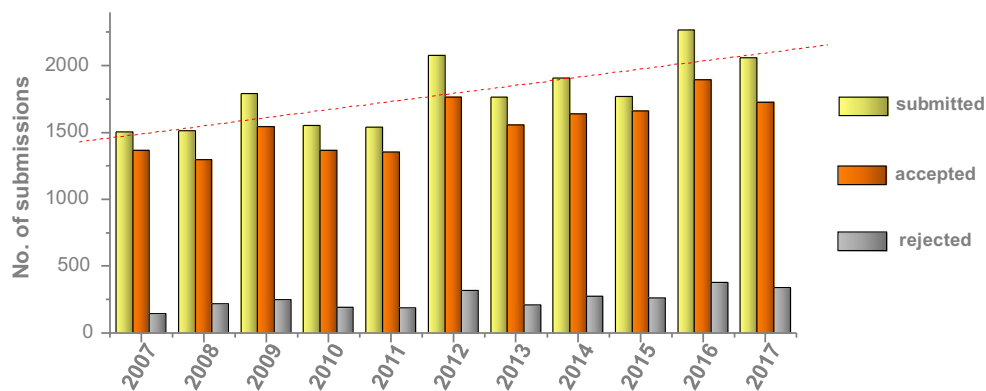
Overall 1,925 scientific abstracts were submitted and finally 1,613 were accepted (16.2% rejection rate). This number was only slightly topped by last year's meeting in Barcelona and the 25th Annual Meeting in Milan 2012 (see Fig. 1). The highest numbers of abstracts were received from Italy (12.3%), Germany (9.1%), Spain (7.1%) and France (5.6%). At 5.5%, Japan was the best-represented non-European country. Overall 75% of abstracts were from Europe, 15% from Asia, 7% from the Americas, and a small fraction from Africa and Australia; details are provided in Fig. 2. Looking at the cities from where abstracts were submitted, most European abstracts came from Munich (321), followed by Rome (265), London (239) and Genova (237). Most abstracts from outside Europe came from Tehran (141), Seoul (131), Beijing (126) and New York (110).

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Fig. 1 Numbers of abstracts submitted, accepted and rejected for presentation at EANM annual meetings since 2007



The abstracts covered a wide range of topics. Even though research related to applications of nuclear medicine in oncology dominated, an equal number of abstracts related to other clinical areas including cardiovascular science (8.1%), neurosciences (8.8%) and basic sciences including physics and instrumentation (12.9%) and radiopharmacy/radiochemistry (11.0%). The translational and multidisciplinary character of the Congress was indicated by the fact that the specific topics Translation (‘from molecule to man’, M2M) included 400 abstracts and Dosimetry & Radionuclide Therapy (DoMoRe) included 457 abstracts.

However, it is not the overall numbers that are important, but the content of the abstracts regarding novel findings and their impact on advances in the field of nuclear medicine. The EANM 2017 Highlights Lecture included a selection of the work presented at the meeting. The abstracts were initially rated by selected experts in the individual categories, who also proposed 421 of the best abstracts for possible inclusion in this year’s Highlights Lecture. From these abstracts, 46 were finally chosen for inclusion as examples of a large number of submissions that were often comparable and scientifically of equal importance. The authors (S.F., C.D.) had the honour to present the Highlights Lecture at this meeting. In preparing for

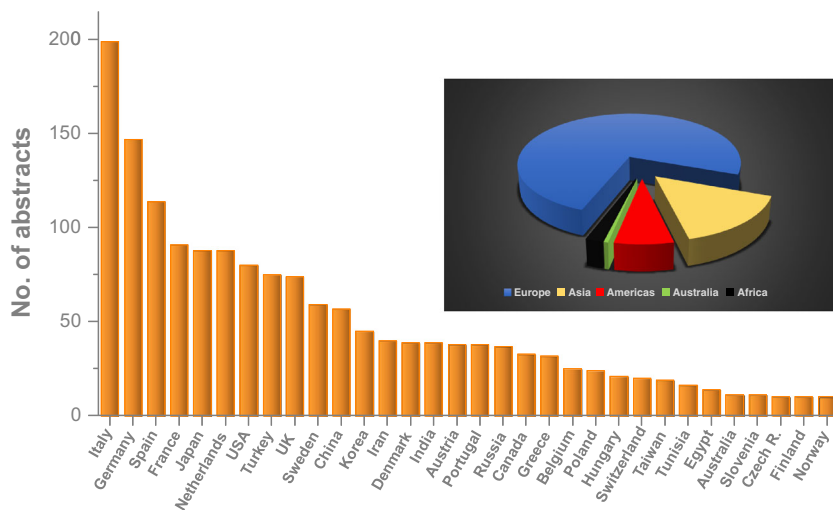
the task of selecting the submissions to be included, they asked: Could this congress “make nuclear medicine great again”? (a reference to the recent campaign of the US president).

We invite readers to judge for themselves and to read this year’s highlights of EANM 2017 in different categories. As there is no Oscar statue awarded, we proposed that the submitting scientists be honoured with a virtual George and Marie statue, in memory of George von Hevesy and Marie Curie-Sklodowska, considered the pioneers of diagnostic and therapeutic nuclear medicine applications, who, we are sure, would have enjoyed the presentations would have acknowledged their scientific quality.

Physics and instrumentation

Novel instrumentation for improved diagnostic procedures has been a major driving force for advancing molecular imaging, in particular for PET applications over recent years, accompanied by applied physics to realize the full potential of the new technologies. This meeting continued this trend and gave a glimpse of what may lie available in years to come.

Fig. 2 Abstracts submitted to EANM 2017 by country and continent



Cates and Levin [1] (Stanford, USA) presented an outstanding paper on a promising new PET detector design. They changed the readout of the detectors from the end to the side which led to lower losses in signal while also allowing a much higher timer resolution down to 100 ps. Implemented in a clinical detector system an increase in signal to noise ratio by a factor of 5 could be achieved. These innovations could be packaged into a practical clinical detector module with mixed analogue-digital multiplexing schemes that require a low number of readout channels, leading to a faster and more sensitive PET detector.

A study in collaboration between Belgian, Dutch and US centres was presented by Vandenberghe et al. [2]. They presented a cost-efficient 2-mm resolution whole-body PET design with a 1 m field of view extendable to 2 m, also based on a cost-effective design. With this approach they were able to achieve a 20 times faster PET acquisition or alternatively a 20-fold reduction in dose, further improving and simplifying PET scans (Fig. 3).

Novel technologies also have to be translated into clinical practice and show their added value in a larger setting. In this respect two presentations reflect such developments. One important topic is to harmonize readouts to establish clinical standards in particular for the quantification of molecular processes, which is one of the advantages of nuclear imaging methods. Kaalep et al. [3] (Tallinn, Estonia) presented a collaborative study (several European and one US centre) investigating the feasibility of harmonizing the performance of state-of-the-art PET/CT systems. They performed phantom studies on the newest time of flight (ToF) PET/CT systems available with resolution modelling/point spread function (PSF) technologies, and showed that it is possible to harmonize the systems' retard, which may provide the basis for a wider study of system harmonization.

Novel tools to utilize the full data generated by novel imaging studies are advancing rapidly. Grueneisen et al. [4] (Essen, Germany) used radiomics to predict the N and M stage of cervical cancer using PET/MRI. They used the PET/MRI signal and both PET and MRI images and applied a radiomics approach to define certain features typical of certain types of tumour. They achieved very high sensitivity and specificity for predicting N and M stage of cervical cancers, providing a number of noninvasive biomarkers and facilitating improved tumour evaluation and treatment planning. Thus, they were also able to show the benefit of PET/MRI over MRI alone.

