Total-body PET kinetic modeling and potential opportunities using deep learning

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Synopsis

The uEXPLORER total-body PET/CT system provides a very high level of detection sensitivity and simultaneous coverage of the entire body for dynamic imaging for quantification of tracer kinetics. This paper describes the fundamentals and potential benefits of total-body kinetic modeling and parametric imaging focusing on the noninvasive derivation of blood input function, multiparametric imaging, and high-temporal resolution kinetic modeling. Along with its attractive properties, total-body kinetic modeling also brings significant challenges, such as the large scale of total-body dynamic PET data, the need for organ and tissue appropriate input functions and kinetic models, and total-body motion correction. These challenges, and the opportunities using deep learning, are discussed.

Keywords

Total-body PET; dynamic imaging; kinetic modeling; parametric imaging; deep learning

Technical Terms

Total-body PET; dynamic imaging; kinetic modeling; parametric imaging; image-derived input function; high temporal resolution; dual-tracer; time-delay correction; model selection; deep learning; convolutional neural network

1. Introduction

Positron emission tomography (PET) is a sensitive molecular imaging method that uses radiolabeled tracers to monitor the biological and physiological function of the scanned subject in vivo¹. PET is commonly used in the clinic to acquire static images of radioactivity distribution at a specific time interval (e.g., 60 minutes) post radiotracer injection. The standardized uptake value (SUV) is provided as a semi-quantitative measure of tracer uptake which, however, is affected by many factors, including body habitus, dietary preparation,
and scan timing. Dynamic PET imaging can also be performed by taking multiple consecutive frames, typically starting at the time of injection. The acquired four-dimensional (4D: 3D space and 1D time) data from dynamic PET may reflect a broad spectrum of physiological and metabolic information, including blood flow, tracer delivery, transport, and metabolism. Dynamic PET data are commonly analyzed using compartmental modeling to quantify physiological parameters in a region of interest (ROI) or at the image voxel level (i.e., parametric imaging)\textsuperscript{2,3}.

Although dynamic PET with tracer kinetic modeling has many potential advantages over static PET imaging, its widespread use and clinical implementation have been hampered by several factors such as high noise, the need of blood input function, long scan time, etc\textsuperscript{4}. Dynamic PET also has been largely restricted to single-organ imaging, e.g., for brain and heart, due to the short axial field-of-view (AFOV) of conventional PET scanners (usually \~20 cm)\textsuperscript{5}. Implementation of whole-body dynamic imaging has become feasible on conventional PET scanners\textsuperscript{6-8}, but it is challenging to achieve high temporal resolution, and the data is sparse in the temporal domain due to the need to move the subject relative to the scanner to image different parts of the body. The recent advent of the uEXPLORER total-body PET system (and other long AFOV scanners that cover all or most of the vital organs\textsuperscript{9,10}) provides unprecedented levels of detection sensitivity and simultaneous coverage of the entire body for dynamic imaging\textsuperscript{5,11,12}. The total-body kinetic modeling and parametric imaging enabled by this system may have potential in both molecular imaging research and clinical applications\textsuperscript{13-16}. Meanwhile, challenging technical problems co-exist with the opportunities given the large scale of total-body dynamic data sizes and the need for consideration of physiological heterogeneities in different organs and of the presence of motion that occurs throughout the dynamic series of images. This paper provides a brief introduction of total-body dynamic PET imaging and kinetic modeling (section 2.1-2.3), describes its potential benefits and limitations (section 2.4-2.6), and discusses potential directions to address the challenging problems using deep learning (section 3). Image examples shown in this paper are mainly from the widely used tracer \textsuperscript{18}F-fluorodeoxyglucose (FDG), but the concept is also applicable to most other radiotracers.

