Evaluating Treatment Response of Radioembolization in Intermediate-Stage Hepatocellular Carcinoma Patients Using 18F-Fluoroethylcholine PET/CT

Markus Hartenbach1, Stefan Weber1, Nathalie L. Albert2, Sabrina Hartenbach3, Albert Hirt1, Mathias J. Zacherl2, Philipp M. Paprotka4, Reinhold Tiling2, Peter Bartenstein2, Marcus Hacker1, and Alexander R. Haug1,2

1Division of Nuclear Medicine, Department of Biomedical Imaging and Image Guided Therapy, Medical University of Vienna, Vienna, Austria; 2Department of Nuclear Medicine, Ludwig-Maximilians-University of Munich, Munich, Germany; 3Institute of Pathology, German Armed Forces Hospital Ulm, Ulm, Germany; and 4Department of Clinical Radiology, Ludwig-Maximilians-University of Munich, Munich, Germany

The aim of this study was to evaluate 18F-fluoroethylcholine PET/CT as a metabolic imaging technique for the assessment of treatment response to 90Y radioembolization in patients with locally advanced hepatocellular carcinoma (HCC). Methods: Thirty-four HCC patients undergoing 78 18F-fluoroethylcholine PET/CT scans were identified for this study. Patients with initial or follow-up metastatic disease (n = 9) were excluded at the time point of the metastatic occurrence as well as patients with negative α-fetoprotein (AFP; n = 1), resulting in 24 patients and 57 scans that were eligible. All patients were scheduled for radioembolization and underwent 1 pretherapeutic and at least 1 posttherapeutic 18F-fluoroethylcholine PET/CT scan. Volume-of-interest analysis and volume-of-interest subtractions were performed. Maximum, mean, and peak standardized uptake value (SUV) analysis was performed, and the total intrahepatic 18F-fluoroethylcholine positive tumor volume (FEC-PTV) and tumor-to-background ratio were assessed. Statistical analysis was performed using a decreasing AFP of at least 20% as a standard of reference for therapy response including receiver-operating-characteristic analyses as well as descriptive and correlation analyses and multiple logistic regression. Results: Fourteen follow-up examinations were categorized as responder and 19 follow-up examinations as nonresponder. Absolute AFP values did not correlate with SUV parameters (P = 0.055). In receiver-operating-characteristic analyses, the initial mean SUV, maximum SUV, and tumor-to-background ratio demonstrated the highest area under the curve, 0.84 (P = 0.009), 0.83 (P = 0.011), and 0.83 (P = 0.012), respectively, resulting in a positive prediction of 82%, 83%, and 91% at the respective cutoff points. When multiple logistic regression analysis was applied, this result was achieved in an area under the curve of 0.90 (P = 0.001), with a positive prediction of 94% and a sensitivity of 94%. The FEC-PTV did not reach significance in the presented dataset. Conclusion: 18F-fluoroethylcholine PET/CT demonstrates a high potential for follow-up assessment in the context of radioembolization in patients with locally advanced, but nonmetastatic, HCC and initially elevated AFP, possibly enabling early therapy monitoring independent of morphology.

Key Words: oncology; GI; PET/CT; radionuclide; therapy; FEC; PET; hepatocellular carcinoma; radioembolization; therapy

a subgroup analysis (16). Choline is transported into the cells via choline transporters or diffusion and phosphorylated by the choline kinase (17). Radiolabeled phosphocholine is then trapped in the cells and not further metabolized. Therefore, it can be used as a suitable surrogate for cell membrane biosynthesis.

According to the Barcelona Clinic Liver Cancer (BCLC) classification, liver-directed therapies such as transarterial chemoembolization are indicated for patients with local multinodular disease (BCLC stage B) (12). In these patients, who represent approximately 20% of HCC patients with a median survival time of 20 mo, radioembolization is also of increasing significance (18). For the purpose of our study, an advantage of this subgroup with advanced but liver-confined disease is the possibility to evaluate treatment-induced quantitative metabolic alterations of the liver lesions by PET imaging and to compare them with blood tumor markers, such as the AFP, not being affected by metastatic disease. Therefore, the assessment of local therapy response and the potential of 18F-fluoroethylcholine PET/CT can be evaluated more independently from RECIST or mRECIST.