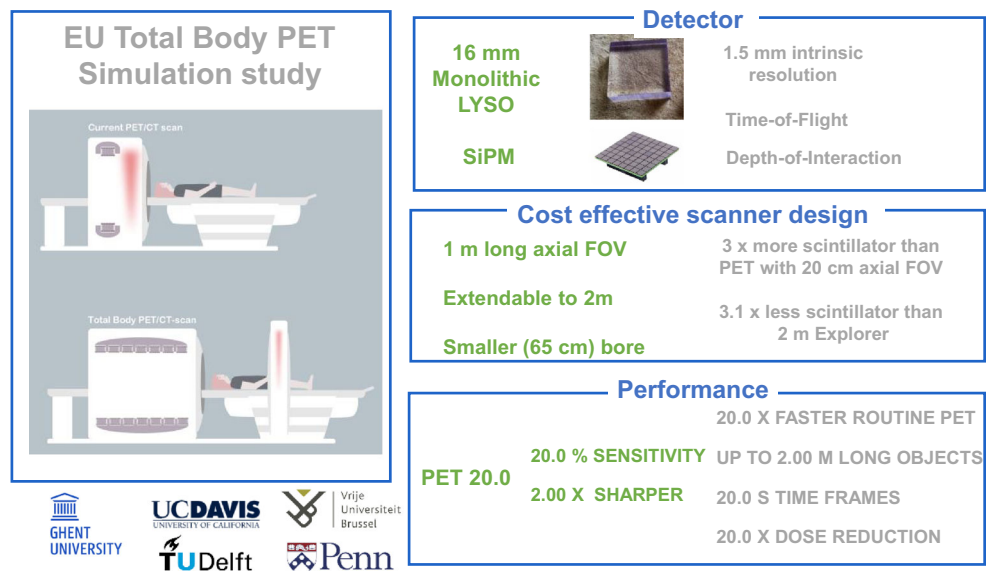
Novel technologies also have to show benefits in terms of patient safety. In this respect the presentation of Petoussi-Hens et al. [5] (Neuherberg, Germany) should be mentioned. These authors described a voxel-based dosimetry program able to accommodate the new dosimetric International Commission on Radiological Protection (ICRP) framework for simplified implementation in the clinical setting to provide more accurate estimation of radiation dose in individual patients.

Cardiovascular science

In this category a large number of submitted abstracts were on the subject of the diagnosis of myocardial infarction. Even though the topic as such is not novel, new opportunities in this area are emerging, in particular for imaging plaques and thrombosis. Another novel topic, which was dealt with in several presentations, is PET imaging of cardiac amyloidosis.

Rischpler et al. [6] (Munich, Germany) evaluated the relationships among myocardial inflammation, the area at risk (AAR), oedema and tissue damage using different approaches. In particular, they compared data derived from ^{99m}Tc-sestamibi SPECT and FDG PET/MRI and other

Fig. 3 PET 20.0: a cost-efficient, 2.00-mm resolution total body monolithic PET system with very high sensitivity and an adaptive axial field of view up to 2.00 m, presented by Vandenberghe et al. of the University of Ghent in collaboration with other Dutch, Belgian and US institutions [2]



MRI information. They found that the AAR was quite similar between MRI and sestamibi SPECT, but at the same time there was a poor correlation between oedema and AAR, and the association between oedema and irreversible tissue damage and inflammation indicates that MRI should not be used for AAR assessment.

Novel targets for cardiovascular applications were also presented. A very innovative exploratory study involving collaboration between the two universities of Munich and a Belgian centre was presented by Varasteh et al. [7]. They targeted the mannose receptor (CD206) expressed on activated reparative macrophages with a ^{68}Ga -NOTA anti-CD206 Nanobody for imaging the mannose receptor in mice with experimental myocardial infarction. With this approach they were able to depict the area where activated macrophages accumulate, which is strictly associated with the infarction healing process. In a second presentation by the same group [8], the use of the same radiotracer (^{68}Ga -NOTA anti-CD206 Nanobody) for a different application, to identify atherosclerotic plaque in apolipoprotein E-knockout (ApoEKO) mice, was reported, suggesting that NOTA anti-CD206 Nanobody is also a promising tracer for non-invasive detection of atherosclerotic plaques.

An exploratory open-label clinical study was presented by Kim et al. [9] (Seoul, Korea). They reported that it may be possible to diagnose deep venous thrombosis and pulmonary embolism using ^{18}F -GP1 PET/CT (Fig. 4). They targeted a glycol-protein receptor (GPIIb/IIIa) overexpressed on activated platelets and they were able to show the presence in vivo of pulmonary embolism as well as deep venous thrombosis with an extremely high sensitivity in the range 94–100%. They were also able to detect additional sites of venous thrombosis that were not detected by conventional imaging methods.

A representative study of cardiac amyloidosis imaging is that of Genovesi et al. [10] (Pisa, Italy) who investigated the use of ^{18}F -florbetaben PET/CT in this indication. They used dynamic scans and were able to clearly discriminate the patterns of

accumulation in the two different types of cardiac amyloidosis, transthyretin-related amyloidosis (ATTR) and light chain-related amyloidosis (AL). On the basis of the different wash-in and wash-out characteristics of the tracer in the ATTR and AL groups, ^{18}F -florbetaben PET/CT was able to differentiate the two types of amyloidosis indicating that brain involvement in the context of systemic amyloidosis in ATTR patients could not be due to the same type of amyloid deposits, which involve other organs including the heart and bone marrow.

Basic science and preclinical developments

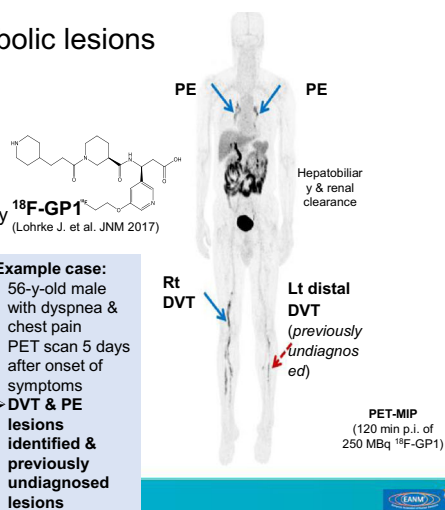
Nuclear medicine has the unique potential to become great because of translation of developments in basic biomedical sciences into clinical applications. A number of presentations exemplify this in an impressive way with novel radiotracers and basic studies on novel, clinically relevant targets.

Amor-Coarasa et al. [11] (New York City, USA) reported on the development of a new ligand, ^{18}F -RPS-544, that targets the CXCR4 receptor, and is relevant for the evaluation of tumour progression and metastatic potential. They performed PET studies in nude mice bearing bilateral (CXCR4+ and CXCR4-) PC3 xenograft tumours and showed excellent tumour targeting, a sevenfold improvement over existing ^{18}F -fluorinated ligands reported to date resulting in high tumour to background ratios. Baranski et al. [12] (Heidelberg, Germany) presented a study on a novel dual labelled compound based on PSMA-11 (Fig. 5) evaluating the potential of a combination of preoperative staging by PET/CT and fluorescence-guided surgery for detecting metastasis or neoplastic lymph nodes in patients with prostate cancer. They showed that this compound (^{68}Ga -PSMA-HBED-CC-IRDye800CW) provides PSMA-specific PET and fluorescence signals with promising pharmacokinetic properties in the surgical setting indicating a high potential for future clinical translation.

Fig. 4 Diagnosis of deep venous thrombosis and pulmonary embolism using ^{18}F -GP1 positron emission tomography. An exploratory open-label study from Seoul [9] showing excellent targeting of GPIIb/IIIa receptors on activated platelets leading to high sensitivity and specificity

^{18}F -GP1 PET identified thromboembolic lesions in all patients recruited

- **Clinical proof of concept obtained for detection of thromboembolic lesions with ^{18}F -GP1 PET**
- Target: GPIIb/IIIa receptors on activated platelets
- Recruitment based on acute symptoms of pulmonary embolism (PE) or deep vein thrombosis (DVT)
- 19 patients examined, many patients with both events
 - 15/16 (94%) patients with DVT positive with GP1
 - 14/14 (100%) patients with PE positive with GP1
- Seen with GP1 and not seen on standard imaging:
 - 23 additional DVT lesions in 13 patients
 - 3 additional PE lesions in 3 patients
- GP1 lesion detection rate on the artery / vein level may depend on the thrombus age



Technology: Dual-labeled PSMA- Inhibitors

In vivo Proof of Concept Studies

- High affinity to PSMA and specific internalisation
- High tumor-to-background ratio



