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Therapeutic implications of molecular imaging with PET in the combined modality treatment of lung cancer

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ABSTRACT

Molecular imaging with PET, and certainly integrated PET-CT, combining functional and anatomical imaging, has many potential advantages over anatomical imaging alone in the combined modality treatment of lung cancer. The aim of the current article is to review the available evidence regarding PET with FDG and other tracers in the combined modality treatment of locally advanced lung cancer. The following topics are addressed: tumor volume definition, outcome prediction and the added value of PET after therapy, and finally its clinical implications and future perspectives.

The additional value of FDG-PET in defining the primary tumor volume has been established, mainly in regions with atelectasis or post-treatment effects. Selective nodal irradiation (SNI) of FDG-PET positive nodal stations is the preferred treatment in NSCLC, being safe and leading to decreased normal tissue exposure, providing opportunities for dose escalation. First results in SCLC show similar results. FDG-uptake on the pre-treatment PET scan is of prognostic value. Data on the value of pre-treatment FDG-uptake to predict response to combined modality treatment are conflicting, but the limited data regarding early metabolic response during treatment do show predictive value. The FDG response after radical treatment is of prognostic significance. FDG-PET in the follow-up has potential benefit in NSCLC, while data in SCLC are lacking. Radiotherapy boosting of radioresistant areas identified with FDG-PET is subject of current research.

Tracers other than ¹⁸FDG are promising for treatment response assessment and the visualization of intratumor heterogeneity, but more research is needed before they can be clinically implemented. © 2011 Elsevier Ltd. All rights reserved.

Background

Lung cancer accounts for 219,000 new cancer cases and 159,000 deaths a year in the United States, representing 15% of cancer cases and 28% of cancer deaths in 2009.¹ As patients often present with primary irresectable disease, the majority of patients with localized disease is currently treated with multimodality treatment using a combination of surgery, chemotherapy (CTx), radiotherapy (RT) and targeted agents. Although survival has significantly improved with combined modality treatment, still, about one third of locally advanced lung cancer patients experience local failure as their first site of relapse.² Furthermore, these combined treatment strategies are often associated with dose limiting toxicity,

prohibiting further intensification of treatment. This could potentially be overcome by targeted antitumor therapy with increasingly conformal RT techniques and targeted agents. Furthermore, progress is made in strategies directed at individualization and early adaptation of therapy dependent on the treatment response, which may lead to optimization of the therapeutic ratio in each individual. After completion of curative treatment, improvement of outcome could be accomplished by an early detection of local progression, increasing the possibility for those patients to be offered salvage therapy.

With the introduction of these techniques, however, accurate definition of the tumor volume to be treated becomes increasingly important. This emphasizes the need for imaging techniques enabling accurate definition of the presence and extent of tumor before, during and after curative treatment in cancers of the respiratory tract. While CT and MRI are the most accurate imaging modalities with respect to anatomical information, they often lack the potential to distinguish between vital tumor and non-malignant tissue. Here, molecular imaging with positron emission tomography (PET), providing metabolic information, has additional





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value. Different radiopharmaceuticals have been evaluated for the imaging of malignant tumors, of which ¹⁸F-fluorodeoxyglucose (¹⁸F-FDG) is by far most commonly used. FDG-PET scanning utilizes the difference in accumulation of FDG between normal and cancerous tissues, based on an enhanced glucose metabolism in cancer cells. Other PET tracers, visualizing specific molecular pathways in tumors such as proliferation (e.g. ¹¹C-methionine, ¹¹C-choline, ¹⁸F-fluorothymidine) hypoxia (e.g. ¹⁸F-FMISO) or expression of certain receptors (Her2Neu, EGFR) are increasingly being used in the evaluation of malignancies.

Thus, metabolic imaging with PET, and certainly integrated PET-CT, combining functional and anatomical imaging, has many potential advantages over anatomical imaging alone in the combined modality treatment of lung cancer. PET using FDG is at present most widely applied in the clinical practice of non-small cell lung cancer (NSCLC). Its use in small cell lung cancer (SCLC) is rapidly emerging.

The aim of the current article is to review the available evidence on the use of PET in the combined modality treatment of locally advanced lung cancer. For NSCLC and SCLC separately, the role of PET imaging will be addressed with respect to the following topics:

- 1. Definition of the tumor volume to be treated, both with respect to the primary tumor and the locoregional lymph nodes
- 2. Outcome prediction on basis of PET before or early after the start of treatment, and the added value of PET after therapy

With regard to both these topics, the clinical implications of the use of PET are addressed, and future perspectives are provided.

Because ¹⁸F-FDG is by far most commonly applied in clinical practice, the majority of evidence comes from this tracer. Therefore, where "PET" is used in this article, this refers to "¹⁸F-FDG-PET", unless otherwise stated. Wherever other tracers have shown additional value, or are regarded as promising in the near future, they will be discussed.

Search strategy

A comprehensive literature search was conducted using the "Pubmed" database. Included search terms were: "Non small cell lung cancer", "Small cell lung cancer", "NSCLC", "SCLC", "Target volume definition", "Delineation", "Gross tumor volume (GTV)", "Clinical target volume (CTV)", "Tumor heterogeneity", "Selective nodal irradiation", "Prognostic value", "Outcome prediction", "Follow-up", "Combined modality treatment", "Chemotherapy", "Radiotherapy" or "Radiation" in combination with "PET", "Positron emission tomography" or "Molecular imaging".

Reference lists of relevant articles were searched for further studies.

Only publications in the English language and published online before February 1, 2010 were included.

Non-small-cell lung cancer (NSCLC)

NSCLC represents more than 80% of lung cancer cases.³ Combined chemoradiotherapy is the standard treatment for locally advanced (stage III), inoperable NSCLC.⁴ The added value of PET to select patients for combined modality treatment has been studied extensively^{5–7}, and it was shown that PET staging results in superior outcome due to stage migration: up to 30% of stage III patients are diagnosed with distant metastases.^{8,9} This clearly affects patient outcome as it withholds toxic therapy in individuals who will not benefit from it.

Below, we will discuss the role of PET in the RT planning and evaluation of combined chemoradiotherapy for stage III NSCLC.

