Consensus Recommendations on the Use of $^{18}$F-FDG PET/CT in Lung Disease

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PET with $^{18}$F-FDG has been increasingly applied, predominantly in the research setting, to study drug effects and pulmonary biology and to monitor disease progression and treatment outcomes in lung diseases that interfere with gas exchange through alterations of the pulmonary parenchyma, airways, or vasculature. To date, however, there are no widely accepted standard acquisition protocols or imaging data analysis methods for pulmonary $^{18}$F-FDG PET/CT in these diseases, resulting in disparate approaches. Hence, comparison of data across the literature is challenging. To help harmonize the acquisition and analysis and promote reproducibility, we collated detailed acquisition protocols and analysis methods from 7 PET centers. From this information and our discussions, we reached the consensus recommendations given here on patient preparation, choice of dynamic versus static imaging, image reconstruction, and image analysis reporting.

Key Words: pulmonary imaging; FDG; PET/CT; lung disease

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ET has been explored in lung diseases, mostly in a research setting, as an imaging indicator of molecular changes for monitoring disease progression and treatment effects. The pathogenesis of these diseases leads to reduced gas exchange through alterations of the pulmonary parenchyma, airways, or vasculature ($1–4$). However, the development of effective therapies remains disappointingly slow, hampered by the limited ability to quantify molecular changes in the lungs and assess drug binding and activity. $^{18}$F-FDG is the most widely available and commonly applied PET tracer and is predominantly used to assess lung inflammation and fibrosis-related processes for both research and clinical purposes. New molecularly targeted PET tracers are also being developed to support the respiratory drug development process, with a recent publication demonstrating the additional potential for PET imaging to assess drug target engagement in the lungs ($5$). Therefore, we anticipate that the application of PET imaging in lung diseases will continue to grow across multiple centers.

In light of this growing use, standardization of acquisition protocols and analysis methods will facilitate data comparisons from multicenter studies and across the literature. To date, a range of analysis methods has been applied to disparately acquired static and dynamic datasets in studies of lung disease and related processes ($6–13$). Given that the lungs uniquely contain varying amounts of air depending on the disease, are the source of respiratory motion, and have a relatively high fractional blood volume, special considerations are needed when applying quantitative imaging approaches. Because of the variation in protocols among centers, however, comparability of the results across centers is somewhat limited.
Our group has previously published a summary of the conceptual approaches for lung PET imaging and highlighted some of the issues with clinical and research applications (2). To build on these applications, ensure comparability of data across centers, and enable cross-center data comparisons, we identified a need for harmonized protocols. Therefore, we convened representatives from all academic and commercial centers active in PET lung imaging to develop and present consensus recommendations for patient preparation, scanning protocol design, and imaging data analysis. The primary aim is to improve standardization to support uniform data collection and interpretation, thus improving the potential to compare results and pool data across research studies and enable advancement of the field. Moreover, it is hoped that a better understanding of the origin of the PET signal, gained from studies using dynamic acquisition and complex analyses, will enable the development of simpler static measurement protocols that can be more readily applied in clinical practice. Our consensus recommendations are timely given the current higher level of attention on uniform protocols to enhance the reproducibility of quantitative PET, as evidenced by recently published consensus papers on cancer and neuroimaging (14–17).

The acquisition protocols and image analysis details (Supplemental Tables 1 and 2; supplemental materials are available at http://jn.nmijournals.org) were collated from 7 participating centers that have conducted PET studies on lung diseases (Cambridge University Hospitals NHS Foundation Trust and University of Cambridge, Invicro, Massachusetts General Hospital, University College London, University Hospitals Coventry and Warwickshire, University of Edinburgh, and Washington University in St Louis; Supplemental Table 3). The acquisition protocol information included patient preparation, image acquisition, and image reconstruction. The analysis details included the whole and regional lung delineation methods and imaging endpoint derivations from both static and dynamic PET acquisitions. Between March 2018 and August 2019, the collated information was reviewed and the consensus recommendations developed over the course of 3 meetings that we attended in person or via teleconference. The collated details of the acquisition protocols were reviewed by all of us. Protocol differences among the centers were discussed and resolved to produce the consensus recommendations. When considering data analysis details, it was clear from the start that the desired imaging endpoints would vary among studies depending on the individual study objectives. Therefore, we defined minimum reporting requirements with a sufficient level of detail to enable comparison of published studies from different centers and to improve the reproducibility of analyses. All authors reviewed and approved the following recommendations.