3/25/2018 | Ann-Christin Baranski

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Fig. 5 ⁶⁸Ga-PSMA-HBED-CC-IRDye800CW, a novel dual labelled PSMA inhibitor with promising properties that may allow PET/CT to be combined with fluorescence-guided surgery for the evaluation of prostate cancer [12]

Heskamp et al. [13] (Nijmegen, The Netherlands) presented a study exploring the potential of PD-L1 microSPECT/CT imaging for monitoring changes in PD-L1 expression in tumours during radiotherapy (RT), of relevance in patients receiving immune checkpoint inhibitor treatment. They analysed an animal model which was irradiated and 1 day later received 25 MBq ¹¹¹In-labelled anti-PD-L1 antibody followed by microSPECT/CT imaging, in comparison with non-irradiated control mice. They showed that expression of the PD-L1 target increases after irradiation. Xu et al. [14] (Beijing, China) presented a similar study. These authors used an anti-PD-L1-labelled antibody, ⁶⁴Cu-NOTA-Ab1881, in an animal model expressing the PD-L1 target. Using this labelled antibody they were able to predict therapy outcome, distinguishing PD-L1-positive and PD-L1-negative xenografts in the mouse model with PET imaging, and also demonstrated therapeutic efficacy, which was highly correlated with the tumour uptake of ⁶⁴Cu-NOTA-Ab1881. These studies suggested that targeting PD-L1 could be used to monitor PDL1 expression during immune checkpoint inhibitor administration to select patients benefiting from treatment and to develop optimal treatment strategies.

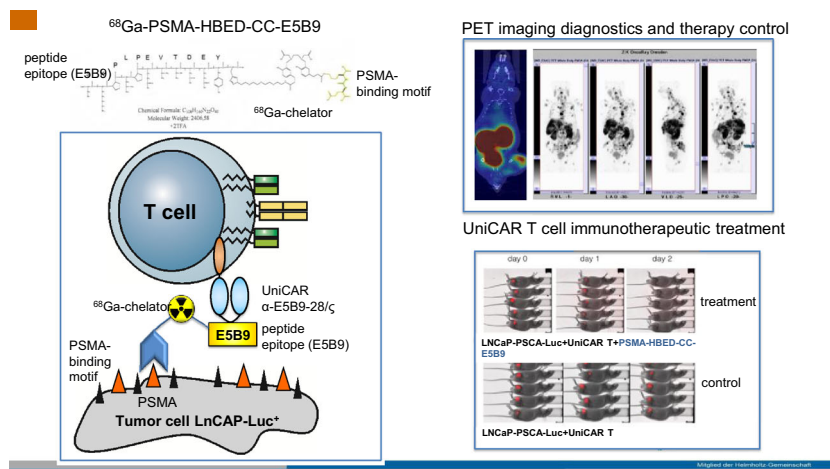
D'Alessandria et al. [15] (Munich, Germany) presented the results of a study comparing galectin-3 immunotargeting and radioiodine imaging in thyroid orthotopic tumour models. These authors showed that galectin-3 is expressed in thyroid carcinomas but is not related to the sodium iodide symporter in animals models. This approach could allow evaluation of

thyroid cancer independently from the sodium iodide transporter status, and is a promising approach to changing the management of patients with thyroid cancer who lose radioiodine uptake in the clinical setting.

In relation to radionuclide therapy, a presentation by Altai et al. [16] (Uppsala, Sweden) focused on pretargeted radionuclide therapy. They used a HER-2-targeting Affibody modified with peptide nucleic acids (PNAs) that allowed pretargeting by first injecting the PNA-Affibody followed by ¹⁷⁷Lu-PNA in the therapeutic setting. The use of Affibody-based PNA-mediated pretargeting was able to deliver high doses to tumours with a very good therapeutic effect in a preclinical model, while sparing the kidneys, which has so far been the problem with this therapeutic approach.

With a special kind of theranostic approach, Bergmann et al. [17] (Dresden, Germany) addressed a new concept that involved targeting PSMA-expressing tumours using a novel immunotherapeutic technique (UniCAR T). Using a ⁶⁸Ga-labelled modified PSMA 11 molecule with a recognition site for T cells (HBED-CC-PSMA-E5B9, PSMA-TM), the therapeutic efficacy of T cells is switched on leading to site-directed therapy that was demonstrated in an experimental animal model (Fig. 6). PSMA-TM was shown to be a useful PET imaging agent for noninvasively following the progress of individualized PSMA-directed immunotherapeutic tumour treatment. This is a nice proof of concept for the use of nuclear imaging to plan, monitor and optimize patient treatments based on novel immunotherapy concepts.

Fig. 6 A theranostic approach, presented by Bergmann et al. [17], used to monitor a novel immunotherapeutic technique (UniCAR T) based on site-directed target molecules that upon binding selectively turn on the immunotherapeutic action of T cells. A ^{68}Ga -PSMA targeting molecule is able to initialize the therapeutic effect and at the same time allows the targeting efficiency to be monitored by PET



Neuroscience

A great variety of presentations covered novel applications in neuroscience, reflecting the high potential of nuclear medicine for visualizing numerous targets for a variety of diseases. In particular, interesting contributions in relation to Parkinson's disease (PD), cognitive impairment and neuroinflammation were presented.

Hooshyar Yousefi et al. [18] (Munich, Germany) reported a first imaging study investigating a therapeutic target in PD, alpha-synuclein (α -syn), using a fluorinated compound, ^{18}F -DABTA. They were able to demonstrate in an animal model a significantly increased PET signal in regions with abundant α -syn (substantia nigra, thalamus, cerebellum and brainstem). This selective α -syn tracer may allow early diagnosis of α -synucleinopathies such as PD and dementia with Lewy bodies.

Another target, glutamate receptor type 5 (mGluR5), is a potentially important regulator of the most common excitatory neurotransmitter system in the brain. Kang et al. [19] (New York, USA) explored the possibility of a correlation between mGluR5 and PD using the selective mGluR5 PET probe ^{18}F -FPEB in nine patients with PD compared with a control group, and showed a significant enhancement of binding of the radiotracer in PD patients compared with the healthy controls. This study indicates that the response to dopaminergic denervation in PD is mediated in part by the mGluR5 system.

A very interesting first-in-human study was presented by Arlicot et al. [20] (Tours, France). They demonstrated a very reliable and convenient PET protocol for dopaminergic dysfunction imaging using a novel fluorinated compound, ^{18}F -LBT-999, a neuronal dopamine transporter ligand, that could potentially replace current SPECT imaging. ^{18}F -LBT-999 PET imaging showed the conventional depiction of the basal ganglia in healthy volunteers, but showed a significant decrease in uptake in patients with PD, demonstrating easy quantification and excellent discrimination between patients with early PD and healthy controls.

Another highly interesting abstract was presented by Unterrainer et al. [21] (Munich, Germany). They reported a study of the mitochondrial translocator protein (TSPO) expressed in neuroinflammation (Fig. 7) and upregulated in high-grade glioma (HGG), and ^{18}F -GE-180, a novel third generation TSPO receptor ligand. In this first-in-human study performed in 12 patients with HGG, ^{18}F -GE-180 PET was shown to be a promising tool for assessing the extent of the malignancy before and after radiation therapy, and also for evaluating response to radiation therapy.

Tau radiotracers mimic intracellular tau deposition, a pathological feature of the main neurodegenerative disorders. Seibyl et al. [22] (New Haven, USA) presented a new fluorinated compound, ^{18}F -PI-2620. They performed first-in-human PET studies in patients with Alzheimer's disease (AD) and progressive supranuclear palsy (PSP), in comparison with a healthy group, revealing a different pattern of accumulation in AD patients, especially in the temporal lobes, and different patterns of distribution in PSP patients, demonstrating a high target specificity and high signal in regions of expected tau pathology.

