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1  **PMOD Kinetic Modeling Tool (PKIN)**

The PMOD kinetic modeling tool represents a flexible environment for the simulation and fitting of models over time. While initially aimed at the compartment models employed in Positron Emission Tomography (PET) and Single Photon Emission Computed Tomography (SPECT), models for other modalities such as magnetic resonance or optical imaging can also be easily incorporated due to the general approach and the plug-in structure for the models.

PKIN includes the following features:

- **import of time-vectors from measurements of blood and tissue activity via text files or directly from the PMOD Volume-Of-Interest (VOI) tool,**
- **fitting of plasma fraction curves to derive tracer activity in plasma from whole blood activity,**
- **fitting of metabolite correction curves to derive free tracer activity from total plasma activity,**
- **representation of the plasma and whole-blood activity by a model function for data smoothing and extension beyond the last measurement,**
- **selection of a comprehensive list of compartment models, reference models and graphical plots,**
- **weighted and non-weighted fitting of the selected model to the measurement whereby the parameter can be enabled or fixed, and can optionally be restricted within a physiological range,**
- **coupled fitting of measurements from different tissues to improve the accuracy for parameters assumed to be identical across tissues,**
- **calculation of goodness-of-fit criteria usable for model comparisons,**
- **Monte Carlo simulations to assess the identifiability of model parameters using a standard or user-defined distribution of the measurement errors,**
- **sensitivity analysis of compartment models to quantify the correlation between the model parameters,**
- **visualization of the relative contributions of the different compartments to the model curve,**
- **generation of synthetic studies representing compartmental kinetics for testing the performance of simplified pixel-wise models,**
- **batch mode operation to fit or Monte Carlo simulate a set of prepared studies,**
- **saving of the fitted parameters into text files and their aggregation for further statistical investigations.**

Please note that the following description is intended as a reference and not as a tutorial. For practical examples how to work with the PKIN software please refer to the tutorial videos available online.
2 **Introduction to Modeling in PET**

For the data processing in PKIN it will be helpful to understand the basic modeling concepts in PET. The following sections present short introductions of compartment and reference models. Other model types are described in the PKIN Model Reference section.

For in-depth understanding of PET Kinetic Modeling we strongly recommend the PMOD users to attend one of the excellent yearly PET Pharmacokinetics Courses which are annually organized prior to the alternating NRM and BrainPET meetings. These courses include theory as well as practical work and are organized by the the top experts in the domain. Another worthwhile course is PET Methodology (King's College London). An additional source of valuable information is the data analysis website of the Turku PET Centre.

### 2.1 Compartment Models

A physiologic system is often described by decomposition into a number of interacting subsystems, called compartments. Compartments should not be understood as a physical volume, but rather as a mass of well-mixed, homogeneous material that behaves uniformly. Each compartment may exchange material with other compartments. Examples of frequently used compartments are

- Authentic (unchanged) tracer in the arterial plasma that can be extracted into the tissue. The concentration of authentic tracer as a function of time during the PET acquisition is called the "Arterial Input Function" (AIF), or simply the "Input Curve".
- Free tracer in tissue that can be bound or that may diffuse back into the blood.
- Tracer in tissue that has been specifically bound, for instance, at the target receptor site.
- Tracer in tissue that has been non-specifically bound to other than the targeted cell components.

Compartment models are visualized by diagrams wherein the rectangles symbolize compartments and the arrows represent material exchange, as illustrated for the different models below. Although the mechanisms of material transport between compartments may differ, models that can be reasonably analyzed with standard mathematical methods assume first-order processes. As a consequence, the change of tracer concentration in each compartment is a linear function of the concentrations in the other compartments. Because of the tiny amounts of tracer material applied in nuclear medicine it is usually assumed that the observed system is not disturbed by the tracer. It is furthermore assumed, that the physiologic conditions do not change during the study, so that the rate of the material exchange can be considered constant.

#### 2.1.1 Model Structure

In the following, the compartment models are illustrated using a tracer that is injected intravenously as a bolus and is not freely diffusible. Please refer to the consensus paper of Innis et al [56] for the details of the nomenclature and the interpretation of the models in the context of brain receptor tracers.

When the tracer arrives in the heart chambers, it is well mixed with blood and distributed by the arterial circulation. It finally arrives at the capillary bed where exchange with the tissue can take place. Some fraction of the tracer is extracted into tissue and metabolized, the rest is transported back to the heart, from where a new circulation starts.