**PATIENT PREPARATION**

**Fasting Period**

To minimize glucose-related inhibition of 18F-FDG uptake, only plain water should be consumed during a fasting period lasting a minimum of 6 h before scanning. This is also the recommendation of the Quantitative Imaging Biomarkers Alliance for oncologic PET imaging (16).

**Blood Glucose Level**

Among the protocols collated, the minimum acceptable blood glucose level before scanning varied. We agreed that a lower limit of 4 mmol/L would help avoid requiring glucose administration for preventing hypoglycemia that might interfere with the scan results. The European Association of Nuclear Medicine guidelines for tumor imaging (14) recommend an upper limit of 11 mmol/L for clinical studies. However, because medication such as steroids and other factors can influence glucose levels, we felt this should be a guide and that upper limits should be determined on a per-study basis.

Similarly, regarding the fasting period and acceptable blood glucose and insulin levels for diabetic patients, there were insufficient data to allow us to make a firm recommendation. We agreed that it would be reasonable to consider excluding patients with diabetes from small studies to minimize confounding factors.

**PATIENT POSITIONING AND COMFORT**

For pulmonary scanning, arm placement within the field of view can cause CT attenuation artifacts that may compromise the accuracy of the PET data. Therefore, having patients’ arms above their head for the duration of the scan is preferred. For patients who cannot tolerate this position, ensuring that the arms fit completely within the CT field of view consistently is recommended to reduce inaccurate attenuation correction across the chest when comparing studies. A maximum body mass index of 35 kg/m² is generally suggested to minimize both variability in positioning of patients within the scanner and body habitus effects. We also recommend that patients with implants that are within the chest field of view, such as pacemakers, be excluded from studies to avoid extreme artifacts.

Prolonged periods of lying down may be difficult for patients with lung diseases and may result in gross movement, causing attenuation correction errors. Measures should be taken to maximize comfort during scanning, including allowing a continuous supply of oxygen when needed, placing a vacuum bag filled with pillows or memory foam padding under the arms to provide support and improve blood circulation, using knee wedges to increase back comfort, and applying blankets for warmth (4). Appropriate padding of pressure points and avoidance of joint overstretching are particularly important for sedated, mechanically ventilated patients. Providing breaks during a dynamic PET scan for patients to sit or stand may be an option but will require rigorous realignment of each scan portion to produce a contiguous time–activity curve.

**ADMINISTERED 18F-FDG ACTIVITY**

18F-FDG doses used for clinical oncologic scans as outlined in the European Association of Nuclear Medicine guidelines (14) are sufficient for a static clinical scan in most lung diseases. Since many factors determine the choice of injected activity, such as static or dynamic image acquisition, frame duration, desired

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**NOTEWORTHY**

- This consensus statement provides recommendations for 18F-FDG PET lung imaging protocols, image analysis, and reporting to facilitate comparison of data acquired at different centers.
- Corrections for the effects of air (air fraction correction) and blood feature prominently in the quantitative analysis methods that are proposed to improve lung tissue–specific 18F-FDG quantification.
- Novel reconstruction methods and other approaches may help overcome the challenge of minimizing the effects of respiratory motion on quantification.
analysis endpoints, and standard practice at the study location (supplemental materials), no recommendation can be made to fit all studies. The introduction of more sensitive PET scanners could also affect the choice of injected dose. For research studies, the minimum activity that provides sufficient image quality to meet study objectives should be used (18), and we recommend that this activity be determined on a per-study basis.