These facts encouraged us in carrying out this study to evaluate pre- and posttherapeutic 18F-fluoroethylcholine PET/CT scans of patients with known HCC being scheduled for radioembolization at our clinic. The aim was to assess the potential of this metabolic hybrid imaging method for monitoring therapy response.

MATERIALS AND METHODS

This retrospective study was approved by the local ethics committee of the University of Munich, and the requirement to obtain informed consent was waived. Thirty-four patients with known HCC according to the EASL criteria or histology were identified for this analysis. All patients were scheduled for radioembolization and received in total 78 pre- and posttherapeutic 18F-fluoroethylcholine PET/CT scans. Nine patients presented initial or follow-up extrahepatic disease, not being categorized stage B anymore and therefore not suitable for this analysis. One patient was AFP-negative on initial and follow-up PET/CT scans. In total, 24 patients with 57 scans (24 before radioembolization and 33 after radioembolization) were eligible and analyzed. The interval for postradioembolization PET/CT examinations was 3 mo. Four patients underwent 3 PET/CT follow-up examinations, and 1 patient underwent 2 PET/CT scans after radioembolization. All additional follow-up PET/CT scans were compared with the initial prereadioembolization PET/CT scans (Fig. 1).

Before radioembolization, all patients underwent angiography to evaluate vascular anatomy and identify relevant aberrant vessels. When necessary, prophylactic embolization of these vessels was performed (19). SIR-Spheres (SIRTex Medical) were applied directly into the arterioles containing contrast agent (120 mL; Iopromide, Imeron 300; Bracco) was intravenously administered at a rate of 2.5 mL/s. Initiation of the CT scan was initiated 60 min after intravenous injection of 18F-fluoroethylcholine (3 MBq/kg of body weight). Emission data were reconstructed with attenuation correction on the basis of concurrent diagnostic CT. FEC was produced and delivered by PETNet GmbH.

For image analysis, Siemens Syngo TrueD workstations were used. Detailed volume-of-interest (VOI) analysis and subtractions were performed using Hermes Hybrid 3D (Hermes Medical Solutions). Liver VOIs for total liver volume (TLV) were drawn manually on CT images using region-of-interest interpolation. Maximum, mean, and peak standardized uptake value (SUVmax, SUVmean, and SUVpeak, respectively) analysis was performed with a 3-dimensional VOI and a 50% isocontour. The analysis of the intrahepatic 18F-fluoroethylcholine positive tumor volume (FEC-PTV) was threshold-based. The lower threshold was defined as being 50% higher than the SUVmax taken from a cubic VOI out of a non-HCC–affected area of the liver (defined by CT) (Fig. 2).

As long as only venous-phase CT scans were obtained in this PET/CT approach, other multiphase CT or multiphase MR images obtained in the posttherapeutic interval for 18F-fluoroethylcholine PET/CT follow-up scans was 12 ± 3 wk, starting after the second therapy in cases of dual-time-point radioembolization. Whole-body PET scans were acquired in 3-dimensional mode using one of two 64-slice PET/CT scanners (Biograph 64 TruePoint [Siemens Medical Solutions] or Discovery [GE Healthcare]). Before the CT scan, mean weight-adapted iodine-containing contrast agent (120 mL; Iopromide, Imeron 300; Bracco) was intravenously administered at a rate of 2.5 mL/s. Initiation of the CT scan was initiated 60 min after intravenous injection of 18F-fluoroethylcholine (3 MBq/kg of body weight). Emission data were reconstructed with attenuation correction on the basis of concurrent diagnostic CT. FEC was produced and delivered by PETNet GmbH.