## 2. Total-body dynamic PET and its potential for kinetic modeling

### 2.1. Basis of dynamic PET and kinetic modeling

Figure 1 shows the flowchart for dynamic PET imaging and kinetic modeling. An example of tracer kinetic modeling with compartmental models\textsuperscript{17} for metabolic imaging is given in Figure 2 for \textsuperscript{18}F-FDG. The FDG net influx rate $K_j = \frac{k_1 k_3}{k_2 + k_3}$ is directly proportional to the metabolic rate of glucose and is a macro parameter of interest \textsuperscript{2,3}. Compartmental modeling usually needs nonlinear curve fitting to estimate the kinetic parameters, which is computationally expensive and also sensitive to noise. As an alternative to compartmental modeling, the Patlak model\textsuperscript{18} is a linear graphical method that can approximate $K_j$ using the slope of a graphical plot of the blood input function and tissue time-activity curve (TAC). It has the advantages of computational efficiency and noise robustness for parametric imaging. Examples of compartmental models and graphical methods for modeling ligand-receptor
kinetics are reviewed by Watabe et al.\textsuperscript{19}. The potential of kinetic modeling and parametric imaging has been demonstrated in both research studies and clinical diagnosis\textsuperscript{20,21}.

One challenge of kinetic modeling is to obtain the input function \( C_p(t) \). Conventionally it would be obtained using arterial blood sampling which, however, is invasive. To reduce invasiveness, population-based input functions (PBIF) can be used in combination with one or two blood samples\textsuperscript{22}. As the spatial resolution of PET scanners has improved over time, it has become feasible to non-invasively derive a blood input function from dynamic PET images if a large blood region (e.g., left ventricle or aorta) is available in the scanner FOV. This type of input function is usually referred to as the image-derived input function (IDIF).

### 2.2 Limitations of dynamic whole-body imaging on conventional PET scanners

Conventional PET scanners have a short AFOV, typically 15-30 cm, as Figure 3(A) shows. To acquire a set of whole-body dynamic images in sequence, the scanner must use multiple bed positions and multiple passes\textsuperscript{8,23}, as shown in Figure 3(B). As a result, early-phase data that has unique information linked to blood flow and blood volume is only available for the organs imaged in the first bed position for conventional scanners but is missing for most of the body. There are also large temporal gaps in dynamic frames at any given scanned location. Furthermore, due to the isotropic nature of the annihilation photon emission, the detection sensitivity within the short AFOV is low\textsuperscript{11}, leading to a high noise level in dynamic images.

The acquisition of an input function \( C_p(t) \) for whole-body kinetic modeling with short AFOV PET scanners is another challenge. If measuring of the IDIF is desired, the location of the first bed position may need to be shifted from the main organ of interest (e.g. the brain) to a location covering the aorta, losing valuable early information in the organ of interest. This means some (complex) kinetic model models (e.g. two-tissue compartment model) can no longer be used for analyzing the tissue of interest.

### 2.3 Total-body dynamic imaging on uEXPLORER

The development of the uEXPLORER total-body PET/CT system\textsuperscript{12}, with a 194 cm AFOV, is an important step in addressing some limitations of conventional PET scanners, as shown in Figure 4. It allows simultaneous dynamic imaging of the entire body. This eliminates the large temporal gaps in conventional dynamic whole-body imaging. The total-body axial coverage also increases the scanner detection sensitivity by 20-40 fold and the image signal-to-noise ratio by 5-6 fold\textsuperscript{5} for imaging the entire body. In addition, the high sensitivity allows dynamic imaging with much higher temporal resolution (HTR), such as 1 s per time frame\textsuperscript{13} or even 0.1 s per frame\textsuperscript{24} compared to 10-40 s per frame in traditional protocols, which may be used to explore novel clinical applications.

### 2.4 Potential benefits of total-body dynamic PET for kinetic modeling: examples

#### 2.4.1 Noninvasive IDIF—Total-body dynamic images acquired from the uEXPLORER system bring several potential benefits for non-invasive IDIF determination. As compared to dynamic whole-body imaging on conventional PET scanners, the total-body coverage of uEXPLORER allows an IDIF to be extracted from major blood pools in the body without
compromising the temporal resolution of imaging any other organs. This is advantageous even for single-organ dynamic PET imaging. For example, for dynamic brain imaging with conventional PET, the carotid arteries are the largest available blood pool present in the FOV from which to derive an IDIF; with a diameter of 5-6 mm, the carotids have little signal and suffer from severe partial volume effects. Total-body dynamic PET overcomes this problem by providing a low-noise IDIF from the large blood pool present in the FOV such as the left ventricle or ascending/descending aorta. Figure 5 shows an example of brain parametric imaging from a healthy subject scan on the uEXPLORER, where carotid partial volume effects result in increased $K_i$ estimates.