### Table 1
Recommendations on Patient Preparation, Acquisition Protocols, and Image Reconstruction

<table>
<thead>
<tr>
<th>Category</th>
<th>Recommendation</th>
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<tbody>
<tr>
<td>Patient preparation</td>
<td></td>
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<tr>
<td>Fasting period</td>
<td>Minimum of 6 h before scanning, with plain water allowed</td>
</tr>
<tr>
<td>Blood glucose level</td>
<td>Minimum of 4 mmol/L, with upper limit determined by study objectives</td>
</tr>
<tr>
<td>Patient positioning</td>
<td>Arms above head when possible, with maximum body mass index of 35 kg/m² allowed for consistent arm positioning</td>
</tr>
<tr>
<td>Administered ¹⁸F-FDG activity and injection duration</td>
<td>Activity no more than needed for sufficient image quality to meet study objectives, with injection duration documented and kept consistent for all patients</td>
</tr>
<tr>
<td>Acquisition protocol</td>
<td></td>
</tr>
<tr>
<td>Static vs. dynamic</td>
<td>Static scans acceptable if compatible with study objectives; dynamic scans recommended when tracer quantification for specific lung compartments is needed; rationale for chosen acquisition method reported</td>
</tr>
<tr>
<td>Respiratory gating</td>
<td>End-expiration gating results reported when used; list-mode data stored to allow future reprocessing as techniques improve</td>
</tr>
<tr>
<td>Accounting for respiratory motion</td>
<td>Breath-hold at end-expiration or mid-expiration most frequently used to match lung volumes for CT and PET; prescan coaching of breathing instructions recommended (coaching patient on breathing instructions for attenuation correction CT can minimize most respiratory motion errors); exploration of approaches such as cine-CT</td>
</tr>
<tr>
<td>Duration and time frame</td>
<td></td>
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<tr>
<td>Static imaging</td>
<td>Starting at 60 min after injection</td>
</tr>
<tr>
<td>Dynamic imaging</td>
<td>45- to 90-min acquisition starting immediately after injection; 60-min scan is routinely tolerated (breaks may be required for improved patient tolerance)</td>
</tr>
<tr>
<td>Time frames for typical 60-min dynamic acquisition (time after injection): 0–2 min: 5–15 s/frame; 2–5 min: 20–30 s/frame; 5–10 min: 60 s/frame; 10–18 min: 120 s/frame; 18–30 min: 180 s/frame; 30–60 min: 300 s/frame</td>
<td></td>
</tr>
<tr>
<td>Image reconstruction</td>
<td>Method harmonized as much as possible in multicenter study; for iterative reconstruction, larger number of iterations should be considered to ensure uniform convergence, regardless of image reconstruction algorithm, followed by suitable filter to control noise if desired</td>
</tr>
</tbody>
</table>

### Table 2
Recommendations on Image Analysis Parameters

<table>
<thead>
<tr>
<th>Modality</th>
<th>Analysis method</th>
<th>Abbreviation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Static</td>
<td>Body-weight SUV&lt;sub&gt;max&lt;/sub&gt;</td>
<td>SUV&lt;sub&gt;max&lt;/sub&gt;</td>
<td>Maximum SUV* (unitless)</td>
</tr>
<tr>
<td>Dynamic</td>
<td>Patlak</td>
<td>K&lt;sub&gt;i&lt;/sub&gt;</td>
<td>Net influx rate of ¹⁸F-FDG (mL plasma/mL lung/min)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>V&lt;sub&gt;ss&lt;/sub&gt;</td>
<td>Steady-state partition coefficient (mL/cm&lt;sup&gt;3&lt;/sup&gt;)</td>
</tr>
<tr>
<td></td>
<td>Compartamental modeling</td>
<td>K&lt;sub&gt;ic&lt;/sub&gt;</td>
<td>Net influx rate of ¹⁸F-FDG (mL plasma/mL lung/min) (from K&lt;sub&gt;i&lt;/sub&gt;, k&lt;sub&gt;2&lt;/sub&gt;, and k&lt;sub&gt;3&lt;/sub&gt;)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>V&lt;sub&gt;b&lt;/sub&gt;</td>
<td>Fractional blood volume (unitless)</td>
</tr>
<tr>
<td>CT</td>
<td>Air fraction determination</td>
<td>V&lt;sub&gt;a&lt;/sub&gt;</td>
<td>Fractional air volume (unitless)</td>
</tr>
</tbody>
</table>

*Air fraction correction may be applied using V<sub>a</sub>.

**Injection Duration**

Very little information has been published on injection duration for dynamic acquisitions, even though this can influence the compartmental modeling results. Observations at several participating centers indicate that a bolus with a shorter injection duration and adequate early-frame time sampling will better characterize the peak of the blood time–activity curve and may avoid a biased estimate of
fractional blood volume ($V_{fb}$, unitless), whereas a longer injection duration may lead to improved fitting of the lung time–activity curves and better estimate the metabolic rate. Until these observations have been validated, a specific injection duration cannot be recommended. Until then, we recommend that the injection technique be defined for each study protocol and reported accordingly, with steps taken to ensure consistency for all scans within a study, particularly when several personnel are involved.