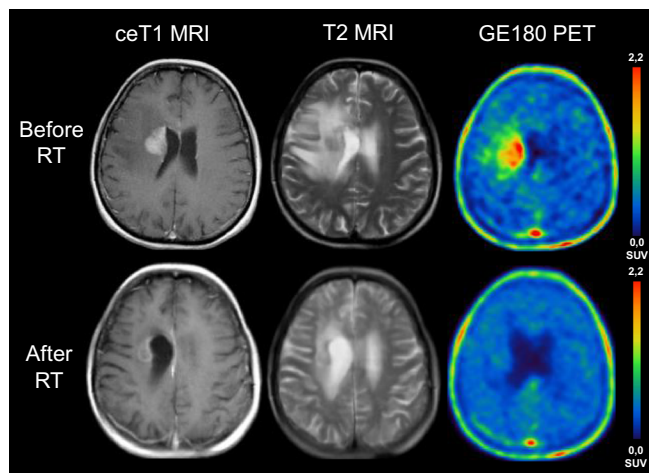
New radiopharmaceuticals: clinical applications

Based on many excellent submissions describing the first clinical applications of novel radiopharmaceuticals, a separate category was dedicated solely to this topic, involving imaging of beta cells, apoptosis, adrenal and neuroendocrine tumours.

Boss et al. [23] (Nijmegen, The Netherlands) used ^{68}Ga -exendin-4 PET/CT to image beta cells in patients after gastric bypass surgery. Those patients who benefited most from bypass surgery had an increased beta-cell mass (BCM), whereas BCM was lower in patient with incomplete remission. Quantifying BCM could therefore predict the outcome of surgery and could be used for personalizing treatment. Dubash et al. [24] (London,

➤ Heterogeneous PET - response to radiotherapy

- 2/8 patients: nearly complete remission of the extent of TSPO expression (92.0% and 94.7%); see *Figure for an example*
- 5/8 patients: reduction between 26.1% and 63.2%
- 1/8 patients: increase of extent of TSPO expression of 34.8% despite the previous radiotherapy



■ CONCLUSION

- [^{18}F]GE-180 PET provides a next ordinary high tumor-to-background contrast in HGG patients
- [^{18}F]GE-180 PET might represent a promising tool for HGG delineation with a potential influence on treatment planning
- TSPO expression assessed with [^{18}F]GE-180 PET might be a possible imaging biomarker for response assessment in HGG patients
- Correlation of [^{18}F]GE-180 PET with progression-free-survival and overall-survival is ongoing

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Fig. 7 TSPO PET imaging of high-grade glioma (HGG) using the novel ligand ^{18}F -GE-180. First-in-human results in the course of radiotherapy [21]

UK) reported a first-in-human study with ^{18}F -ICMT-11, targeting caspase 3/7, which is involved in apoptosis, as a novel marker for programmed cell death. They evaluated 17 patients with breast and lung cancer, and analysed the pattern of expression of this caspase. They concluded that fluorinated ICMT-11 can be used as a potential biomarker for chemotherapy-induced apoptosis. Schirbel et al. [25] (Wurzburg, Germany) presented a new compound called IMAZA, labelled with ^{123}I and ^{131}I , for targeting advanced adrenocortical carcinoma. Compared to the first-generation iodometomidate compound (IMTO), ^{123}I -IMAZA had higher metabolic stability resulting in highly improved image contrast, which allowed translation to therapy of this rare tumour with ^{131}I -IMAZA with very promising therapeutic efficacy and few side effects (Fig. 8).

Rischpler et al. [26] (Munich, Germany) presented data on LMI1195, a novel ^{18}F -labelled radiotracer for noninvasive PET imaging of the norepinephrine transporter, which is expressed in pheochromocytoma and paraganglioma. LMI1195 demonstrated better imaging properties than ^{123}I -MIBG, and is highly promising for the confirmation, exclusion and staging of tumours of the adrenal medulla or the sympathetic trunk with the potential to move from SPECT to PET in this indication. An interesting study was presented by Zheng et al. [27] (Beijing, China). They reported the first-in-human application of a dual-targeted radiopharmaceutical, ^{68}Ga -TATE-RGD, that binds to both the somatostatin receptor and integrin $\alpha\text{v}\beta3$. In 37 patients with non-small-cell lung cancer and neuroendocrine tumours (NET) they found that ^{68}Ga -TATE-RGD provided better quality images than other single-targeted radiopharmaceuticals.

Clinical oncology

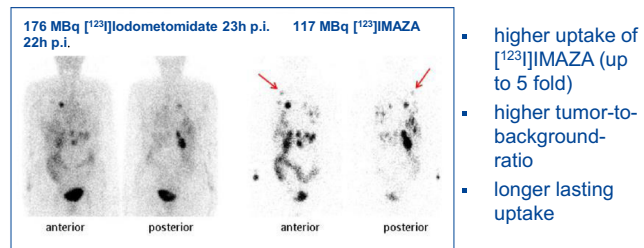
The broad progress in clinical applications in oncology was thoroughly represented in the meeting. A variety of topics, exemplified by an excellent presentation on radiomics, CXCR4 targeting, breast cancer imaging and hypoxia, were covered.

Liu et al. [28] (New York, USA) reported the identification of early small lung cancer with aggressive features (solid and micropapillary patterns) using a combination of PET and CT radiomics analysis. They extensively reviewed 170 radiomics features to identify aggressive components, and found a very interesting correlations between 29 CT and 65 PET radiomics features. Both PET and CT radiomics features were found to be potential biomarkers of aggressive lung adenocarcinoma subtypes with the potential to discriminate between patients requiring rather extensive surgery and those in whom a conservative surgical approach would be sufficient. An interesting study was presented by Watts et al. [29] (Chandigarh, India), who investigated the use of ^{68}Ga -pentixafor PET/CT, targeting the chemokine receptor CXCR4, in 25 lung cancer patients. They found that this tracer shows different degrees of uptake in biopsied lung carcinoma depending on the histology, and this result is correlated with CXCR4 expression. This novel PET tracer therefore seems to be useful for imaging and also for stratification, with prognostic implications.

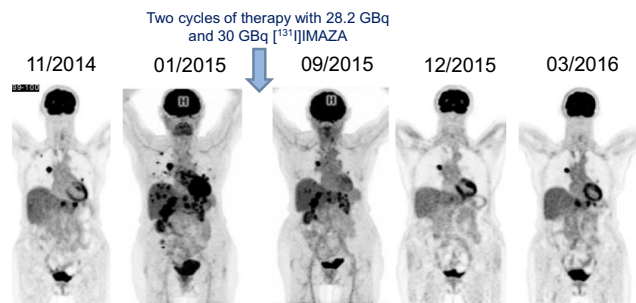
Paquette et al. [30] (Sherbrooke, Canada) reported the use of the combination of ^{18}F -FDG PET/CT and oestrogen receptor imaging with ^{18}F -4FMFES in patients with breast cancer, in particular in 13 patients with the luminal subtype, which is

Fig. 8 $^{123}\text{I}/^{131}\text{I}$ -IMAZA as a new theranostic tool in patients with advanced adrenocortical carcinoma [25]

Direct comparison of [^{123}I]IMTO and [^{123}I]IMAZA in an ACC-patient



[^{18}F]FDG-PET before and after [^{131}I]IMAZA-therapy



known to be difficult to assess using FDG alone. The combination of the two tracers overcame the limitation of FDG and achieved a better detection rate and diagnostic confidence, and more accurate staging. Medina Ornelas et al. [31] (Mexico City, Mexico) reported an interesting study of the use of ^{68}Ga -PSMA PET/CT in the evaluation of metastatic breast cancer. This tracer is a marker of angiogenesis in many types of solid tumour, and in this particular setting was found to be very accurate with expression of PSMA in metastatic sites observed in 100% of 11 patients. This result could help select patients with tumours with high expression of PSMA for targeted therapy with ^{177}Lu -PSMA.