The ability to acquire low-noise input functions, non-invasively, and measured using the same device as the tissue TACs, aids in unbiased kinetic parameter estimation. While most organs are supplied by a single blood input from the arterial system, some organs have dual blood supplies. The liver, for example, is supplied by both the hepatic artery and the portal vein. However, IDIF extraction from the portal vein is difficult with conventional dynamic PET imaging, due to the combined effect of limited spatial resolution, the small size of the portal vein (about 10 mm), and high noise levels present in a small ROI. This challenge can be reduced by total-body dynamic PET imaging on uEXPLORER which with much higher sensitivity and higher spatial resolution will allow better estimation of the IDIF from such vessels.

### 2.4.2 Multi-parametric imaging

Limited by the short AFOV, conventional PET scanners cannot simultaneously capture total-body kinetic signals, especially in the early phase of radiotracer bolus distribution. Thus, parametric imaging has usually been limited to the linear Patlak plot method in dynamic whole-body PET studies with conventional scanners. This method provides a slope image for the macro kinetic parameter $K_i$ and an intercept image related to a combination of fractional blood volume $v_b$ and volume of distribution. However, the full potential of compartmental modeling that allows quantification of micro kinetic parameters (e.g., the tracer delivery rate $K_1$ and fractional blood volume $v_{fb}$) are difficult to explore with such systems. For example, a whole-body $K_i$ image can be obtained with conventional scanners, whereas a whole-body $K_1$ image cannot.

Total-body dynamic imaging with the uEXPLORER system has the potential to address this shortcoming and enable high-quality total-body kinetic modeling and parametric imaging of micro kinetic parameters. Figure 6 shows an example for parametric imaging of FDG uptake rate $K_i$, fractional blood volume $v_f$, and FDG delivery rate $K_1$ from a uEXPLORER scan.

Among the different kinetic parameters measured, the tracer delivery rate $K_1$ is of particular clinical interest due to its connection to blood flow. This parameter is different from the influx rate $K_i$ and hence can provide complementary information, e.g., potentially for simultaneous evaluation of perfusion-metabolism mismatch using FDG for myocardial viability. Parametric imaging of the relative delivery rate $R_1$ (ratio of $K_1$ between a tissue region and a reference region) of tau tracers and beta amyloid tracers is also being explored as a surrogate of cerebral blood flow to simultaneously assess both blood flow and misfolded protein changes in neurodegenerative disease using a single tracer. Readers are
referred to the section III.D of a recent review article on parametric imaging for more
details\(^4\).

Total-body parametric imaging of the fractional blood volume \(v_b\) may also add useful
physiological and pathological information. For example, chronic obstructive pulmonary
disease (COPD) changes lung blood volume\(^27\). \(v_b\) may also reveal the local blood supply
and microenvironment of a tumor, thus may be helpful to improve tumor diagnosis and
characterization\(^28\).

**2.4.3 High-temporal resolution kinetic modeling**—The ability of uEXPLORER for
HTR imaging (e.g., 1 s per frame) allows not only better temporal sampling of the blood
input function, but also more accurate modeling of fast tracer kinetics. For example, after
bolus injection, radiotracer signal in the lungs is supplied by the pulmonary artery and
bronchial artery. However, differentiation of the contributions from the two blood supplies is
challenging if the temporal resolution of dynamic imaging is insufficient. With total-body
dynamic PET, it becomes feasible to measure lung TACs with high temporal resolution and
derive the bronchial arterial input function and pulmonary arterial input function from the
left ventricle and right ventricle, respectively. Our preliminary results have demonstrated
HTR imaging has a significant effect on the quantification of \(^{18}\)F-FDG \(K_1\) and \(v_b\) in the
lungs\(^29\).