Definition of the tumor volume to be treated

Primary tumor

Although FDG-PET has a high sensitivity for the detection of the primary tumor, an important drawback is its lack of anatomic detail, which limits its ability to define the exact tumor boundaries. The spatial resolution of current PET scanners is limited to 4–6 mm¹⁰, which is far lower than that of modern CT scanners, with a resolution down to 1 mm. There are cases, however, where anatomic imaging modalities such as CT are compromised in their ability to define the exact tumor border, such as in patients with atelectasis or tumors near the thoracic wall.¹¹ Multiple studies have shown a large interobserver variation in delineation of the tumor on CT.^{11,12} This interobserver variation is significantly diminished by using the information of a FDG-PET scan, co-registered with CT.^{13–16} Overall, volumes delineated using PET-CT are smaller.¹⁴ Differences between PET and CT were mainly found in the regions with atelectasis.^{14,17,18}

Various quantitative methods have been developed for automatic tumor delineation using PET instead of visual interpretation of the PET signal. The most straightforward method uses an absolute threshold of the standardized uptake value (SUV). The SUV_{max} threshold of 2.5 is often used for this purpose.¹⁹ An absolute threshold should be used with caution, however, as the SUV is associated with considerable variety due to both technical and biological factors.²⁰ An alternative method is the use of a relative threshold, e.g. a certain percentage of the SUV_{max}. Recently, more complex methods have been developed, including the application of an individualized threshold based on the source-to-background ratio (SBR) or the watershed clustering method.²¹⁻²³ An example of the difference in interobserver variation between manual and autocontour based delineation is provided in Fig. 1. Nestle et al. compared absolute (SUV_{max} \ge 2.5), relative (40% SUV_{max}) and individual (SBR algorithm) quantitative methods and a visual interpretation method with CT-volumes.²⁴ There were large differences in the resulting volumes, particularly in patients with a heterogeneous pattern of FDG-uptake.

Although autocontour based delineation methods have thus proven their utility in reducing interobserver variability, all quantitative methods harbor the risk of including metabolic active but not cancerous tissue in the GTV. Therefore, it has been suggested that those methods should be used complementary to visual interpretation, and not as a substitute for it.^{25,26}

Ideally, validation of the delineated tumor volume should be obtained by correlating it with the tumor volume at pathologic examination, being the gold standard. The currently available data are based on two-dimensional correlations in early stage disease.^{27–29} With the use of a relative threshold, a better correlation was found for CT than for PET (correlation coefficient 0.87 vs. 0.77).²⁷ Yu et al. found the best correlation with integrated PET-CT based on an absolute threshold (SUV_{max} ≥ 2.5).²⁸ A correlation coefficient of 0.90 was found between the maximal tumor diameter obtained with SBR-based autodelineation and pathology.²⁹ Promising attempts are made to develop a three-dimensional model, but results in large patient cohorts are to be awaited.^{30–32}

The methods described above are aimed at an accurate definition of the gross tumor volume (GTV) in order to ensure that this region is adequately covered by the RT treatment fields. Characteristics associated with radioresistance, such as hypoxia, cell density and proliferation, however, are known to be heterogeneous across the tumor.^{33–35} FDG-PET scans may allow the identification of therapy-resistant areas within the tumor. It would be logical to selectively boost the radioresistant areas, whilst decreasing the dose to the less resistant zones, resulting in higher tumor control with similar side effects.^{36–39} It has been demonstrated that regions with high FDG uptake prior to radiotherapy correspond well with the location of recurrent/persistent tumor after sequential



Fig. 1. Example of a manual (a) and autocontour based (b) delineation of the primary tumor For autocontouring, the SBR based method was used. Arrows indicate changes in interobserver variation in delineation between the two methods. Reprinted from: van Baardwijk A, Bosmans G, Boersma L, et al. PET-CT-based auto-contouring in non-small-cell lung cancer correlates with pathology and reduces interobserver variability in the delineation of the primary tumor and involved nodal volumes. *Int J Radiat Oncol Biol Phys* 2007;68:771–778. Copyright 2010, with permission from Elsevier.



Fig. 2. Correlation between pre-treatment high FDG-uptake areas and location of residual disease. Representative FDG-PET-CT images of three patients pre- and postradiotherapy. The light gray lines indicate the 50% SUV_{max} FDG high-uptake area pre-radiotherapy. The dark lines indicate the residual metabolic-active areas postradiotherapy, also transposed on the pre-radiotherapy scan. Visual evaluation shows a large correspondence between the residual areas post-radiotherapy with the high FDGuptake areas pre-radiotherapy. Reprinted from: Aerts HJ, van Baardwijk AA, Petit SF, et al. Identification of residual metabolic-active areas within individual NSCLC tumors using a pre-radiotherapy (18)Fluorodeoxyglucose-PET-CT scan. *Radiother Oncol* 2009;91:386–392. Copyright 2010, with permission from Elsevier.

chemo-radiotherapy or RT alone (Fig. 2).^{40,41} Furthermore, those regions remain stable during a course of RT (Fig. 3).⁴² Thus, selective boosting of areas of assumed radioresistance identified with FDG-PET before the start of RT appears to have a good rationale. It remains important, however, to consider that other biologic characteristics within the GTV, such as inflammation, may be associated with increased FDG-uptake as well, that are not directly related to increased radioresistance.⁴³Therefore, further research in this field is strongly encouraged.

The additional value of FDG-PET scanning in defining the primary tumor volume is thus beyond doubt. However, the drawbacks of FDG-PET should be kept in mind. Due to the poor resolution, blurring does occur, particularly at the tumor edges.²⁰ Those blurring effects at the tumor boundary are even more pronounced by motion artefacts. Although the long acquisition time of PET is disadvantageous with respect to defining an absolute tumor edge and quantitating metabolic activity, it may have additional value in determining the extent of tumor motion. The acquisition time of several minutes results in a tumor volume incorporating the averaged position of the tumor over multiple respiratory and cardiac cycles. In a phantom study, PET-based treatment volumes resulted in an adequate coverage of the tumor, while CT-based volumes harbored the risk of a geographical miss.⁴⁴ Respiratory gating or 4D imaging techniques allow the incorporation of the extent of tumor movement, while optimizing image quality and quantitation as the blurring effect is reduced.⁴⁵ Those techniques are presently being evaluated in clinical studies.^{45,46}

Microscopic disease extension

The poor spatial resolution of PET precludes a direct evaluation of the presence and extent of microscopic disease around the macroscopic tumor border. Definition of the area of potential microscopic spread in patients treated with chemoradiotherapy is important as this region should be covered in the radiation field. The only way to quantitate microscopic spread beyond the tumor border visible on imaging is by correlating imaging with the findings at pathologic examination. Until now, this correlation has only been performed between CT and pathology.^{47–49} Furthermore, no correction was applied for deformation of the lung lobe after surgery. Methods for the correlation of both PET and CT with pathology, which do take into account deformation, are under development.^{30,31} First results indicate an average microscopic spread in vivo of 9 mm³¹, suggesting that currently applied margins might be too small to cover microscopic disease.