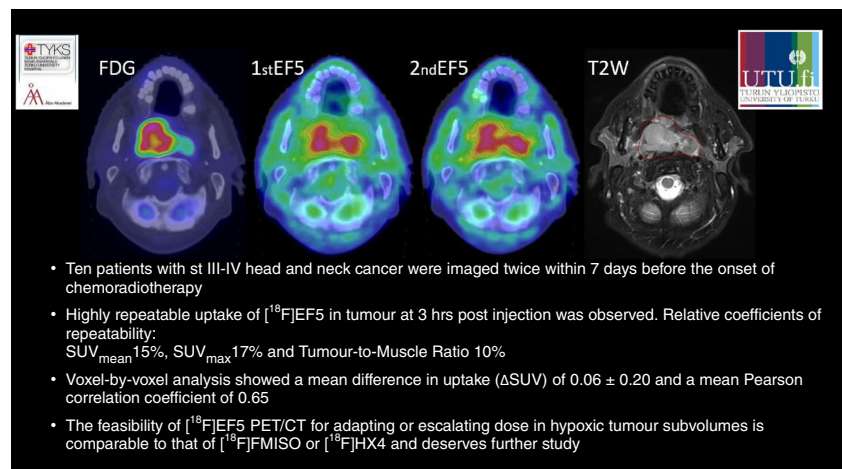
Silvoniemi et al. [32] (Turku, Finland), presented a study that focused on hypoxia imaging, because lack of oxygen contributes to RT resistance and more aggressive behaviour

of several types of cancer (Fig. 9). In 11 patients with head and neck cancer, PET/CT with ^{18}F -EF5 showed very good repeatability in the imaging of tumour hypoxia, which is very important especially for planning hypoxia-targeted treatment interventions.

Prostate

Prostate cancer imaging and therapy, driven by the success of PSMA ligands, has been a well-represented topic throughout all EANM meetings over recent years. The high interest and scientific focus is exemplified by a number of excellent presentations dealing with RT planning, use of PET/MRI and novel prostate cancer targeting radiopharmaceuticals.

Fig. 9 Repeatability of tumour hypoxia imaging using ^{18}F -EF5 PET/CT in head and neck cancer [32]



The first excellent study was presented by Calais et al. [33]. This was a collaboration between the US and Germany. The authors reviewed 252 patients with early biochemical relapse after prostatectomy who had been scanned with ^{68}Ga -PSMA-11 PET/CT to define the impact on RT planning (Fig. 10). ^{68}Ga -PSMA-11 PET/CT affected salvage radiation therapy planning in 55% of patients with early biochemical recurrence (PSA <1 ng/ml) after radical prostatectomy.

Schwarzenboeck et al. [34] (Rostock, Germany) analysed the use of ^{68}Ga -PSMA PET/CT in 38 patients with prostate cancer with a high risk or persistence or biochemical recurrence who were candidates for radiation therapy. In about 70% of these patients there was a clear direct impact on radiation therapy in terms of a change in the planning target volume and the addition of booster doses. Another interesting study was presented by Freitag et al. [35] (Munich, Germany) on the use of ^{18}F -PSMA-1007 PET/MRI combined with integrated multiparametric PET/MRI for imaging prostate cancer. This promising high-resolution coregistered PET/MRI protocol was completed in 1 h and was found to be feasible for the patients, and also appeared to result in fewer artefacts in the prostatic fossa than ^{68}Ga -PSMA 11 (due to low bladder accumulation of ^{18}F -PSMA-1007).

Schmidkonz et al. [36] (Erlangen, Germany) presented clinical data on the use of $^{99\text{m}}\text{Tc}$ -MIP-1404, a SPECT PSMA ligand, in prostate cancer patients. They enrolled 380 subjects and demonstrated high accuracy in detecting PSMA-positive lesions with a very good detection rate of 89% in patients with a PSA level ≥ 2 ng/ml and 40% in patients with a PSA level <2 ng/ml. Another example of outstanding work was presented by Haefliger et al. [37] (Lausanne, Switzerland). They compared the use of ^{18}F -choline (FCH) and ^{68}Ga -NODAGA-MJ9 (MJ9), a bombesin analogue and GRP receptor antagonist, for PET/CT imaging in 15 patients with histologically proven prostate cancer

with a mean Gleason score (GS) of 7 ± 1 (6–9) and a mean PSA level of 40 ± 73 ng/ml. MJ9 uptake was significantly higher in the prostate and lymph nodes than FCH uptake, while FCH uptake was significantly higher than MJ9 uptake in bone lesions, confirming the role of bombesin analogues in the initial staging of prostate cancer patients.

Batra et al. [38] (New York, USA) reported a study of the use of ^{89}Zr -df-IAB2M for PET/CT imaging of prostate cancer in a safe administration in nine men with a presurgery median PSA level of 8 ng/mL and a GS of ≥ 7 (range 6–9) in 89% of patients. IAB2M is an anti-PSMA minibody excreted via the hepatic route and in this phase 2a clinical trial showed excellent targeting of prostate cancer lesions and a very low urinary excretion in comparison with small-molecule PSMA ligands, with which bladder activity can compromise visualization in the prostatic bed.

Radionuclide therapy and dosimetry

Radionuclide therapy is becoming an ever more important part of nuclear medicine practice with optimization of therapies, alpha therapies showing excellent clinical results, and new therapeutic approaches entering clinical trials.

A multicentre study presented by Sharma et al. [39] (Hershey, USA), that pooled the data from three randomized studies, evaluated the efficacy and safety of adding selective internal radiation therapy (SIRT) using ^{90}Y resin microspheres to first-line mFOLFOX chemotherapy in 1,103 patients with liver metastases from colorectal cancer (mCRC). The addition of SIRT to first-line chemotherapy in patients with liver-only and liver-dominant mCRC led to an improvement in objective response rate ($p = 0.001$) and liver-specific progression-free survival ($p < 0.001$).

Fig. 10 ^{68}Ga -PSMA PET/CT mapping of early biochemical recurrence after primary surgery in 270 Patients with PSA < 1.0 ng/ml, indicating the high potential clinical impact of PSMA PET on salvage radiotherapy planning [33]

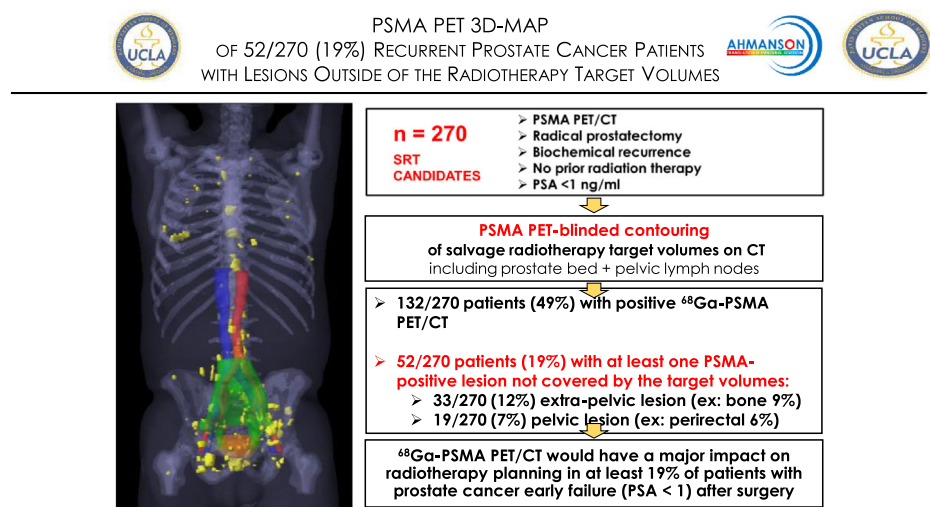
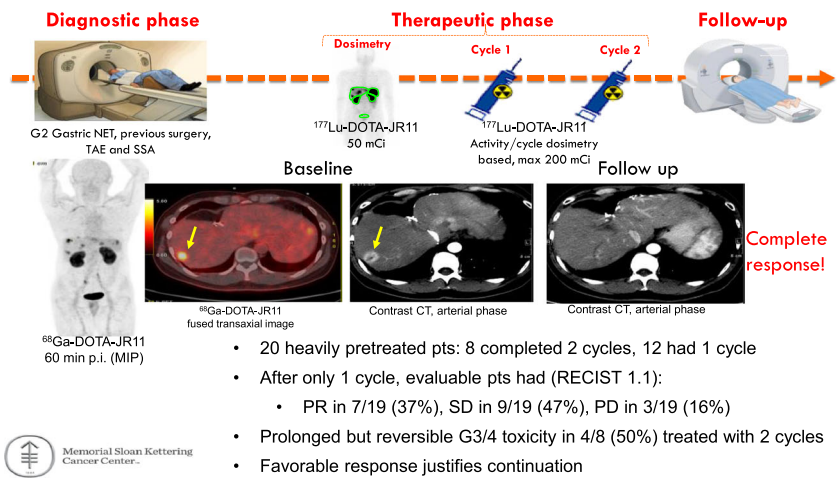


Fig. 11 Theranostic trial of somatostatin antagonists ⁶⁸Ga-OPS201 and ¹⁷⁷Lu-OPS201 in well-differentiated neuroendocrine tumours (NET), revealing high safety and excellent response rates even after only one cycle of therapy with ¹⁷⁷Lu-OPS201 [41]



Dizdarevic et al. [40] (Brighton, UK) presented a multicentre study involving various centres in Europe and the USA called REASSURE, an observational study on ²²³Ra, a targeted alpha therapy, in which 1,106 patients with metastatic castration-resistant prostate cancer (mCRPC) were enrolled. The interim analysis included data from 583 patients who had completed chemotherapy. Patients not previously treated with chemotherapy had less advanced disease at baseline than those who had received chemotherapy, and side effects (drug-related serious adverse events and treatment-emergent drug-related adverse events, most frequently gastrointestinal or haematological) were

more frequent in patients who had had prior chemotherapy, influencing the dropout rate of alpha therapy.