### IMAGE ACQUISITION

#### Static Versus Dynamic

Static PET imaging is the most common acquisition protocol used clinically. Static image endpoints have been shown to correlate with lung physiology and quality-of-life measurements (19) and thus may be sufficient to meet the study objectives. For example, static acquisitions may be sufficient for studies focused on areas of higher density, such as fibrotic regions or inflammatory nodules, that often demonstrate relatively high uptake levels. Using a reference region may also provide sufficient correction for differences in the blood signal due to systemic factors, such as variations in the metabolism of white blood cells, without the need for a full kinetic model, as explored for idiopathic pulmonary fibrosis (12). Delayed postinjection static images may also help reduce the influence of blood activity by allowing it to clear, as previously explored using images obtained at 180 min after injection (20). However, this delay may make quantification in normal lung tissue challenging because of the very low signal. Major advantages of static acquisitions include a short scanning duration that is well tolerated, as well as broader accessibility, as every clinical site will have the capacity to acquire and analyze the data.

Dynamic imaging enables estimation of kinetic parameters and can add insights into the origins of the PET signal. Kinetic modeling of blood and lung cell compartments (parenchyma, airway wall, vascular wall, and immune cells (2)) to enable corrections for $V_{fb}$ contributions may be achievable (11,21), thus making dynamic acquisitions an important consideration when attempting to account for such effects with a single tracer. Additionally, dynamic scan parameters have been shown to be more sensitive than static endpoints in quantifying the presence of low-level lung inflammation under certain conditions, such as in acute lung injury models (22), and in discriminating the sources of increased $^{18}$F-FDG uptake in different mechanisms of injury (23). The aims of each research study will define the optimal approach, and the choice should be justified when the study is reported.

The partial volume of air in each voxel will affect both static and dynamic acquisitions and may vary among lung regions. This effect can be corrected using appropriately acquired CT images matched to the PET images for respiration, referred to here using the term air fraction correction. As new tracers are introduced for lung imaging, the kinetics and dynamic range of uptake will also be important factors in deciding between static and dynamic acquisitions.

#### Respiratory Motion

Because of the unique function of the lungs, the large variation in air volume with normal respiration can reduce the accuracy of PET quantitation. Clinical CT examinations are routinely performed during a breath-hold, often at full inspiration. Since respiration must occur during the longer static and dynamic PET scans, locally varying displacement and compression of tissues will potentially contribute to errors in both attenuation correction and air fraction correction. The displacement causes blurring and increases the partial-volume effect, particularly near the diaphragm, whereas the compression of tissues affects density and radiotracer concentration (24). A mismatch between the point in the respiratory cycle at which the CT portion of the PET/CT scan is acquired and the average position represented by the PET data can therefore lead to artifacts from attenuation correction and air fraction correction and is an important source of error.

Multiple approaches to removing or limiting respiratory motion have been investigated. Shallow breathing during the CT acquisition or breath-holds at gentle end-expiration, as for cancer imaging (25), or at mid-expiration for idiopathic pulmonary fibrosis patients (26,27) can minimize the effects of misregistration of PET and CT images due to respiratory motion. Visual feedback systems to monitor and continuously display the lung volume to the subject have been used but are not widely adopted (28). Acquisitions using repeated breath-holds have also been suggested (29) but can be difficult for those with reduced lung function. Longer CT scans acquired over the duration of the respiratory cycle, such as cine-CT or low-pitch helical CT, can be used to create averaged CT scans (30). A 4-dimensional CT dataset sorted according to the respiratory signal can be used for gated PET data (31), with several studies investigating various respiratory gating strategies for PET (32). Interpolated averaged CT scans constructed from full-inspiration and end-expiration scans may also be considered (33), a method that could also be applied to CT scans obtained from dynamic imaging protocols in which breaks are built into the dynamic acquisition (34).

The approach most used among the centers to minimize PET and CT misregistration is breath-hold at gentle end-expiration or at mid-expiration for the CT scan. Prescan coaching to familiarize the patient with breathing instructions improved compliance and is thus recommended. Since many of the advanced techniques described above are not widely available and require postimaging offline processing, we recommend that, for research studies, list-mode acquisitions be acquired and stored for future reprocessing when respiratory gating and other advanced techniques to minimize respiratory motion become readily available.