For image analysis, Siemens Syngo TrueD workstations were used. Detailed volume-of-interest (VOI) analysis and subtractions were performed using Hermes Hybrid 3D (Hermes Medical Solutions). Liver VOIs for total liver volume (TLV) were drawn manually on CT images using region-of-interest interpolation. Maximum, mean, and peak standardized uptake value (SUVmax, SUVmean, and SUVpeak, respectively) analysis was performed with a 3-dimensional VOI and a 50% isocontour. The analysis of the intrahepatic 18F-fluoroethylcholine positive tumor volume (FEC-PTV) was threshold-based. The lower threshold was defined as being 50% higher than the SUVmax taken from a cubic VOI out of a non-HCC–affected area of the liver (defined by CT) (Fig. 2). As long as only venous-phase CT scans were obtained in this PET/CT approach, other multiphase CT or multiphase MR images obtained in the posttherapeutic interval for 18F-fluoroethylcholine PET/CT follow-up scans was 12 ± 3 wk, starting after the second therapy in cases of dual-time-point radioembolization. Whole-body PET scans were acquired in 3-dimensional mode using one of two 64-slice PET/CT scanners (Biograph 64 TruePoint [Siemens Medical Solutions] or Discovery [GE Healthcare]). Before the CT scan, mean weight-adapted iodine-
each patient before radioembolization were considered to confirm these nonaffected areas. The relationship between the FEC-PTV and the TLV, measured on CT images and defined as volumetric 18F-fluoroethylcholine tumor-to-liver ratio, was also determined according to the formula:

\[
\text{Volumetric FEC tumor-to-liver ratio} = \frac{\text{FEC-PTV}}{\text{TLV}}
\]

Additionally, tumor-to-background ratios (TBRs) were calculated. As far as the spleen is an organ with relatively high but homogeneous fluoroethylcholine uptake, we defined a cuboid VOI of the spleen (5-voxel side length) and calculated TBRs for SUV$_\text{max}$, SUV$_\text{mean}$, and SUV$_\text{peak}$ according to the formula:

\[
\text{TBR} = \frac{\text{SUV (tumor)}}{\text{SUV (spleen)}}
\]

where SUV is standardized uptake value. According to the $\Delta$AFP (AFP alteration between the PET/CT scans), the $\Delta$ of all PET/CT parameters was also determined, for example, the $\Delta$ of SUV$_\text{max}$ was calculated according to the formula:

\[
\Delta\text{SUV}_\text{max} = \text{SUV}_\text{max} \text{ postRE} - \text{SUV}_\text{max} \text{ preRE},
\]

where RE is radioembolization. Statistical analysis was performed using SigmaPlot 12.0 (Systat Tech Inc.). Data derived from SUV measurements were analyzed for correlation with AFP using Pearson correlation. Descriptive analysis comprised the determination of the respective mean, maximum, minimum, and median values. Patients were defined as therapy responder when AFP decreased more than 20% after radioembolization and as therapy nonresponder when the decrease was less than 20%. Sensitivity, specificity, and negative and positive prediction values were calculated. ROC cutoff values were derived by respective Youden indices. Values below cutoff indicate therapy response, defined as greater than 95% confidence intervals.

The multiple logistic regression analysis of these data resulted in the equation:

\[
\text{Logit}(P) = 2.96 \times \text{SUV}_\text{mean} - 0.95 \times \Delta\text{SUV}_\text{max} + 0.19 \times \Delta\text{SUV}_\text{maxTBR}
\]

Applying this equation to the present dataset resulted in an AUC of 0.90 ($P = 0.001$), with an improved positive prediction of 94% and an also increased sensitivity of 94% ($P < 0.001$; Fig. 4; Table 1). However, when comparing the AUC of SUV$_\text{mean}$ with the AUC of Logit(P), the level of significance was slightly not reached ($P = 0.073$).