Reidy et al. [41] (New York, USA) reported a phase I theranostic trial evaluating the safety and radiation dosimetry of the SSTR2 antagonist JR11 in 20 patients with metastatic well-differentiated NET. The preliminary data showed that all tumours were detected by ⁶⁸Ga-OPS202 and therapy with ¹⁷⁷Lu-OPS201 was associated with a very impressive response in more than 80% of patients, even after one cycle, indicating that this theranostic combination is highly promising for imaging and therapy (Fig. 11).

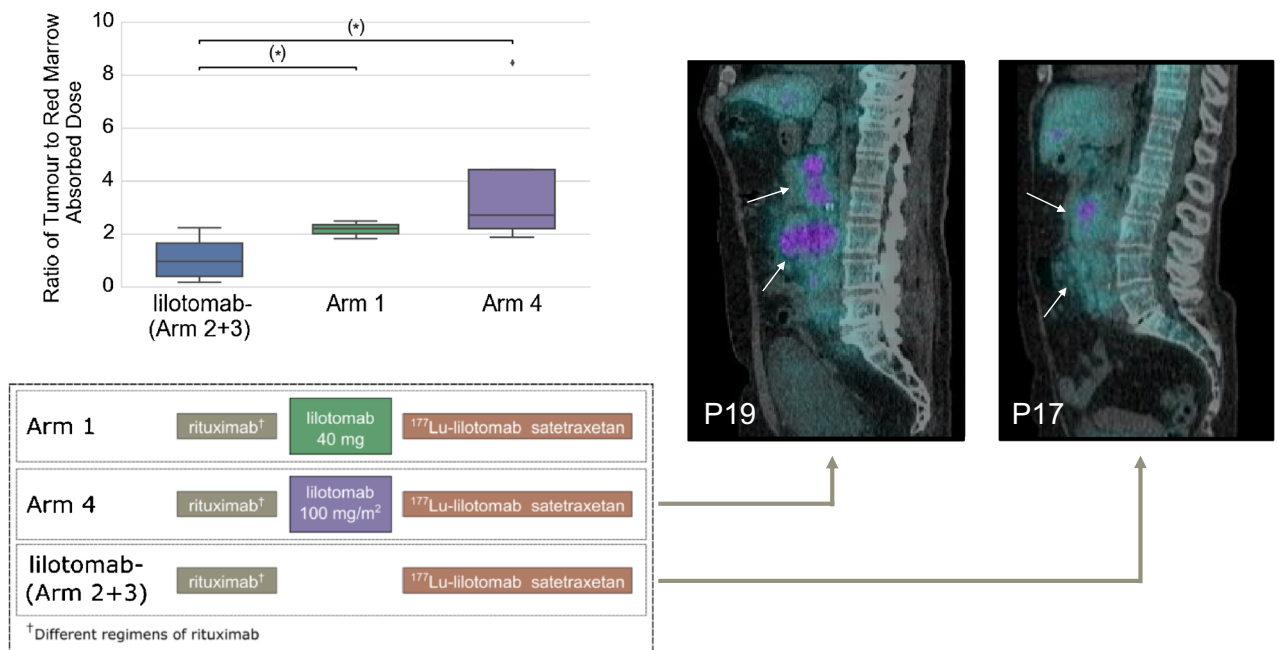
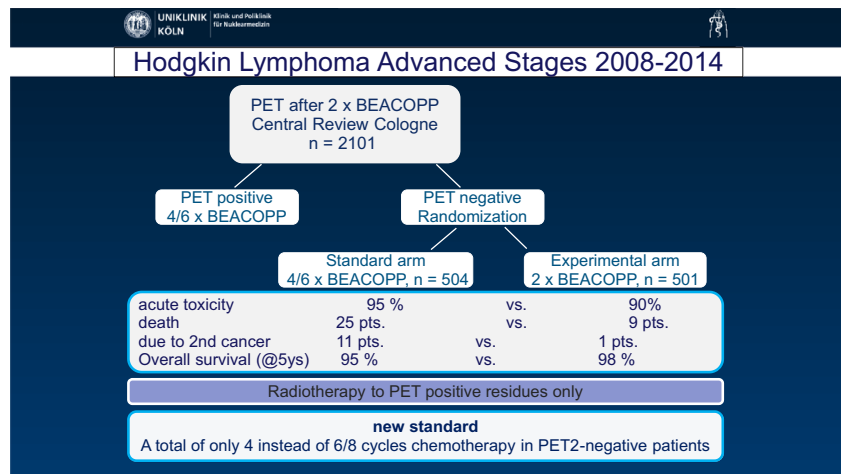


Fig. 12 Results of early phase clinical trials with ¹⁷⁷Lu-lilotomab satetraxetan targeting the CD37 antigen. Predosing with lilotomab prior to antibody–radionuclide conjugate therapy with ¹⁷⁷Lu-lilotomab

satetraxetan significantly increases the ratio of tumour to red marrow absorbed dose in patients with non-Hodgkin lymphoma [42]

Fig. 13 Treatment reduction in patients with advanced stage Hodgkin lymphoma and a negative interim PET scan. Final results of the international, randomized phase 3 HD18 trial by the German Hodgkin Study Group [47]



Blakkisrud et al. [42] (Oslo, Norway) reported the results of early phase clinical trials of the use ¹⁷⁷Lu-lilotomab satetraxetan (Fig. 12), a novel anti-CD37 antibody–radionuclide conjugate, for the treatment of non-Hodgkin lymphoma. Four different combinations of predosing and pretreatment were used. They enrolled 16 patients (pretreated with different regimens of rituximab) and there was a significantly higher red marrow dose in patients without predosing with anti-CD37 antibody radionuclide than in those with predosing ($p = 0.04$ and $p = 0.05$), indicating that predosing had a mitigating effect on red marrow absorbed dose, which would allow an increase in the tumour to red marrow absorbed dose ratio.

Scheidhauer et al. [43] (Munich, Germany) presented results of a pilot study in 12 patients evaluating the feasibility, safety and therapeutic efficacy of intravesical instillation of ²¹³Bi-anti-EGFR antibody in reducing recurrent bladder cancer. All patients showed excellent tolerance of this treatment without any side effects. SPECT/ CT monitoring clearly revealed the location of the ²¹³Bi-anti-EGFR antibody immunoconjugate in the bladder, and treatment resulted in a documented complete eradication of tumour cells in 25% of patients.

EANM 2017 specials

This category was introduced in this year’s Highlights Lecture to point out some particularly memorable submissions and presentations that stress the speciality of nuclear medicine in various aspects. We selected the most unusual abstracts, one abstract with the highest clinical impact, to show that science in the end improves patient care, and the radiopharmaceutical of the year indicating the potential of nuclear medicine to expand into new fields in years to come.