The second example is the multiphase Patlak plot. Patlak plots derived with standard
temporal resolution commonly show a single late-time linear phase. With HTR imaging,
preliminary results from Zuo \textit{et al.}\(^30\) demonstrated two additional approximately linear
phases: one at around 20-30s (first-pass) and the other at around 1-2 minutes (early-time)
(Figure 7). Total-body parametric images of the slopes of the first-pass, early-time and
standard late-time linear phases also demonstrate different spatial patterns. It is worth noting
that Patlak plot has been used with high temporal resolution in dynamic contrast-enhanced
magnetic resonance imaging for assessing blood brain barrier permeability\(^31\). Thus, we
postulated that the first-pass and early-time Patlak slopes may be related to blood flow and
tracer permeability, though the precise meaning and physiological basis for these earlier
linear phases remain to be determined.

Another example of HTR kinetic modeling is for the separation of blood flow and tracer-
specific transport from the overall tracer delivery rate \(K_1\), e.g., through time-varying kinetic
modeling that is currently under development\(^32,33\). For \(^{18}\)F-FDG, HTR kinetic modeling
may make it possible to derive blood flow, glucose transport, and glucose metabolism
simultaneously from a single dynamic scan.

**2.4.4 Dual-tracer dynamic PET imaging**—Single-scan dual-tracer (or multi-tracer)
PET imaging has attracted a lot of interest over the past decades (see the review paper from
Kadma\(\text{\textit{s}}\) and Hoffman\(^34\)). To recover separate images of each tracer from the same scan,
dynamic imaging and kinetic modeling can be used to separate the two tracer signals from
each other. The robustness of single-scan dual-tracer methods has typically been limited by
data noise. The increased sensitivity of total-body PET scanners is offering new
opportunities to make this framework more robust and feasible for clinical investigation. The
total-body coverage will also allow simultaneous dual-tracer dynamic imaging of the entire body.

2.5 Potential clinical impact

Compared to the semi-quantitative SUV used routinely in the clinic today, total-body kinetic modeling on the uEXPLORER opens up the window for studying systemic diseases quantitatively and in a multi-parametric fashion using a single radiotracer. One obvious example is studying the heterogeneity of metastatic tumor characteristics, both before and after treatment. Total-body PET allows for kinetic data to be acquired simultaneously for all metastatic lesions in the body, and multi-parametric imaging may provide a sensitive assay for assessing the response or likely response of each lesion to treatment. For example, dysregulated cellular metabolism and angiogenesis are both hallmarks of cancer\(^{35}\). High-quality multi-parametric imaging of FDG \(K_i\) (reflecting metabolism) and \(K_1\) (reflecting perfusion and glucose transport) has the potential to provide complementary information of tumor metabolism and angiogenesis for more accurate tumor characterization for response assessment.

Similar approaches could be used to quantify inflammatory or infection burden across the entire body with appropriate radiotracers. For example, in arthritis, total-body dynamic imaging allows all the joints of the body to be assessed simultaneously with kinetic modeling approaches. Finally, absolute quantification of perfusion and blood volume across the entire body would likely have a broad clinical impact, especially if scans can be performed quickly and at relatively low radiation dose. Clinical applications would include a range of cardiovascular diseases (for example, peripheral arterial disease) where the vascular health of the entire body can be assessed and quantified. Whole-body parametric imaging of rapid physiological processes and tracer distributions is particularly challenging on conventional short axial field of scanners as there simply is insufficient time to move the bed and collect data across the entire body.

2.6 Limitations of total-body dynamic PET

While total-body dynamic PET brings several potential benefits, some limitations of conventional dynamic PET imaging still remain. For example, parametric imaging of FDG \(K_i\) usually requires a one-hour long scan, which is one of the main hurdles that limit the widespread adoption of dynamic FDG-PET in clinics. This hurdle remains with total-body dynamic PET imaging. Many radiotracers have the issue of being metabolized during the imaging time. Metabolite correction is needed for determining the true parent plasma input function\(^{36}\), which remains a challenge for total-body kinetic modeling, though new noninvasive methods are being explored. Another limitation of total-body dynamic PET imaging is the limited availability of total-body PET systems. At the time of writing there are approximately 10 installations of total-body or long (> 1 meter) AFOV scanner worldwide. This implies that initially total-body dynamic PET studies will be only conducted in a small number of institutions, although the number of these scanners is expected to increase quite quickly in the coming years.
Total-body kinetic modeling also brings new significant challenges, including the large scale of total-body dynamic datasets and wide physiological heterogeneity in different organs that should be accounted for by using organ and tissue appropriate models. Total-body parametric imaging is further complicated by non-rigid subject motion that occurs throughout the dynamic acquisition. Kinetic parameter estimation with compartmental modeling is also highly nonlinear, suffers from local minima, and is sensitive to noise, resulting in a challenging task for conventional ROI-based modeling that is exacerbated when performing parametric imaging where models are applied on a voxel-by-voxel basis. In addition, the computational efficiency of nonlinear parametric imaging needs to be further improved for practical use because up to 10 million image voxels per acquisition are typically processed in total-body parametric imaging.