Lymph nodes

Accurate identification of nodal metastases has become of particular importance since routine elective nodal irradiation, i.e. the prophylactic irradiation of clinically uninvolved lymph nodes, is no longer recommended in NSCLC.^{50,51} FDG-PET has a higher sensitivity and specificity for the detection of lymph node involvement in NSCLC than CT (sensitivity: 83% vs. 62%; specificity: 97% vs. 91%, respectively).⁵² Both PET- and CT-based selective irradiation of involved lymph nodes has proven its safety in NSCLC, with the occurrence of isolated nodal failures (INF) in less than 5% of



Fig. 3. Stability of high FDG-uptake areas during a course of RT PET-CT images of three patients before treatment (Day 0) and during treatment (Days 7 and 14). Lines indicate 60% of maximal standardized uptake value (SUV_{max}) threshold. Visual inspection showed that location of the hotspot remained at the same location during treatment; however, the volume of the hotspot changed. Reprinted from: Aerts HJ, Bosmans G, van Baardwijk AA, et al. Stability of 18F-deoxyglucose uptake locations within tumor during radiotherapy for NSCLC: a prospective study. *Int J Radiat Oncol Biol Phys* 2008;71:1402–1407. Copyright 2010, with permission from Elsevier.

patients.^{53–56} One study found INF in up to 15% with PET-based SNI.⁵⁷ However, the accuracy of the identification of lymph nodes in this study was questionable, as only visual interpretation of non-coregistered FDG-PET images was used. In general, the PET-based treatment volumes are smaller than CT-based volumes.^{58,59} Selective nodal irradiation (SNI) has shown not only to be safe, but also to result in a reduction of radiation fields based on CT, and even further based on FDG-PET.⁵⁵ A modeling study showed that treating only FDG-positive mediastinal areas decreases radiation exposure of the lungs and the esophagus sufficiently as to allow for radiation dose-escalation.^{55,59} An example of the difference resulting from RT planning with PET-CT compared to CT alone is illustrated in Fig. 4.

Although PET-defined SNI appears to be safe, ideally, pathological confirmation should be obtained. Pathological validation of the CT-and PET-based nodal treatment volumes was performed in 998 lymph nodal stations from 105 patients.⁶⁰ The coverage of all pathologic lymph nodes was 89% with PET-based treatment volumes compared to 75% with CT (p = 0.005). Nevertheless, a false negative rate with PET up to 14% has been reported in operable patients.⁶¹ A possible explanation for the low amount of isolated nodal failures is the incidental irradiation of clinically negative lymph node stations through coverage by the beam penumbra of conventional RT fields. The Michigan group showed that risk factors of nodal metastases, such as a large tumor size and central location, were associated with a considerable dose to the high-risk nodal regions.⁶² Therefore, caution is warranted with the application of new RT technologies, such as stereotactic body radiotherapy (SBRT), intensity modulated radiotherapy (IMRT) and particle therapy, as they are associated with a more conformal dose distribution.



Fig. 4. Example of the effects of RT planning with PET-CT compared to CT alone Projection of the planning target volume (PTV) of a 66-yearold female with a large cell carcinoma of the right lower lobe with pathological lymph nodes on CT scan in areas 4R and 3R and on FDG-PET scan in area 7. Although the lung exposure was lower with PET-CT than with CT (V20 25 vs. 30% and MLD 15.4 vs. 19.3 Gy, respectively), the esophageal exposure was higher with PET-CT because of the involvement of level 7 (MED 16.9 vs. 14.1 Gy, V55 18 vs. 4%, D_{max} 60.1 vs. 58.6 Gy, respectively, for PET-CT and CT). Reprinted from: De Ruyscher D, Wanders S, Minken A, et al. Effects of radiotherapy planning with a dedicated combined PET-CT-simulator of patients with non-small-cell lung cancer on dose limiting normal tissues and radiation dose-escalation: a planning study. *Radiother Oncol* 2005;77:5-10. Copyright 2010, with permission from Elsevier.

Different methods can be used for delineation of the involved lymph nodes on PET. Nestle et al. compared the nodal volumes resulting from visual delineation and absolute (SUV_{max} \ge 2.5),

relative (40% SUV_{max}) and individual (SBR-based) thresholds.⁶³ There were no clinically relevant differences in resulting volumes. SBR-based contouring of lymph nodes generally showed a good correlation with pathology.²⁹ Regardless of the delineation method, the question remains how to incorporate the lymph nodes in the treatment volume. There are no data available on the microscopic extension of disease outside lymph nodes. This residual uncertainty can be overcome by encompassing the whole anatomical mediastinal region in the treatment volume, as was done in the Maastricht studies described above.^{55,58,59}

As for the primary tumor, the exact anatomical localization of the mediastinal lymph nodes may be blurred because of respiratory motion. Ideally, individually determined margins should be applied to cover the lymph nodes in all respiratory phases, as there is a large intra- and inter-individual variation in lymph node motion, not related to the motion of the primary tumor.^{64–66}

Clinical implications and future perspectives

The incorporation of PET in RT planning has shown the potential for dose escalation through a reduction of the radiation fields, mainly because of avoidance of irradiating PET negative lymph nodes.^{58,59,67} With individualized radiation dose escalation based on normal tissue constraints, patients treated with sequential chemoradiation had survival rates comparable to results with concurrent chemoradiation schedules while less toxicity was observed.⁶⁸ These results imply that PET-based RT planning might ultimately lead to higher cure rates, and randomized prospective studies are warranted to investigate this further.

Further optimization of the treatment volume could be obtained by accurate definition of the appropriate margin around the delineated tumor to cover microscopic disease. This information should become available from pathology correlation studies.

Recently, an increasing tendency has emerged to move away from the concept of homogeneous irradiation. Studies have been performed to investigate the feasibility of selectively boosting areas with residual FDG-uptake after 40–60 Gy, with diverging results.^{69,70} Prospective trials are awaited to investigate whether radiation dose redistribution leads to better treatment outcome, for which preparations are currently being performed.^{37,71} In the studies mentioned before, ¹⁸FDG was used as a tracer for radioresistant areas. Other tracers, e.g. for hypoxia (¹⁸F-FMISO, ¹⁸F-HX4) or proliferation (¹⁸F-FLT) could be used complementary to or instead of FDG.^{72–76} These tracers deserve further investigation for this purpose.