A very unusual abstract, was submitted by Amor-Coarasa et al. [44] (New York, USA). They evaluated the use of a 3D printer to print a synthesis module for radiopharmaceutical preparations. By this method they established a complex multistep radiosynthesis including distillation that was sufficiently robust and reliable for routine clinical use, and thereby reduced costs considerably. Another unusual abstract was a “naturalistic study”. This was a multicentre study presented by Guedj et al. [45] (France) evaluating the use of florbetaben PET/CT for amyloid plaque imaging as compared with other approaches in the work-up of dementia patients. The authors demonstrated that the

Fig. 14 An impressive case of treatment response after only one cycle of ²²⁵Ac-PSMA 617, the radiopharmaceutical of the year at EANM 2017. Data on therapeutic efficacy [48]

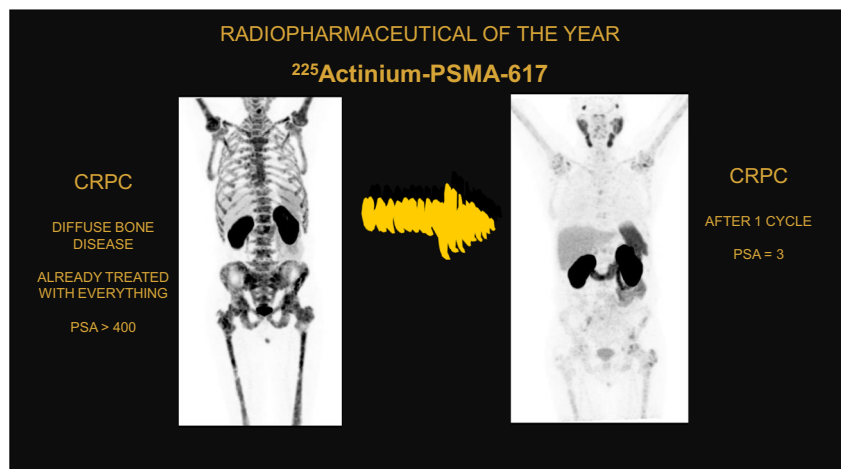


Table 1 Institutions submitting more than 20 abstracts to EANM 2017 (in alphabetical order according to the city of origin)

Centre	City	Country
Sant Orsola	Bologna	Italy
Universitätsklinik Essen	Essen	Germany
San Martino	Genova	Italy
University Medical Center (UMC)	Groningen	The Netherlands
Ludwig Maximilian University	Munich	Germany
Technische Universität	Munich	Germany
Radboud University	Nijmegen	The Netherlands
Uppsala University	Uppsala	Sweden
Medizinische Universität	Vienna	Austria

use of amyloid plaque imaging is superior to cerebrospinal fluid analysis, which is not only definitely more invasive and aggressive but is also less effective. Shiri et al. [46] presented a most unusual and scientifically highly interesting paper. These authors sought to predict lung metastases in patients with soft tissue sarcoma applying advanced machine learning to radiomic features. The unusual aspect, however, was the collaboration between Iranian universities and universities in the US, showing that science is above politics.

The abstract with the highest clinical impact was presented by Kobe et al. [47] (Cologne, Germany). This was a multicentre randomized double-blind study in which 2,101 patients with advanced stage Hodgkin lymphoma were enrolled. The aim of the study was to investigate the possibility of reducing the number of chemotherapy cycles in patients with a negative interim PET scan (Fig. 13). The study showed that the standard six or eight courses of eBEACOPP therapy can be reduced to only four courses in patients with a negative interim FDG PET scan without loss of lymphoma control. The improved tolerability of the de-escalated treatment strategy also resulted in a significant increase in overall survival (97.7%, $p = 0.006$)

^{225}Ac -PSMA 617 was chosen for the special award of the radiopharmaceutical of the year. The data for this promising compound for alpha radiation therapy in advanced prostate cancer were presented by Kratochwil et al. [48] (Heidelberg, Germany). They enrolled patients with castration-resistant prostate cancer already treated with every available drug and, for comparison, patients treated with this novel alpha therapy. This new therapeutic radiation-approach showed favourable low haematological toxicity and also remarkable antitumour activity in terms of objective response and progression-free survival (Fig. 14).

Conclusion

EANM 2017 again showed the high potential of nuclear medicine to be involved in the translation of basic science into clinical applications and in novel ways to diagnose and treat

an ever greater number of diseases. This could only have been achieved by great contributions from all over the world.

We particularly mention the most active institutions, all of them submitting more than 20 abstracts to the meeting (summarized in Table 1), and to acknowledge the authors most active during EANM 2017: Peter Bartenstein from Munich, Gianmario Sambuceti from Genova, and Markus Schwaiger from Munich.

We asked whether our scientific community at EANM 2017 could “Make Nuclear Medicine Great Again”; we can definitely answer “Yes We Can”.

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Compliance with ethical standards

Conflicts of interest None.

Ethical approval No human or animal studies by the authors are cited in this review.

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References

1. Cates CS, Levin CS. A promising PET detector design that achieves 100 ps FWHM coincidence time resolution. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):S302.
2. Vandenberghe S, Mikhalyova E, Brans B, Defrise M, Lahoutte T, Muylle K, et al. PET 20.0: a cost-efficient, 2mm spatial resolution total body PET with point sensitivity up to 22% and adaptive axial FOV of maximum 2.00m. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):S305.
3. Kaalep A, Sera T, Rijnsdorp S, Yaqub M, Talsma A, Lodge MA, et al. Feasibility of the state-of-the-art PET/CT system performance harmonization. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):S223.
4. Grueneisen J, Nensa F, Herrmann K, Bariye A, Forsting M, Umutlu L. Radiomics analysis predicts – and M-stage of primary cervical cancer using multiple PET/MR-derived quantitative features. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):S313.
5. Petoussi-Hens N, Ocampo Ramos J, Zankl M, Li W, Rühm W. Voxel based internal dosimetry of radiopharmaceuticals in diagnostic nuclear medicine. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):S356.
6. Rischpler C, Handwerker U, Dirschinger R, Kunze K, Kossmann H, van Marwick S, et al. Segmental comparison of myocardial