3. Opportunities using deep learning for total-body kinetic modeling

Deep learning has attracted broad attention for its huge potential in almost every field, including PET\(^{37,38}\). One major advantage of deep learning is its ability for end-to-end training of a mapping from problem inputs to answers. Once the model is trained, the prediction is fast. In this section, we discuss a few examples for potential deep learning-based solutions to address the specific challenges in total-body kinetic modeling.

3.1 Voxel-wise corrections for blood input function

In total-body dynamic PET, the IDIF is usually extracted from an ROI selected within a large central blood pool. For a peripheral tissue region that is far from the blood ROI, the arrival time of the radiotracer is delayed compared to the start time of the extracted blood input function. Thus, this time delay of the input function needs to be corrected to achieve accurate kinetic quantification. Time delay correction has been considered impactful only for high-temporal resolution imaging in conventional dynamic PET but is also essential in total-body imaging because the time delay can be 30-40 s in peripheral tissues. In addition, a dispersion correction to account for the mixing of the injected radiotracer in the blood may also be critical to model the input function for a specific organ\(^{14,39}\).

Corrections for these factors can be pursued by joint estimation which estimates the parameters simultaneously with other kinetic parameters during the fitting of TACs. A recent example is given by Feng et al.\(^4\) in total-body imaging of early FDG kinetics. However, the joint estimation method can be computationally expensive. Alternatively, time delay can be corrected voxel-wise before parametric imaging\(^40\). Such a pre-correction method has the advantage of reducing the computational burden. While analytical approaches such as the leading-edge method may not achieve the best accuracy, one of our ongoing efforts is to explore deep learning for regression to train a model that can predict the time delay and/or dispersion of the blood input from the tissue TAC data in combination with the blood input function. The labeling data for training can be built from patient data using the joint estimation method.
3.2 Learning-based input function

For tracer kinetic modeling, a plasma input function is what is actually needed, while an IDIF represents the tracer activity in the whole blood. The tracer fraction in plasma may be pre-determined based on population data, but it is challenging to adapt this to individual patients\textsuperscript{41}. In addition, an IDIF may suffer from partial volume and spill-over effects, which leads to a potential discrepancy between the IDIF and true blood input function.

Machine learning has the potential to provide a robust and fast prediction of the input function from tissue TACs. For example, Kuttner et al. estimated the IDIF from multiple blood pool TACs or multiple tissue TACs using a Gaussian Process model and long short-term memory network\textsuperscript{42}. Instead of requiring manual ROI placements, machine learning can also estimate the IDIF directly from dynamic images\textsuperscript{43}.

Metabolite correction is needed for some radiotracers\textsuperscript{36} and can be done by arterial blood sampling (or if the metabolism is slow and the arterial-venous difference is small, venous blood sampling may be sufficient) followed by high-performance liquid chromatography analysis, which is complex and invasive, and therefore is often mainly used to derive a population-based model. Mathematical metabolite correction\textsuperscript{44-46} is an alternative solution that jointly estimates the parent plasma input function during TAC fitting\textsuperscript{47}. However, this method requires a complex optimization due to the high nonlinearity of the model and can easily get stuck at a local minimum and has a high computational cost. One possible solution is to use deep learning to learn an end-to-end solution from the TAC data by using the global optimization or arterial sampling result as the training labels. Once trained in this manner, the model can be efficient to use.