Conclusion

FDG-PET has an important additional role to anatomic imaging in defining the primary tumor volume. Automatic delineation with adaptive techniques, such as SBR-based methods is to be preferred above absolute thresholding. Models are under development that correlate imaging findings with pathology in three dimensions. These could finally allow validation of different thresholds for SUV-based contouring and evaluation of microscopic spread and intra-tumor heterogeneity. Selective boosting of radioresistant areas identified with FDG-PET is subject of current research.

Selective irradiation of FDG-PET positive nodal stations is safe and leads to decreased normal tissue exposure, providing opportunities for dose escalation. With the increased use of more conformal radiation techniques, the safety of PET-based SNI should be re-evaluated. Disparities in treatment volumes resulting from different contouring methods are smaller than for the primary tumor.

Outcome prediction on basis of PET before or early after the start of treatment, and the added value of PET after therapy

Despite the improved outcome of inoperable stage III NSCLC achieved with combined chemoradiotherapy, the majority of inop-

erable stage III NSCLC patients still show disease progression after treatment, with 23–43% having an isolated local recurrence as their first site of progression.^{77–81} As the treatment is associated with considerable toxicity, it would be of great value to select patients before or early during treatment, with the highest probability to benefit from treatment and to adjust the treatment in the other patient group.

Early response assessment with conventional chest X-ray and CT is limited by their poor discriminating capacity between residual tumor and treatment induced changes.^{82,83} PET scanning allows for the assessment of changes in glucose consumption of the tumor during chemo- and radiotherapy. Several studies have shown a correlation between the SUV and tumor cell proliferation^{84–86}, supporting the hypothesis that an early change in FDG uptake has predictive value.

Prognostic value of pre-treatment PET

Most evidence regarding the prognostic value of pre-treatment PET comes from studies in heterogeneous patient populations with both early and late stage disease, treated with different modalities. A meta-analysis was performed within the IASLC lung cancer staging project. 11/13 eligible studies in stage I–IV NSCLC identified a high SUV as a poor prognostic factor for survival, with a combined HR for survival of 2.27 (95% CI: 1.43–3.04) for low vs. high SUV.⁸⁷ The threshold was variable between the studies, ranging from 5 to 20. Those differences are due to both technical and patient related factors, such as different scanners, time intervals between injection and scanning and fasting times. Furthermore, the relationship between SUV and prognosis is rather gradual than fixed at a single cut-off point.

Regarding the ability of PET to predict response to combined modality treatment, the first study in patients with advanced disease treated with (chemo-)radiotherapy showed a positive correlation between the tumor to muscle ratio (TMR) and response, but no significant correlation with outcome.⁸⁸ Two later studies revealed SUV_{mean} and SUV_{max} to be significantly associated with overall survival.^{89,90} In the first population, both tumor grade and UICC stage showed a stronger correlation with survival than the SUV⁸⁹, while in the second study, SUV_{max} was the strongest predictor.⁹⁰ By contrast, the most recent study in the largest cohort of stage III and IV NSCLC patients thus far (n = 214), did not show a significant relationship with survival.⁹¹ This study was not included in the IASLC meta-analysis mentioned above.

Outcome prediction on basis of early PET response during combined modality treatment

Because FDG is preferentially accumulated in viable tumor cells^{92,93}, FDG-PET imaging is an attractive method to visualize early treatment response. In advanced NSCLC, the predictive value of an early metabolic response to palliative chemotherapy, as well as to radical treatment with (chemo)radiotherapy has been evaluated. Prospective observational studies have consistently shown that in advanced NSCLC treated with palliative chemotherapy, the metabolic response after 1–3 cycles of chemotherapy is strongly correlated with outcome.^{94–97}

With respect to radical treatment, metabolic response to induction chemotherapy prior to radiotherapy or surgery in locally advanced NSCLC patients has been shown to correlate with outcome in multiple studies.^{95,98-100} One study, however, did not show a predictive value.¹⁰¹ While the evidence regarding the predictive value of a metabolic response to induction chemotherapy is abundant, far less is known about its value early during the course of radical treatment itself. Two studies investigated the predictive value of response during radiotherapy alone or chemoradiation. The first study was a pilot study in 15 patients treated with RT alone or chemoradiotherapy. A significant correlation was found between the response after 45 Gy of RT and the response 3 months after RT.¹⁰² The second study, investigating the predictive value of response during concurrent chemoradiotherapy¹⁰³, showed a significant difference in long-term survival between patients with and without a metabolic response after 3 weeks of concurrent chemoradiotherapy. An overview of the studies evaluating outcome prediction on basis of early PET response to combined modality treatment is provided in Table 1. A study evaluating response during radical RT revealed a large intra-patient heterogeneity in the evolution of SUV_{max} during and after radical RT.¹⁰⁴ Different time patterns were seen for responders and non-responders, but due to the limited patient numbers, the predictive value of the SUV_{max} changes could not be assessed.

To make FDG response assessment a valuable tool in routine clinical practice, a clear definition of response should be prescribed, as the intra-patient variability of repeated tumor SUVmeasurements is in the range of 10–15%.^{105–107} Furthermore, early response should be assessed at a fixed time interval. Ideally, the interval should be short enough to switch to a potentially more successful treatment as early as possible, but with a time interval sufficient to allow for a reliable response assessment. In 1999, the EORTC published consensus guidelines on which cut-off points should be used to define response at different time intervals¹⁰⁸. which are still widely applied in clinical practice. Weber et al. defined a metabolic response after the first cycle of chemotherapy as a decrease in FDG uptake of more than twice the standard deviation, calculated to be 20%. This definition correlated with final response according to RECIST, as well as with time to progression and overall survival.⁹⁶ The Melbourne group demonstrated that visual response assessment on PET with the use of standardized response criteria correlated with survival and was superior to response assessment on CT using WHO response criteria.¹⁰⁹ In 2009, the PET Response Criteria in Solid Tumours (PERCIST version 1.0) have been proposed resulting from a review of qualitative and quantitative methods of metabolic response assessment.¹¹⁰ PER-CIST recommends to correct SUV for lean body mass (SUL) as this accounts for variations due to differences in body composition.