### RESULTS

Of the 34 patients undergoing pre- and posttherapeutic 18F-fluoroethylcholine PET/CT, 11 had HCC proven by histology and 23 were diagnosed according to the EASL criteria. After exclusion of the 10 patients with metastases of negative AFP, 24 patients with 33 follow-up PET/CT scans were analyzed. Nineteen follow-up scans in patients who presented an increasing AFP were categorized as nonresponse. Fourteen follow-up scans with decreasing AFP were categorized as therapy response (Fig. 3). Two of the patients with an additional 2 PET/CT scans had an increasing AFP—1 patient had an initially decreasing and then again increasing AFP (one turned into reference ranges and remained normal), and the patient with 1 additional PET/CT scan had a decreased AFP on both scans. Absolute AFP values did not correlate with SUV parameters in Pearson correlation analysis ($P = 0.055$). When ROC analyses were applied, the initial pretherapeutic SUV$_\text{mean}$, SUV$_\text{max}$ (pre- to posttherapeutic), and SUV$_\text{max}$ (pre- to posttherapeutic) demonstrated the highest AUC (Fig. 4), with 0.84 ($P = 0.009$), 0.83 ($P = 0.011$), and 0.83 ($P = 0.012$), respectively, resulting in a positive prediction of therapy nonresponders of 82% ($P = 0.009$), 83% ($P = 0.002$), and 91% ($P = 0.002$) at the respective cutoff points. Results are presented in Table 1.

The multiple logistic regression analysis of these data resulted in the equation:

\[
\text{Logit}(P) = 2.96 \times \text{SUV}_\text{mean} - 0.95 \times \Delta\text{SUV}_\text{max} + 0.19 \times \Delta\text{SUV}_\text{maxTBR}
\]

where SUV is standardized uptake value. According to the $\Delta$AFP (AFP alteration between the PET/CT scans), the $\Delta$ of all PET/CT parameters was also determined, for example, the $\Delta$ of SUV$_\text{max}$ was calculated according to the equation:

\[
\Delta\text{SUV}_\text{max} = \text{SUV}_\text{max} \text{ postRE} - \text{SUV}_\text{max} \text{ preRE},
\]

where RE is radioembolization. Statistical analysis was performed using SigmaPlot 12.0 (Systat Tech Inc.). Data derived from SUV measurements were analyzed for correlation with AFP using Pearson correlation. Descriptive analysis comprised the determination of the respective mean, maximum, minimum, and median values. Patients were defined as therapy responder when AFP decreased more than 20% after radioembolization and as therapy nonresponder when the decrease was less than 20%. Sensitivity, specificity, and negative and positive prediction values were calculated. ROC cutoff values were derived by respective Youden indices. Values below cutoff indicate therapy response, defined as greater than 95% confidence intervals.

The multiple logistic regression analysis of these data resulted in the equation:

\[
\text{Logit}(P) = 2.96 \times \text{SUV}_\text{mean} - 0.95 \times \Delta\text{SUV}_\text{max} + 0.19 \times \Delta\text{SUV}_\text{maxTBR}
\]

Applying this equation to the present dataset resulted in an AUC of 0.90 ($P = 0.001$), with an improved positive prediction of 94% and an also increased sensitivity of 94% ($P < 0.001$; Fig. 4; Table 1). However, when comparing the AUC of SUV$_\text{mean}$ with the AUC of Logit(P), the level of significance was slightly not reached ($P = 0.073$).
The SUV\textsubscript{max} tumor-to-liver ratio in our cohort was 3.21 ± 1.62 (mean ± SD), with a minimum ratio of 1.67 (background SUV\textsubscript{max}, 9.24 ± 2.82). Nevertheless, the analysis of the volumetric data including the FEC-PTV as well as the volumetric \(^{18}\text{F}\)-fluoroethylcholine tumor-to-liver ratio did not reach significance in the presented dataset. The highest AUC was reached by the ΔFEC-PTV and the ΔFEC-to-liver-volume ratio (both AUC, 0.68; \(P = 0.175/0.164\); Fig. 5).