3.3 Voxel-wise kinetic model selection

In conventional parametric imaging with short AFOV scanners, one kinetic model is commonly used for the kinetic parameter estimation for all image voxels. However, the physiology varies among organs in the body and is beyond the descriptive ability of a single kinetic model. An oversimplified model causes biased quantification, while a too complex model leads to unstable parameter estimation. This issue can be addressed by voxel-wise model selection which chooses the best of two or more candidate models. As shown in Figure 8(B), the parametric image generated by a standard two-tissue irreversible compartment model produces artifacts in the ventricular blood pool and blood vessels (pointed out by arrows). These artifacts can be suppressed by the appropriate model selection, as shown in Figure 8(C).

Conventional methods for model selection use statistical metrics to balance the trade-off between the curve-fitting error and the model complexity. Golla et al. compared several model selection metrics, including Akaike Information Criterion, Model Selection Criterion, and Schwartz Criterion, and found strong agreement across them\textsuperscript{48}. These methods can be implemented in a voxel-wise manner but are time-consuming for total-body imaging.

Deep learning may bring an efficient solution to kinetic model selection by formulating model selection as a binary or multi-class classification problem. The spatial correlation of neighboring voxels can also be incorporated into the learning model. Such an approach has
been recently explored by Klyuzhin et al.\textsuperscript{49} and Fuller et al.\textsuperscript{50} for detecting transient responses in neurotransmitter PET.

### 3.4 Fast and robust prediction of kinetic parameters

Machine learning has also been applied for directly predicting kinetic parameters from the TAC data. A trained model has the potential to generate parametric images very efficiently and be robust to noise. Early attempts were demonstrated in 2001 using a shallow neural network for $^{13}$N-ammonia perfusion parametric imaging by Golish et al.\textsuperscript{51}, in 2011 using a support vector machine for kinetic prediction with a shortened dynamic FDG-PET protocol\textsuperscript{52,53}, and recently using a deep convolutional neural network for predicting the parametric images of macro parameters\textsuperscript{54}.

While learning a direct mapping from the TAC space to kinetic parameter space remains a big challenge, it is more feasible to predict high-quality parametric images from low-quality parametric images generated by conventional kinetic modeling. For example, parametric imaging commonly uses an indirect approach that reconstructs the dynamic images first and then performs kinetic modeling voxel by voxel to generate parametric images, as discussed in the sections above. In comparison, the ‘direct method’ combines the kinetic model into the forward projection model of PET image reconstruction and estimates the parametric images directly from the projection data. Direct parametric image reconstruction has been demonstrated to outperform the indirect method\textsuperscript{55}. However, direct parametric image reconstruction requires a more complex optimization and is time-consuming. Feng et al.\textsuperscript{56} and Xie et al.\textsuperscript{57} used convolutional neural network learning to recover high-quality Patlak parametric images (estimated by the direct method) from the Patlak images estimated by the indirect method.

### 3.5 Single-Subject Deep Learning

One general challenge with deep learning is the requirement of collecting a large number of patient scans for building the training dataset. This is particularly challenging for deep learning for kinetic modeling because each patient has only one blood input function, though many tissue TACs are available for building a database. With a limited number of blood input functions, the generalization capability of a trained model remains a concern.

Single-subject deep learning may be a feasible alternative solution and is being pursued in our group. In this method, a deep learning model is trained on the fly with a low computational cost by using a small fraction of voxels labeled with conventional kinetic modeling. The remaining voxels are unlabeled but will be predicted efficiently using the trained model. One advantage of this learning method is all the training and testing samples share the same blood input function. The labeled and unlabeled data can be further combined through a semi-supervised learning framework\textsuperscript{58}.

### 3.6 Voxel-wise motion correction

As the image data size and FOV increase with total-body PET, various types of subject motion, involuntary (e.g. respiratory, cardiac, bladder filling) and voluntary (e.g. head tilting) are always present in the FOV. For the purposes of total-body parametric imaging,
positioning changes in the tissue will be reflected in the individual voxel TACs, which in turn may affect kinetic parameter estimation. Motion tracking hardware$^{59,60}$ and data-driven methods$^{61}$ such as the centroid of distribution method$^{62-64}$ can be applied to total-body dynamic imaging. Note that total-body motion is generally non-rigid and requires time-consuming algorithms for motion correction. Deep-learning based solutions may have the advantage to efficiently and non-rigidly register respiratory-gated PET images$^{65}$, and may provide improved results with high-quality images generated from total-body PET.