A comparison between the EORTC criteria and PERCIST is provided in Table 2. Overall and most importantly, the same definition of response criteria should be used by different groups to be able to compare metabolic response studies across different centers.

Concerning the type of measurement, semiquantitative methods, such as the relative change in SUV, appear to perform equally well as more complex quantitative methods such as change in the net-influx constants (Ki) or metabolic rate of glucose (MRglu).^{95–97} This facilitates the use of early PET response for outcome prediction in daily clinical practice.

Added value of PET after combined modality treatment

The accuracy of PET after treatment is assumed to be lower than at initial staging because of therapy induced inflammatory and perfusion changes.¹¹¹ Nevertheless, PET still has a high accuracy in detecting recurrent lung cancer, with a sensitivity up to 98% and a specificity of 62–92%^{111–113}, and is more accurate than CT in the distinction of tumor from post-RT effects.^{109,114,115} Here, the added value of a post-treatment PET is addressed with regard to the prognostic value of a PET early after treatment and the role of PET in the follow-up after combined modality therapy.

We identified four studies addressing the prognostic value of PET after radical treatment in locally advanced NSCLC patients, consisting of (chemo-)RT.^{79,88,104,116} Four studies evaluated the predictive value of PET after induction chemoRT before surgery.^{100,117–119} Details of the studies evaluating the prognostic value after radical (chemo)RT and the predictive value after induction (chemo)RT are provided in Table 3. These studies were unambiguous in their conclusion that the FDG response after radical treatment has prognostic value. Mac Manus proved the superiority of PET response above CT. Response on PET and CT was identical in only 40% of patients. In multivariate analysis, only the PET response was significantly associated with survival.¹⁰⁹

Clear cut-offs should be prescribed to define the different prognostic subgroups. In the aforementioned studies, however, there is a large heterogeneity in the way FDG-uptake after therapy was measured. Some studies reported an absolute threshold post-treatment^{88,100,117,119}, while others stratified patients according to the relative change in SUV.^{79,104,109,116,118} In none of the studies a direct comparison was made between the different methods. Until more data are available we recommend the use of the EORTC criteria for PET response¹⁰⁸ for prognostic stratification, as the results of the larger studies are mainly based on these criteria.

Another important aspect is the timing of the PET-CT. The median time interval in the studies described above was 14–70 days. It is recommended to perform a PET-CT scan not earlier than 3– 6 months after treatment to avoid false positive results due to post-therapy inflammatory changes.^{111,120} The time interval should not be excessive either, as the final aim is to select patients for further therapy. Hicks et al. observed no confounding effect through post-RT inflammatory changes for response assessment with a PET-CT scan performed 70 days after radical RT.¹²¹ As different time-points have not been compared directly, we recommend the use of the time point 70 days post-treatment.

It should be noted that the results described above only apply for patients treated with conventional or hyperfractionated RT with or without chemotherapy. In hypofractionated SBRT, where 3–5 large fractions are applied, persistently elevated SUV_{max} of >3.5 have been described up to one year post treatment.^{122,123} These different findings may be explained by localized normal tissue changes induced by SBRT, such as segmental atelectasis or focal fibrosis, not distinguishable from persistent or recurrent tumor.

PET in the follow-up of NSCLC could improve outcome when progressive disease can be detected early enough to allow radical retreatment. There are no convincing data supporting that early

Table 1

Prediction of outcome on basis of early PET response to combined modality treatment

Study	Ν	Stage	Timepoint of PETscan	Radical treatment	Definition of cut-off	Predictive value
Hellwig (2004) ⁹⁹ Hoekstra (2005) ⁹⁵ Pottgen (2006) ¹⁰⁰ Kong (2007) ¹⁰²	47 47 50	IIB-III IIIA III I-III	After induction therapy ^a 1 and 3 Cycles 3 Cycles 45 Cyc	Surgery Surgery or RT ChemoRT ± Surgery (chemo)RT	SUV _{max} = 4 Residual MR _{glu} = 0.13 NR	OS: yes OS: yes Histopathologic tumor response: yes CMP/PMR 3 months after treatment: yes
(Decoster) 2008 ⁹⁸ Tanvetyanon (2008) ¹⁰¹	31 89	I–III III I–III	3 Cycles 2 Cycles	RT	CMR/FWR CMR 30% Decrease in SUV _{max}	PFS: yes OS: trend OS: no
Zhang (2009) ¹⁰³	46	III	40–50 Gy	ChemoRT	50% Decrease in SUV_{max}	OS: yes

N, number of patients; NR, not reported; CMR, complete metabolic response; PMR, partial metabolic response; OS, overall survival; PFS, progression free survival.

^a Data on number of cycles are not provided. Induction therapy consisted of chemotherapy only or chemotherapy followed by RT.

Table 2

Comparison of response criteria according to EORTC and PERCIST.

	EORTC	PERCIST
Progressive Metabolic Disease (PMD)	>25% Increase in SUV of tumor defined on pre-treatment scan, or >20% increase in the longest dimension of FDG- uptake, or Appearance of new FDG-uptake in metastatic lesions	>30% Increase in SUL peak and absolute increase of SUL units ≥0.8 from baseline scan in pattern typical of tumor and not of infection /treatment effect, or Visible increase in extent of FDG-uptake (> 75% increase in total lesion glycolysis), or Appearance of new FDG-avid lesions typical of cancer and not related to infection /treatment effect
Stable Metabolic Disease (SMD)	<25% Increase or <15% decrease in SUV of tumor defined on pre-treatment scan, and <20% increase in the longest dimension of FDG-uptake	No CMR, PMR or PMD
Partial Metabolic Response (PMR)	 >15% Decrease in SUV of tumor defined on pre-treatment scan (after 1 cycle) > 25% decrease in SUV of tumor defined on pre-treatment scan (after >1 cycle) 	>30% Decrease in SUL peak and absolute drop in SUL units ≥0.8 of the most intense lesion before and after treatment (not necessarily the same lesion) No new FDG-avid lesions typical of cancer
Complete Metabolic Response (CMR)	Complete resolution of FDG-uptake within tumor defined on pre-treatment scan, not distinguishable from surrounding normal tissue	Complete resolution of FDG-uptake within measurable target lesion, less than mean liver activity and indistinguishable from surrounding background blood- pool levels Disappearance of all other lesions to background blood-pool levels No new FDG-avid lesions typical of cancer

SUL: Standardized uptake value corrected for lean body mass.