**DISCUSSION**

Therapy monitoring using hybrid imaging is, particularly in complex and systemic diseases, a preferable method, as this may lead to a complete whole-body status including both morphologic and metabolic information. In HCC, however, hybrid imaging results in several challenges for both methods (4,7,10). Assessing an early response to therapy after local or systemic treatment of HCC, however, according to morphologic aspects is limited (13); thus, the focus of our study was on the significance of the metabolic component of hybrid imaging using \(^{18}\text{F}\)-fluoroethylcholine PET/CT.

All consecutive patients with HCC who were scheduled for radioembolization were considered in this study. To generate a homogeneous patient collective and to have a reference parameter for treatment response independent from image morphology, patients with extrahepatic metastases were not included in the analysis. This allowed the use of AFP as outcome measures but required the exclusion of initially AFP-negative patients, because AFP showed high potential in treatment monitoring in earlier studies only insofar as it was initially positive (14,15). Although there is evidence in the literature using an AFP cutoff of 50% decrease for therapy responders with a baseline AFP 200 ng/mL (20), these studies starting with a large patient population had a large drop-out rate due to these criteria. In our cohort of patients in which an unestablished imaging method was applied, only 2 patients would have been regarded as therapy responders with these criteria applied. Therefore, we chose the AFP cutoff of a greater than 20% decrease for therapy responders, which is also discussed in the literature (20). Thus, we were able to evaluate \(^{18}\text{F}\)-fluoroethylcholine PET/CT for the assessment of intrahepatic HCC after radioembolization.

Our study revealed that a TBR, using the spleen as an organ with a relatively high choline uptake as background, in connection with the initial SUV\textsubscript{mean} indicated response to treatment with high accuracy. Although the total number of cases in this pilot study is low and the predictive power is therefore limited, we determined the positive and negative predictive values of the respective semi-quantitative methods because our collective with 38% responders after radioembolization has a representative distribution according to the literature (21). With a 94% positive prediction, the regression analysis showed high potential to evaluate a treatment failure, which could have a direct impact on an early therapy adaptation. Therefore, functional imaging using \(^{18}\text{F}\)-fluoroethylcholine PET/CT seems to be able to evaluate early local tumor alterations and might therefore be a reliable parameter for response assessment in those HCC patients in whom AFP cannot be used for treatment monitoring—that is, those with metastases or initially negative AFP. This assumption needs to be evaluated, but our initial results can provide the basis for further clinical studies (e.g., in patients with sorafenib therapy), even in advanced HCC, in order to reveal the importance of \(^{18}\text{F}\)-fluoroethylcholine PET/CT in the therapeutic management of HCC.

In contrast to the SUVs, the analysis performed on the FEC-PTV and the volumetric FEC tumor-to-liver ratio yielded no significant results in the assessment of the course of the disease. On the one hand, this might be due to the applied image analysis method, as we used a threshold of 50% increased FEC uptake over a defined background VOI in noncancerous liver tissue, which might also involve inflammatory and cirrhotic inhomogeneous areas in the volume calculation (10). On the other hand, the overall metabolic volume might remain largely unchanged after radioembolization while the maximum choline-metabolizing areas increase or decrease and therefore appear to be more responsible for the course of the disease. A further analysis of this inhomogeneity...
in the primary HCC lesion, with regard to both the metabolism and vascularization and the additional parameters of the MR imaging such as diffusion and perfusion, is the subject of current studies in our group. In the course of functional hybrid imaging using PET/MR imaging, these processes can be measured simultaneously in 1 examination and promise to improve the outcome of these patients significantly by early, noninvasive treatment monitoring.