4. Summary

Total-body kinetic modeling on the uEXPLORER system enables quantitative multi-parametric imaging of the entire body simultaneously. With recent developments in deep learning, the remaining challenges of total-body kinetic modeling and parametric imaging may be addressed. Many opportunities are emerging for exploiting total-body kinetic modeling in various research and clinical applications.

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6. References


PET Clin. Author manuscript.


12. Spencer BA, Berg E, Schmall JP, et al. Performance evaluation of the uEXPLORER Total-body PET/CT scanner based on NEMA NU 2-2018 with additional tests to characterize long axial field-of-view PET scanners. J Nucl Med. Published online 2020:jnumed.120.250597. doi:10.2967/jnumed.120.250597


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**Reading List**

**Must Read**


**Nice to Read**


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Key Points

- Total-body PET kinetic modeling on the uEXPLORER PET/CT system addresses several factors that serve as barriers to implementation of dynamic whole-body PET imaging on conventional PET scanners.

- Total-body kinetic modeling with increased detection sensitivity supports multi-parametric imaging which has clinical potential, but also brings several technical challenges.

- Deep learning provides multiple opportunities in total-body kinetic modeling, including non-invasive input function estimation, kinetic model selection, and kinetic parameter estimation. Applications of deep learning can help with further improvement of accuracy, robustness, and efficiency.
Figure 1.
Flowchart of PET kinetic modeling and parametric imaging. Raw PET projection data in the format of sinogram or list-mode are acquired and reconstructed into dynamic images. For each image voxel or a region of interest (ROI), a time activity curve (TAC) $C_T(t)$ is extracted from the dynamic sequence and fitted using a kinetic model and a blood input function to estimate kinetic parameters. The input function can be either noninvasively derived from the dynamic images or invasively measured by arterial blood sampling or is a population-based input function (PBIF). Kinetic modeling can be ROI-based or voxel-based (i.e., parametric imaging).
Two-tissue compartment model for $^{18}$F-FDG. FDG is transported from plasma to tissue cells with the delivery rate $K_1$ and from tissue to plasma with the rate $k_2$. FDG is phosphorylated in cells into FDG 6-phosphate with the rate $k_3$ and the process can be reversed at the rate $k_4$. The total activity measured by PET is $C_T(t)$ that is a sum of different compartments. $v_b$ denoting the fractional blood volume and $C_{wb}(t)$ is FDG activity in the whole-blood.
Figure 3.
Illustration of dynamic whole-body imaging on a conventional PET scanner (A) with a multi-bed multi-pass strategy (B).
Figure 4. Total-body dynamic PET imaging using uEXPLORER. (A) The uEXPLORER total-body PET system. (B) Total-body imaging. (C) Total-body dynamic $^{18}$F-FDG images of a healthy subject. The images were reconstructed with no point spread function modeling and no post-reconstruction smoothing.
Figure 5. Total-body IDIF extraction.
(A) Maximum intensity projection (MIP) image of the dynamic, uEXPLORER dataset of a healthy volunteer. (B) Example IDIFs derived from different blood pool ROIs. (C) Parametric images of FDG net influx rate $K_i$ using different ROIs for IDIF extraction. The carotid-extracted IDIF results in much higher $K_i$ estimates than the LV-extracted IDIF.
Figure 6.
Total-body parametric images estimated from a 60-minute dynamic $^{18}$F-FDG scan of a patient with metastatic cancer on the uEXPLORER: (A) SUV, (B) FDG net influx rate $K_i$ (C) fractional blood volume $v_b$, and (D) FDG delivery rate $K_1$. 
Figure 7.
High temporal resolution Patlak plot and parametric imaging. (A) Patlak plot of one-hour dynamic FDG data; (B) zoom in of the first 2-minute data; (C) parametric images of the slope at three different phases – standard $K_i$ (30-60 minutes), first-pass $K_i$ (20-30s) and early-time $K_i$ (1-2 minutes). Shown are the maximum intensity projection images. The unit of $K_i$ is mL/min/mL.
Figure 8.
Total-body kinetic model selection. (A) The model selection map. (B) Total-body $K_i$ image using the two-tissue irreversible model without model selection. (C) Total-body $K_i$ image with model selection.