Table 3

Value of post-treatment PET.

Study	Ν	Stage	Interval ^a	Radical treatment	Definition of threshold	Prognostic value	
Post (chemo)RT							
Ichiya (1996) ⁸⁸ Hebert (1996) ¹⁶⁶ Mac Manus (2005) ⁷⁹	20 12 88	III–IV NR I–III	<3 weeksks NR 70 days (median)	RT or chemoRT RT RT or chemoRT	TMR > 5 CR, visually interpreted CMR	RFS: yes Probable ^b OS: yes Distant M: yes Local failure: yes	
Van Baardwijk (2007) ^{29,93}	20	I–III	71 days (median)	RT or chemoRT	CMR or PMR	OS: yes	
After Neoadjuvant treatment							
Choi (1998) ¹¹⁷	29	IIB- IIIA	2 weeks	ChemoRT	$MR_{glu}\leqslant 0.040$	Predictive value pTCP ≥ 95%	
Ryu, 2002 ¹¹⁹	26	III	2 wks	ChemoRT	SUV _{mean>} 3	Pathological complete response:	
Pottgen (2006) ¹⁰⁰	43	III	NR ^c	ChemoRT	SUV _{maxpost} /SUV _{maxpre} ^d =0.38-0.55	yes Histopathologic tumor response: yes	
Eschmann (2007) ¹¹⁸	70	III	2 weeks	ChemoRT	CR, PR, SD, PD, visually interpreted > 80% decrease in $\mathrm{SUV}_{\mathrm{max}}$	OS: yes	

TMR, tumor to muscle ratio; RFS, relapse free survival; NR, not reported; OS, overall survival; CMR, complete metabolic response; PMR, partial metabolic response; MR_{elu}, metabolic rate of glucose; pTCP, probability of pathologic tumor control; CR, complete response; PR, partial response; SD, stable disease; PD, progressive disease. Between end of treatment and PETscan.

^b Patients with a complete response remained locally controlled, while 50% of the patients with a partial or no response showed progression.

^c Interval between the end of induction treatment was not reported. The interval between the start of induction chemotherapy and the PETscan was around 83 days.

^d Ratio between the SUV_{max} post induction chemoradiotherapy and the SUV_{max} after 3 cycles of induction chemotherapy.

detection of progression with chest X-ray or CT scan improves survival.^{82,124-126} This might be different for FDG-PET scanning, as PET is more accurate than CT in the distinction of tumor from post-RT effects^{109,114,127}, and is known to be prognostic for out-come.^{87,93,128–130} A prospective study was performed to evaluate whether PET-CT 3 months after therapy can detect potentially curable progression in locally advanced NSCLC¹³¹, which was the case in a small proportion (3%) of patients, who were all asymptomatic. An economic evaluation showed that a PET-CT scan 3 months after (chemo-)radiotherapy is potentially cost-effective, and is more cost-effective than CT alone.¹³² As the advantage was confined to the asymptomatic patients, a PET-CT scan in this group only is probably as effective and more cost-effective.

Other tracers

As response assessment early during and after therapy is complicated by the limited ability of FDG to discriminate between inflammation and tumor activity, it is worthwhile to investigate alternative tracers, corresponding more specifically with tumor proliferative activity.

The uptake of ¹⁸F-fluorothymidine (¹⁸F-FLT), a marker of DNA synthesis, has been correlated with tumor proliferative activity in various tumor sites including NSCLC.^{133–138} Recent (pre)clinical studies have demonstrated that FLT can detect changes in proliferation during and after irradiation in colorectal tumors and breast cancer cell lines.^{139–141} One pilot study in NSCLC has shown the feasibility of FLT to image proliferation during chemo-radiotherapy.⁷⁶

Amino-acid tracers theoretically have an advantage over FDG in that it more specifically accumulates in viable cancer cells.¹⁴² However, data on its usefulness in evaluating treatment response are scarce. One study compared ¹⁸F-FDG and the amino-acid tracer ¹¹C-methionine (¹¹C-Met) for evaluation of treatment response in lung cancer, but this study focussed on early stage patients treated with stereotactic RT.¹²³ In this study, FDG and Met showed an equal accumulation in inflammatory tissue, a finding supported by the results of other research groups.^{143–145 18}F-fluoromethyltyrosine (¹⁸F-FMT), another amino-acid tracer, has recently been put forward. Two animal studies showed a rapid response to antitumor therapy, and less accumulation in inflammatory cells.^{146,147}

Clinical implications and future perspectives

The clinical impact of patient selection before or early during treatment is beyond doubt, as it avoids ineffective treatment with the associated side-effects and enables alternative therapy in case of an inadequate early response. Additional research is needed to define clear cut-off points for FDG uptake to stratify patients into different treatment modalities. With regard to early response assessment, attention should be paid to other tracers, in particular proliferation markers, as they are less susceptible to uptake in inflammatory tissue.⁷⁶

A final option for improvement in outcome is the use of PET in the follow-up. Currently available data do show a potential benefit of PET in the follow-up compared to CT. Ideally, different follow-up strategies should be compared in a randomized controlled trial to provide definitive insight in the added value of PET in the followup of NSCLC patients after combined modality therapy.

Conclusion

In the overall NSCLC patient group, pre-treatment FDG-uptake is of prognostic value. Results on its ability to predict response to combined modality therapy in advanced stage NSCLC are conflicting. The most recent study in the largest patient cohort did not show a significant correlation with survival.

The predictive value of an early metabolic (FDG) response during induction chemotherapy has been established. Less is known about the predictive value of a metabolic response during radical (chemo-)RT, but the limited available data show a correlation with survival. The FDG response after radical treatment is of proven prognostic significance. A time interval of 70 days after the end of treatment is recommended for response assessment on basis of the EORTC criteria.

An FDG-PET scan in the follow-up after combined modality therapy can detect progression amenable for radical retreatment in a limited number of patients.

Tracers other than FDG are promising for treatment response assessment, but more research is needed before they can be clinically implemented.