Some limitations of our study beyond the AFP cutoff have to be mentioned. First, only patients without metastases were included in order to ensure the comparison with a reliable, not-imaging-based follow-up marker; therefore, additional studies are needed to assess the value of 18F-fluoroethylcholine PET/CT in those patients, as discussed above. Although the patient cohort has a representative responder-to-nonresponder distribution, the total number of responders is low and therefore limits the ability of comparing the sensitivities of the respective semiquantitative methods. Another limitation in the subanalysis of FEC-PTV is the choice of the HCC-nonaffected liver background VOI. Although it was confirmed by multiphase CT or MR imaging, a HCC involvement not detected by these methods might have been possible and could have affected this analysis, which was not significant in our patient cohort. This analysis might also be limited in smaller tumors with lower uptake as far as only areas with 18F-fluoroethylcholine uptake 50% higher than the background SUV max were considered. In our patient cohort with locally advanced tumor stages and a mean tumor-to-liver ratio of 3.21 ± 1.62, this method did not produce drop-outs. On the other hand, applying automated methods of drawing VOIs based on background thresholds, rather than manually drawn VOIs, enables reproducibility. However, as our data demonstrate that the alterations of tumor areas with the highest FEC uptake (applying a 50% iso-contour of the SUV max) are reliable surrogates for therapy response, the coverage of the exact extent of advanced HCCs might be of inferior value for therapy monitoring. Additionally, our results on metabolic response are not compared with RECIST or mRECIST.

The main reason for that limitation is the acquisition of only venous contrast-enhanced CT data in the applied PET/CT protocol and only limited availability of follow-up multiphase CT or MR imaging investigations, which were acquired in an acceptable time span close to the PET/CT follow-ups. However, this study aimed at evaluating the accuracy of metabolic imaging and not the comparison between 2 imaging modalities. Nevertheless, prospective studies assessing radiographic and metabolic response criteria simultaneously are highly demanded and should be performed, for example, using dedicated PET/MR imaging scanners.

In addition, this type of metabolic hybrid imaging harbors the advantage of a more reliable whole-body staging than CT or MR imaging alone, so distant metastases can be detected earlier, enabling an early multimodal therapeutic approach. 18F-fluoroethylcholine PET/CT, however, did already demonstrate promising results in the detection of metastatic disease of HCC (10,16).

CONCLUSION

In the selective group of patients with locally advanced, but nonmetastatic, HCC and initially elevated AFP, 18F-fluoroethylcholine PET/CT demonstrated a high accuracy for response assessment in the context of radioembolization. This result might allow for early monitoring of local therapies independently from morphologic imaging and even in patients in whom serum tumor markers cannot be used for local treatment evaluation (e.g., in the case of metastatic disease or initially negative AFP). To confirm our findings, prospective randomized studies with multiparametric functional hybrid imaging as a basis for decisions are needed.

DISCLOSURE

The costs of publication of this article were defrayed in part by the payment of page charges. Therefore, and solely to indicate this fact, this article is hereby marked “advertisement” in accordance with 18 USC section 1734. No potential conflict of interest relevant to this article was reported.
REFERENCES


Evaluating Treatment Response of Radioembolization in Intermediate-Stage Hepatocellular Carcinoma Patients Using $^{18}$F-Fluoroethylcholine PET/CT

Markus Hartenbach, Stefan Weber, Nathalie L. Albert, Sabrina Hartenbach, Albert Hirtl, Mathias J. Zacherl, Philipp M. Paprottka, Reinhold Tiling, Peter Bartenstein, Marcus Hacker and Alexander R. Haug

Published online: September 24, 2015.
Doi: 10.2967/jnumed.115.158758

This article and updated information are available at:
http://jnm.snmjournals.org/content/56/11/1661

Information about reproducing figures, tables, or other portions of this article can be found online at:
http://jnm.snmjournals.org/site/misc/permission.xhtml

Information about subscriptions to JNM can be found at:
http://jnm.snmjournals.org/site/subscriptions/online.xhtml