#### Duration and Time Frame

Clinical static $^{18}$F-FDG PET imaging protocols have been used successfully to image lung disease (19). Typically, data are acquired over a short interval starting approximately 60 min after
tracer injection. For dynamic imaging studies, acquisitions start at the same time as tracer administration and range from 45 to 90 min in duration. The scan durations reported by the centers are routinely tolerated without discomfort or safety issues and allow robust kinetic analysis. Typically, short frames are used initially to capture the early dynamics of the tracer in blood and are followed by longer frames later in the acquisition. For a typical 60-min scan, our recommended time frames are 5–15 s/frame for the first 2 min after injection, 20–30 s/frame at 2–5 min, 60 s/frame at 5–10 min, 120 s/frame at 10–18 min, 180 s/frame at 18–30 min, and 300 s/frame at 30–60 min. We again recommend retaining the list-mode data to allow future exploration of different time frames.

Most modern PET/CT scanners can cover the entire lung at gentle end-expiration at a single bed position. For those requiring 2 bed positions to cover the entire lung, a dynamic data acquisition initially over 1 bed position followed by a static data acquisition at 2 bed positions may be used. Alternating 2 bed positions over the scan duration, such as alternating seven 4-min acquisitions at each position, is another approach that more easily accommodates breaks between the bed-position scans. Breaks should be scheduled in such a way that disruption of acquisition of the time–activity curve peak during the vascular phase after injection is avoided (Fig. 1). The recently developed total-body PET scanner (35) will eliminate the need for such protocols but is not yet widely available.

**IMAGE RECONSTRUCTION**

Various PET image reconstruction algorithms are available, with different algorithms leading to different noise and image characteristics, depending on the choice of parameters, filters, or a priori assumptions (36). Further, for iterative algorithms, the convergence rate of the values in low-count regions such as the lungs is often lower than in high-count regions (37,38). This difference may affect the observed radiotracer concentration in the lung; therefore, we recommend using a large number of iterative updates (i.e., number of iterations times number of subsets) to ensure uniform convergence (38,39). We agreed that more investigation is needed to define optimal reconstruction specifically for lung PET imaging and that we therefore cannot make specific recommendations based on current data. Additionally, the development of specific phantoms that model relevant aspects of lung physiology and enable the establishment of harmonization standards will be needed to support multicenter studies.

Time-of-flight (TOF) PET scanners have been shown to reduce noise, especially in large patients (40). In addition, the local effects of attenuation mismatch between reconstructed non-TOF PET and CT images are reduced (41–43), albeit at the expense of nonlocal effects outside the lung (27). Although TOF PET/CT scanners have become more widely available, insufficient data are available to make a firm recommendation on whether TOF is valuable for quantitative lung imaging. For consistent results in multicenter studies, we recommend that either data from scanners with a similar TOF time resolution be used (44) or that only non-TOF reconstructed images be used.

Table 1 summarizes our recommendations on patient preparation, acquisition protocols, and image reconstruction.

**IMAGE ANALYSIS REPORTING**

Quantitative PET pulmonary image analysis involves extracting volumes of interest (VOIs) and quantifying the \(^{18}\text{F}-\text{FDG}\) signal within the VOIs. As different study requirements preclude an absolute recommendation of a single methodology for all future studies, we agreed that consistent reporting, in accordance with the guidelines below, would enable better comparison of studies and interpretation of findings from future studies.

**Whole Lung Analysis**

Use of the attenuation correction CT for lung segmentation is recommended. For consistency, performing a quality control check on the segmented lung mask is essential to exclude the chest wall, major airways, bullae, heart, and liver. The lung mask created in CT space should be resampled to PET resolution, for which we recommend using the nearest-neighbor method to obtain a binary mask suitable for VOI processing. In the presence of any gross movement of the patient during the PET and CT acquisitions, the images should be registered to minimize the differences. Details should be provided on methods to confirm that the lung mask segmentation PET and CT are aligned.

**Regional Lung Analysis**

To investigate regional \(^{18}\text{F}-\text{FDG}\) distribution, the whole lung mask can be subdivided into multiple regions either by anatomic lobes, such as through fissure detection (45), or by simple geometric division into upper, middle, and lower lung zones based on either length or volume (10,11). Similarly, anterior–posterior subdivisions and regions with different densities have also been used. There is insufficient information to allow recommendation of a single optimal approach, as the approach may also vary with the disease. Therefore, we recommend reporting all regional analysis details, including definition and selection of VOI location and size.
Quantification of $^{18}$F-FDG Uptake

Static and dynamic analysis techniques have been investigated to account for various lung-specific issues in PET quantification as described above. Below, we describe our consensus recommendations for reporting these quantities.