Small cell lung cancer (SCLC)

SCLC is a tumor with a poor prognosis, characterized by a rapid growth rate. Traditionally, staging of those patients has been limited to the distinction between limited (LD) and extensive disease (ED).¹⁴⁸ Approximately 25% of patients present with limited-disease (LD), defined as disease confined to one hemithorax, including the mediastinum and bilateral supraclavicular fossae.^{149–151} Even in this patient group, surgery is rarely an option because of the advanced stage of locoregional disease. Concurrent chemo-radiation is currently the first choice treatment. Literature is sparse on the role of PET in LD-SCLC. The available literature suggests that FDG-PET has additional value above standard staging procedures in SCLC^{152–159}, with a reported sensitivity and specificity up to

100% and 95%.¹⁶⁰ Staging with PET can positively influence the outcome of chemoradiotherapy for LD-SCLC patients by means of stage migration. Upstaging from LD to ED by FDG-PET scanning occurs in 6-33%^{152-154,161-164} and downstaging in 3-40%.^{153,154,161,163}

Definition of the tumor volume to be treated

Primary tumor

In order to define the tumor volume. PET should be assessed for its ability to distinguish malignant from surrounding normal tissue. Studies addressing this issue are focussed on NSCLC.¹³⁻¹⁶ The same holds true for correlation studies with pathologv.^{28,30,31,48,49} Therefore, we can only assume that similar caveats apply as described above for NSCLC. In short, a major limitation of PET is the low spatial resolution. Hence, the major gain is to be expected in regions where anatomical imaging techniques lack the capacity to discriminate malignant from normal tissue, e.g. in areas with atelectasis. Another question refers to which method should be applied for PET-based tumor delineation. Again, comparison and validation of different methods has exclusively been performed in NSCLC.^{24,29} No conclusions can be drawn regarding which method is to be preferred, except that adaptive techniques are likely to be more accurate than the use of an absolute or relative SUV-based threshold.^{22,23} Obviously, blurring effects due to motion hinder exact tumor delineation. Respiration correlated imaging techniques have the potential to include individual tumor motion in the treatment volume, in conjunction with optimal image quality, as blurring effects are significantly reduced.^{45,46}

Microscopic disease extension

As described previously, the only way to define microscopic disease extension beyond the tumor border visible on imaging is to correlate imaging with pathology. There are no data on image correlation with pathology available for SCLC.

Lymph nodes

The available data suggest that an FDG-PET scan can identify metastases to regional lymph nodes in 14–25% of patients whose mediastinal CT scan is negative.^{152,156,163} With the high sensitivity and specificity of PET in SCLC, it is likely that the use of PET scans improves the coverage of mediastinal lymph node areas in LD-SCLC.

Until recently, few prospective data concerning selective nodal irradiation (SNI) in SCLC were available. A report from the International Atomic Energy Agency (IAEA) meeting emphasized the need for prospective clinical evidence regarding SNI in SCLC.¹⁶⁵ CTbased SNI resulted in an unacceptable amount (11%) of isolated nodal failures outside the treatment volume.¹⁶⁶ These findings imply that results on the safety of SNI in NSCLC cannot straightforwardly be extrapolated to SCLC. Since the publication of the IASLC report, two studies have become available evaluating FDG-PET-based SNI in SCLC. In a planning study, a difference in the treatment plan resulting from PET- and CT-based SNI was found in 24% of patients.¹⁶⁷ Radiation fields increased in 10% and decreased in 14% of patients, respectively. No significant changes in the radiation exposure of the normal tissue were observed. In the subsequent prospective study, 3% of the patients experienced an isolated nodal failure after a minimal follow-up of 18 months, comparable to results in NSCLC. A remarkably low percentage (12%) of grade III esophagitis was found, while this occurs in about 30% of patients receiving elective nodal irradiation or CT-based SNI.^{166,168} This finding deserves further investigation. The low rate of isolated nodal failures and toxicity thus supports the use of PET-based SNI in LD-SCLC.

A few points of caution should be taken into consideration. First, incidental irradiation of surrounding nodal stations might partially explain the low rates of isolated nodal failures with SNI. Therefore, results should be re-evaluated with the application of more conformal techniques (SBRT, IMRT, particle therapy). A second point of attention consists of the methods for target volume definition. In the available study, the mediastinal nodal regions involved on PET were included in the treatment field.¹⁶⁹ As our literature search did not yield any study evaluating autocontouring methods for lymph node delineation in SCLC, we recommend SNI of the whole mediastinal nodal station involved on PET.

Clinical implications and future perspectives

As in most SCLC cases, the bulk of disease is located in the hilar and mediastinal regions, reduction of the treatment volume can mainly be reached by omitting elective nodal irradiation. If the finding of low esophageal toxicity, as described in the first study with PET-based SNI¹⁶⁹ holds true, PET based SNI indeed provides opportunities for treatment intensification. With regard to RT planning, another point of consideration is the concept of subboosting areas of supposed radioresistance. FDG-PET, as well as PET with other tracers, could help to identify those regions within the tumor. Although this concept is readily evolving in NSCLC, no such trend is observed until now in SCLC. Although it is reasonable to assume that characteristics associated with radioresistance are also heterogeneous in SCLC^{33–35}, the distribution of the disease load in NSCLC is different from SCLC, as for most SCLC cases, the majority of the tumor load is found in the nodal stations. Studies on the evolution and stability of regions with high FDG-uptake in NSCLC are entirely focussed on the primary tumor, and no such information is available with respect to lymph nodes. These issues should be addressed when heterogeneous dose escalation is taken into consideration in SCLC.

Conclusion

There are no data available on the role of FDG-PET in defining the borders of the primary tumor. In contrast with CT-based SNI, first results indicate that SNI of FDG-PET positive nodal stations appears to be safe and results in remarkably limited toxicity. With the increased use of more conformal radiation techniques, the safety of PET-based SNI should be re-evaluated. It is recommended to encompass the whole anatomical mediastinal region containing FDG-positive nodes in the treatment volume.

Outcome prediction on basis of PET before or early after the start of treatment, and the prognostic value of PET after therapy

The majority of SCLC patients still shows disease progression short after the completion of chemoradiotherapy, with over 30% having an isolated local recurrence as their first site of progression.¹⁶⁸ Furthermore, the treatment is associated with considerable toxicity, with grade 3 esophagitis in up to 27% of patients.^{117,170–172} Therefore, the ability to predict the benefit from treatment would be of great clinical value. Recent data have made clear that the traditional staging system with two categories (limited and extensive disease) is on its own not an adequate predictor of survival and is not sufficient to stratify patients for the most optimal therapy.^{173,174} Since recently, it is recommended to use the TNM staging for SCLC, as it has proven to result in a better stratification of patients in prognostic subgroups.^{175,176} FDG uptake on PET before, during or after therapy could have a role as additional prognostic and predictive marker in SCLC.