- inflammation, area at risk, edema and irreversible tissue damage after acute myocardial infarction. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):S189.
7. Varasteh Z, Bartels A, Mohanta S, Steinsiek A, Li Y, Braeuer M, et al. Non-invasive visualization of healing phase 2 after myocardial infarction (MI) using 68Ga- NOTA-anti-CD206-Nb: targeting mannose receptor (MR, CD206) on M2 macrophages. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):S370.
 8. Varasteh Z, Mohanta S, Li Y, López Armbruster N, Braeuer M, Nekolla S, et al. Non-invasive visualization of atherosclerotic plaques using 68Ga-NOTA-anti-CD206-nanobody: targeting mannose receptor (MR, CD206) on M2 macrophages. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):S371.
 9. Kim C, Lee J, Han Y, Chae S, Oh S, Lee S, et al. Diagnosis of deep venous thrombosis and pulmonary embolism using 18F-GP1 positron emission tomography: an exploratory open-label study. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):S373.
 10. Genovesi D, Vergaro G, Giorgetti A, Emdin M, Volpi E, Alduini S, et al. [18F]-Florbetaben PET/CT in cardiac amyloidosis: results from the FLORAMICAR study. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):S248.
 11. Amor-Coarasa A, Kelly JM, Ponnala S, Williams C, Vedvyas Y, Kim D, et al. 18F-RPS-544: an imaging agent targeting CXCR4. Imaging and biodistribution. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):S379.
 12. Baranski A, Schäfer M, Bauder-Wüst U, Roscher M, Schmidt J, Stenau E, et al. Dual-labeled PSMA-11 for PET/CT imaging and precise fluorescence guided intraoperative identification of prostate cancer. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):S235.
 13. Heskamp S, Molkenboer-Kueneen JDM, Sandker GW, Wierstra PJ, Bussink J, Boerman OC. Monitoring tumor PD-L1 expression with microSPECT/CT during radiotherapy. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):S144.
 14. Xu M, Han Y, Liu Z. Ab-1881, an anti-PDL1 immune checkpoint inhibitor serves as a theranostic agent for cancer immunotherapy. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):S143.
 15. D'Alessandria C, De Rose F, Kuhlmann MT, Braeuer M, Reder S, Braesh-Andersen S, et al. Immunotargeting of galectin-3 in thyroid orthotopic tumor models opens new challenges for thyroid cancer imaging and biological characterization in vivo. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):S141.
 16. Altai M, Westerlund K, Konijnenberg M, Mitran B, Oroujeni M, de Jong M, et al. Pretargeted radionuclide therapy of HER2-expressing SKOV-3 human xenografts using an Affibody molecule-based PNA-mediated pretargeting. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):S142.
 17. Bergmann R, Feldmann A, Schäfer M, Liolios C, Koristka S, Berndt N, et al. Low molecular weight target module for PET imaging and UniCAR T cell immunotherapeutic treatment of PSMA expressing tumors. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):S294.
 18. Hooshyar Yousefi B, Shi K, Reder S, Reder S, Herz M, Braeuer M, et al. First in vivo imaging and in vitro studies of 18F-DABTA in rat model with E46K alpha synuclein mutation. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):161.
 19. Kang Y, He B, Verma A, Henschliffé C, Kothari PJ, Schlyer D, et al. PET imaging of mGluR5 with [18F]FPEB in Parkinson's disease. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):194.
 20. Arlicot N, Vercouillie J, Mondon K, Gissot V, Maia S, Barantin L, et al. Validation of a reliable and convenient PET protocol for striatal dopaminergic dysfunction imaging using 18F-LBT-999. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):193.
 21. Unterrainer M, Fleischmann D, Lindner S, Brunegrab A, Vettermann F, Vomacka L, et al. TSPO-PET for high-grade glioma imaging using the novelligand [18F]GE-180 – first in human results in the course of radiotherapy. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):236.
 22. Seibyl J, Barret O, Stephens A, Madonia J, Alagille D, Mueller A, et al. Clinical evaluation of 18F-PI-2620, a next generation tau PET agent in subjects with Alzheimer's disease, progressive supranuclear palsy, and non-demented controls. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):252.
 23. Boss M Sr, Deden LN, Aarts EO, de Boer H, IMC J, Brom M, et al. Imaging beta cells in patients after Roux-en-Y gastric bypass (RYGB) surgery by 68Ga-NODAGA-exendin-4 PET/CT. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):162.
 24. Dubash SR, Merchant S, Mauri F, Kozlowski K, Lim A, Patel N, et al. Clinical translation of the caspase 3/7 specific PET radiotracer [18F]ICMT-11 for measuring chemotherapy induced apoptosis in breast and lung cancer. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):378.
 25. Schirbel A, Blümel C, Heinze B, Plaß A, Fuß CT, Megerle F, et al. [123/131I]IMAZA as a new theranostic tool in patients with advanced adrenocortical carcinoma. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):314.
 26. Rischpler C, Schlitter AM, Herz M, Yousefi B, von Werder A, Tauber R, et al. First experience using LMI1195 in patients with the suspicion of pheochromocytoma or paraganglioma. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):186.
 27. Zheng Y, Wang H, Cui X, Zhang L, Tan H, Yao S, et al. A proof-of-concept study of 68Ga-TATE-RGD PET/CT for dual-target imaging of somatostatin receptor and integrin $\alpha v \beta 3$ to detect lung cancer and neuroendocrine tumor in a single scan. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):149.
 28. Liu C, Choi W, Riyahi S, Lu W, Oh J, Deasy J, et al. Prediction of small early lung adenocarcinoma with aggressive histopathologic subtypes using PET and CT radiomic features. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):411.
 29. Watts A, Singh B, Chutani S, Dhanota N, Singh H, Basher R, et al. 68Ga-pentixafor PET/CT imaging targeting CXCR4 chemokine receptors: the first clinical experience in lung carcinoma subtypes. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):162.
 30. Paquette M, Lavallée E, Phoenix S, Senta H, Guérin B, van Lier JE, et al. Combined FDG and 4FMFES PET imaging in ER+ breast cancer patients for improved diagnostic and prognostic value. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):232.
 31. Medina Ornelas SS, García-Pérez FO. 68Ga-PSMA PET-CT in the evaluation of metastatic breast cancer. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):658.
 32. Silvoniemi A, Suilamo S, Laitinen T, Forsback S, Löyttyniemi E, Saunavaara V, et al. Repeatability of tumour hypoxia imaging using [18F]EF5 PET/CT in head and neck cancer. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):210.
 33. Calais J, Czernin J, Fendler WP, Herrmann K, Rauscher I, Hegemann N, et al. Impact of 68Ga-PSMA-11 PET/CT on salvage radiotherapy planning in post-prostatectomy patients with early biochemical recurrence. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):168.
 34. Schwarzenboeck SM, Schubert L, Rennau H, Kurth J, Krause BJ, Hildebrandt G. Impact of 68Ga-PSMA PET/CT on radiation treatment planning of prostate cancer. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):165.
 35. Freitag MT, Kesch C, Cardinale J, Flechsig P, Floca R, Eiber M, et al. Simultaneous whole-body 18F-PSMA-1007-PET/MRI with integrated high-resolution multiparametrical imaging of the prostatic fossa for comprehensive oncological staging of patients with prostate cancer. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):213.
 36. Schmidkonz C, Ritt P, Hollweg C, Beck M, Goetz TI, Sanders J, et al. Tc-99-MIP-1404 imaging for the detection of PSMA positive lesions. A pilot study in 380 patients with histologically

- confirmed prostate cancer. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):164.
37. Haeffliger L, Mitsakis P, Zilli T, Pozzessere C, Delage J, Maecke H, et al. Comparison study between 18F-choline (FCH) and 68Ga-NODAGA-MJ9 (MJ9, bombesin) PET-CT in prostate cancer initial staging. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):146.
 38. Batra JS, Jhanwar YS, Vallabhajosula S, Niaz MJ, Flynn T, Tagawa ST, et al. 89Zr-df-IAB2M for PET/CT imaging of prostate cancer. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):295.
 39. Sharma NK, Gibbs P, Van Hazel G, Heinemann V, Ricke J, Findlay MP, et al. The FOXFIRE/SIRFLOX/FOXFIRE-global randomized studies of first-line selective internal radiation therapy for metastatic colorectal cancer. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):696.
 40. Dizdarevic S, Meidahl Petersen P, Essler M, Versari A, Bourre J, La Fougère C, et al. First interim results of the radium-223 REASSURE observational study: analysis of patient characteristics and safety by prior use of chemotherapy. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):210.
 41. Reidy D, Pandit-Taskar N, Krebs S, O'Donoghue J, Raj N, Cruz E, et al. Somatostatin antagonist theranostic pair 68Ga-OPS202 and 177Lu-OPS201 for well-differentiated neuroendocrine tumors (NETs). *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):313.
 42. Blakkisrud J, Løndalen A, Dahle J, Martinsen AC, Holte H, Kolstad A, et al. Pre-dosing with lilotomab prior to antibody-radiionuclide conjugate therapy with 177Lu-lilotomab satetrexetan significantly increases the ratio of tumour to red marrow absorbed dose in non-Hodgkin lymphoma patients. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):129.
 43. Scheidhauer K, Seidl C, Bruchertseifer F, Apostolidis C, Autenrieth M, Kurtz F, et al. Bi-213-anti-EGFR-MAb therapy of recurrent bladder cancer – a pilot study. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):130.
 44. Amor-Coarasa A, Kelly JM, Kim D, Qu W, Kothari P, Babich JW. A 3D-printed automated dual reactor synthesizer for challenging multi-step 18F-fluorinations: testing and validation. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):273.
 45. Guedj E, Jonveaux T, Verger A, Krolak-Salmon P, Houzard C, Godefroy O, et al. Incremental value of 18F-florbetaben amyloid PET in the diagnostic work-up of most complex patients with dementia in France: a naturalistic study. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):169.
 46. Shiri I, Rahmim A, Abdollahi H, Geramifar P, Bitarafan- Rajabi A. Applying radiomics and machine learning on PET images to predict lung metastases in soft tissue sarcoma patients. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):741.
 47. Kobe C, Goergen H, Fuchs M, Eich HT, Baues C, Diehl V, et al. Treatment reduction in patients with advanced-stage Hodgkin lymphoma and negative interim FDG-PET: final results of the international, randomized, phase 3 HD18 trial by the German Hodgkin study group. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):312.
 48. Kratochwil C, Bruchertseifer F, Giesel FL, Apostolidis C, Haberkorn U, Morgenstern A. PSMA-targeting alpha-radiation therapy with 225Actinium-PSMA-617: dosimetry, toxicity and duration of tumor control. *Eur J Nucl Med Mol Imaging*. 2017;44(Suppl 2):163.