**Static Data Analysis.** $\text{SUV}_{\text{max}}$ (unitless), defined as the highest single-voxel value within a VOI, is the most widely used measure of $^{18}$F-FDG uptake and warrants investigation despite several potential confounding factors that could influence its measurement, including partial-volume–averaging effects (15). Corrections to the SUV for fractional air volume ($V_a$, unitless) and for lung volume measured, should be applied and reported (46,47). $\text{SUV}_{\text{peak}}$, without or with correction for lean body mass (unitless), measured from an approximately $1 \text{ cm}^3$ VOI with the highest value as defined by PERCIST (48), is less influenced by image noise and may also be considered. Comparing areas of diseased lung to normal lung to determine a target-to-background ratio (unitless), as well as characterizing lung heterogeneity, may also be useful approaches (12,49).

**Dynamic Data Analysis.** Dynamic datasets can be analyzed using either Patlak graphical analysis or compartmental modeling. In both cases, an input function is required to describe the time course of radioactivity concentration in arterial plasma. An image-derived input function may be obtained from the dynamic PET data in preference to blood sampling. Image-derived input functions from the pulmonary artery, ascending aorta, descending aorta, and superior vena cava have been investigated (50). In our combined experience, the use of the pulmonary artery, ascending aorta, and descending aorta reached similar outcomes for rate of transfer of $^{18}$F-FDG (influx constant, or $K_i$ [mL plasma/mL lung/min]) (51,52). The right ventricle can also be used as a source for the image-derived input function (53) but is not useful if not consistently included in the field of view for lung PET scans. Time delays between passage of the tracer through the image-derived input function VOI and the tissue of interest may affect blood volume estimates but are less important for $K_i$. The size of the VOI should be reported and should be consistent for all patients within the same study.

**Patlak Graphical Analysis.** Patlak graphical analysis estimates the net influx rate of irreversibly trapped tracers such as $^{18}$F-FDG from the blood into target tissue ($K_i$) and an approximate steady-state partition coefficient between tissue and plasma of nonphosphorylated $^{18}$F-FDG ($V_{ss}$, mL/cm$^3$) (54). $V_a$ was initially assumed to reflect changes in density, leading to explorations of normalizing $K_i$ with the $V_a$ (6). However, normalizing the $K_i$ by the Patlak intercept has been shown algebraically to remain sensitive to changes in $V_a$ and $V_b$ (11,21). Therefore, caution is advised in its application and interpretation as a measure of the true net influx rate. For accurate comparison, the results of Patlak graphical analysis should include both $K_i$ and $V_{ss}$ individually if normalized $K_i$ is used, as reported previously (6,10–12).

**Compartmental Modeling.** Nonlinear regression models are used to estimate the microparameters of the model ($K_1$ [mL plasma/mL lung/min], $k_2$ [1/min], $k_3$ [1/min]) and the blood contribution ($V_b$) from lung time–activity curves and the image-derived input function, to then estimate $K_i$ using a 2-compartment model (2). Since $^{18}$F-FDG can be considered irreversibly trapped over the time frame of the scan, $k_4$ is assumed to be zero. Air fraction correction may also be included in the compartmental modeling, for which net influx rate is denoted as $K_{eq}$ (11,50). The quality of the fit of the data should be assessed and reported using a metric such as $\chi^2$. Care should be taken to report the compartmental modeling method accurately to enable reproducibility, given the various approaches available. The presence of edema in some conditions such as acute lung injury may require an additional compartment (55).

Table 2 summarizes reporting of image analysis parameters. Table 3 lists compartmental modeling–specific parameters that we recommend to promote reproducibility in reporting (56).

**CONCLUSION**

To improve reproducibility both between and within centers and the potential for data pooling, we urge investigators using PET/CT in studies of lung disease to follow the recommendations presented in this article when designing, conducting, and reporting studies. As highlighted in this article, ample opportunities for investigation exist to improve the methods used to acquire and analyze lung PET/CT images. We hope this summary will serve as a basis for advancements in the field of lung PET imaging.

**DISCLOSURE**

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