Prognostic value of pre-treatment PET

Evidence concerning the prognostic value of FDG-uptake before treatment in SCLC is scarce. One study was identified that addressed this subject.¹⁷⁷ The majority of patients had LD and were treated with concurrent chemoradiotherapy. Overall, as well as

for the subgroup with LD, patients with a high SUV_{max} (i.e., higher than the median) had a significantly worse overall survival than patients with a low SUV_{max} (LD: 20.1 vs. 35.3 months). Three prognostic subgroups could be defined on basis of FDG-uptake and disease stage. Those results imply that different treatment strategies are required for LD with low and high SUV_{max} . Randomized clinical studies are warranted to answer whether FDG-uptake in combination with anatomical staging is predictive of outcome and can be used to select the appropriate therapy for each patient group.

Outcome prediction on basis of early PET response during combined modality treatment

The predictive ability of an FDG response early after the start of chemotherapy has been evaluated in two studies, both after one cycle of chemotherapy.^{178,179} However, patients with both LD and ED were investigated. Therefore, the results reflect the predictive value of an FDG response early during palliative chemotherapy or radical chemoradiotherapy. Furthermore, both studies used CT response after completion of therapy as a reference, and not survival. Both studies concluded that the metabolic response was correlated with the response according to RECIST.

Several important questions need to be addressed in future studies to make early response assessment with PET during treatment a valuable clinical tool. Those questions include the type of measurement, the definition for response, and the most optimal time interval for the measurement of early response. Regarding response criteria, the use of the EORTC recommendations¹⁰⁸, a 20% threshold⁹⁶, as well as the criteria for visual response assessment by MacManus are valid¹⁰⁹: Fischer et al. compared the visual method with the EORTC criteria in SCLC response evaluation after one cycle of chemotherapy and found no significant difference.¹⁷⁸ Regarding the type of measurement and the time interval, no separate data on SCLC are available. As long as no such data are available, the most practical alternative is to adhere to the NSCLC recommendations. Those are the use of relatively simple semiquantitative measurements such as SUV_{max}^{95-97} , and a time interval of 1–3 cycles of chemotherapy.^{95–97}

Caution is warranted, however, when projecting results obtained in NSCLC at SCLC. As mentioned before, those are two distinct types of disease with different clinical behavior. SCLC is characterized by a rapid response to chemo- and radiotherapy. Therefore, a response to therapy could be more rapidly visible on CT than it is in NSCLC, which might restrict the beneficial effect of PET. This hypothesis is supported by the study of Fischer et al., who found that early response assessment after one cycle of chemotherapy with CT and PET showed a comparable correlation with the final evaluation on basis of RECIST.¹⁷⁸

Added value of PET after combined modality treatment

Two retrospective studies evaluated the prognostic value of PET after treatment in SCLC patients^{157,180}, with one specifically aimed at LD.¹⁸⁰ The first study included both LD and ED, and both treated and untreated patients, with treated LD patients representing 50% of the study population. It is hard to draw separate conclusions on this group, but overall, there was a significant negative correlation between PET positivity or SUV_{max} and overall survival.¹⁵⁷

The study evaluating exclusively LD patients has some limitations: only 73% was treated with chemoradiotherapy, the remaining patients with palliative chemotherapy.¹⁸⁰ Furthermore, the time interval between the end of treatment and PET-scanning was variable (3–125 days). Finally, the definition of PET positivity was rather broad. With those limitations in mind, the study showed a significant difference in progression free survival between PET positive and negative patients, with a trend for overall survival. A prerequisite for PET in the follow-up to have a positive impact on the final outcome is that progression should be detected at a time that radical retreatment is still an option. The rapid growth rate and early dissemination make it less likely for progression to be detected in a "curable" stage in SCLC than in NSCLC. Although no studies have addressed the role of FDG-PET in the follow-up of SCLC, it can therefore be questioned whether PET scanning is advantageous with respect to outcome.

Clinical implications and future perspectives

It is obvious that the currently available data are insufficient to modify or adapt treatment on basis of pre-treatment FDG uptake or an early metabolic response in SCLC patients. Research in the field of early response assessment is of particular importance to define the additional benefit of PET above CT given the rapid response of SCLC to chemo- and radiotherapy.

Finally, the role of PET after combined modality therapy of LD-SCLC should be addressed. Given the early dissemination of SCLC, most benefit is to be expected with a tracer that allows response evaluation early after treatment. Given the high uptake in inflammatory regions, FDG might not be ideal for this purpose. Therefore, other tracers should be evaluated.

Conclusion

Studies evaluating the prognostic value of PET, its ability to predict response to combined modality treatment and the added value of PET after treatment in SCLC are scarce. Available results mainly come from studies in patients with both limited and extensive disease. Overall, results do show some predictive value of an FDG response before and during therapy, as well as prognostic value of FDG uptake after treatment. No studies have evaluated the impact of PET in the follow-up of SCLC.

General conclusions

Molecular imaging with PET, using different tracers, has the potential to distinguish between vital tumor and non-malignant tissue and to identify intra-tumor characteristics. The additional value of FDG-PET in defining the primary tumor volume has been established, mainly in regions with atelectasis or post-treatment effects. Three dimensional models that correlate imaging findings with pathology are being developed for NSCLC, which could allow validation of different thresholds for SUV-based contouring, evaluation of microscopic spread and intra-tumor heterogeneity. FDG-PET has the ability to identify regions within the tumor that are associated with radioresistance, and it has been proved that these regions remain stable during a radiotherapy course. Therefore, boosting of radioresistant areas identified with FDG-PET appears to be feasible and is subject of current research. Selective irradiation of FDG-PET positive nodal stations in NSCLC is safe and leads to decreased normal tissue exposure, providing opportunities for dose escalation. For this reason, it is the preferred treatment in NSCLC. First results in SCLC suggest that the same holds true for SCLC. Data on the predictive value of pre-treatment FDG-uptake and an early metabolic response during combined modality treatment are conflicting and limited, respectively. The FDG response after radical treatment is of prognostic significance. A time interval of 70 days between end of treatment and PET scanning is recommended for response evaluation in NSCLC. A PET scan in the follow-up of NSCLC potentially improves survival through the detection of progression with radical treatment options. Data on its value in the follow-up of SCLC are lacking.

Tracers other than ¹⁸FDG are promising for treatment response assessment and the visualization of intra-tumor heterogeneity, but more research is needed before they can be clinically implemented.

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