This simplified "physiologic" model can be translated into a 1-tissue compartment model (1TCM):

![Compartment Model Diagram](image)

Because of mixing it is assumed that, the arterial tracer concentration is the same throughout the body, so that it can be measured in a peripheral artery. The tracer concentration in tissue \( C_1(t) \)
increases by the extraction of tracer from the arterial blood plasma. A usual condition for a quantitative analysis is, that only the unchanged tracer, called the \textit{authentic tracer} or \textit{parent}, can enter tissue, whereas labeled metabolites which may also circulate cannot. Because extraction is described by a first-order process, the transfer of material is proportional to the parent concentration $C_P(t)$. Parallel to the uptake, tracer in tissue is reduced by a backwards transfer (or washout), which is proportional to the concentration in tissue. Both processes compete, so that the change over time of the net tracer concentration in tissue ($dC_1(t)/dt$) can be expressed by the following differential equation

$$\frac{dC_1(t)}{dt} = K_1 C_P(t) - k_2 C_1(t)$$

The two transfer coefficients, $K_1$ and $k_2$, have a somewhat different meaning, indicated by using capital and small letters. $K_1$ includes a perfusion-dependent component and has units of milliliter per minute per milliliter tissue, whereas $k_2$ [min$^{-1}$] indicates the fraction of mass transferred per unit time. For example, if $k_2$ equals 0.1[min$^{-1}$], material leaves the tissue compartment at a rate of 10% per minute. The parent concentration $C_P(t)$ is not a compartment, but considered as a measured quantity and appears as the input curve driving the system.

The differential equation of the 1-tissue compartment model can be analytically solved (integrated), yielding the following equation for the concentration of tracer in tissue:

$$C_1(t) = K_1 C_P(t) \otimes e^{-k_2 t} = \int_0^t C_P(\tau)e^{-k_2(t-\tau)} d\tau$$

Hence the time course of the tissue concentration essentially results from a convolution $\otimes$ of the input curve with a decaying exponential. $K_1$ acts as a scaling factor of the concentration course, whereas the washout coefficient $k_2$ has an impact on its form.

More complex compartment models distinguish different forms of tracer in tissue, which is usually interpreted as follows. After entering a cell, the tracer is available for binding in a free form at a concentration $C_1(t)$. Free tracer can directly be bound to its target molecule, $C_2(t)$, but it also may bind to some cell components that are not known in detail, $C_3(t)$. Considering these pathways, a 3-tissue compartment model (3TCM) with six transfer coefficients is established, whereby $C_2$ and $C_3$ communicate only by $C_1$.

The system of differential equations can be derived in analogy to the 1-tissue compartment model, but is much more complex. The equations describing this model are given by

$$\frac{dC_1(t)}{dt} = K_1 C_P(t) - (k_2 + k_3 + k_5) C_1(t) + k_4 C_2(t) + k_6 C_3(t)$$

$$\frac{dC_2(t)}{dt} = k_3 C_1(t) - k_4 C_2(t)$$

$$\frac{dC_3(t)}{dt} = k_5 C_1(t) - k_6 C_3(t)$$

This system contains six unknown parameters ($K_1$ to $k_6$) that are difficult to assess experimentally. For practical purposes, the system is therefore most often reduced to a 2-tissue compartment model (2TCM) by treating free and non-specifically bound tracer as a single compartment $C_1$ (non-displaceable compartment).
This simplification is justified if the exchange free ↔ nonspecific binding is significantly faster than the exchange free ↔ specific binding. Otherwise specific and nonspecific binding cannot be distinguished, and the resulting transfer coefficients cannot be attributed to specific binding alone. The simplified two-tissue compartment model is given by

\[
\frac{dC_1(t)}{dt} = K_1 C_P(t) - (k_2 + k_3) C_1(t) + k_4 C_2(t)
\]

\[
\frac{dC_2(t)}{dt} = k_3 C_1(t) - k_4 C_2(t)
\]

By using a mathematical method called Laplace transformation, analytic solutions can be derived for linear multi-compartment models, but the mathematical formulas are complicated expressions already for two compartments. However, they demonstrate that generally $K_1$ represents a scaling factor like in the solution given for the one-tissue compartment.

**Model Simplification**

While complex models are a more realistic description of the involved processes, they include many parameters. PET data from dynamic measurements has a limited statistical quality, and represents the summed contributions from all compartments. When trying to estimate a large number of model parameters from such data, the variance of the resulting parameters tends to be too high for reliable interpretation. Therefore, only relatively simple models with 2 to 4 (at most 6) parameters are feasible in practice.

When the 3-tissue compartment model is simplified to a 2-tissue compartment model, and further to a 1-tissue compartment model, the meaning of the transfer coefficients changes (except for $K_1$). The “lumped” parameters of the simplified models can be expressed by the parameters of the complex models as shown in the tables below. Note that $k_2$ and $k_3$ have a different meaning in the different models, as well as $C_1$.

<table>
<thead>
<tr>
<th>Parameters in 1TCM</th>
<th>Expressed by parameters of the 2TCM</th>
<th>Expressed by parameters of the 3TCM</th>
</tr>
</thead>
<tbody>
<tr>
<td>$K_1$</td>
<td>$K_1$</td>
<td>$K_1$</td>
</tr>
<tr>
<td>$k_2$</td>
<td>$\frac{k_2}{1 + k_3 / k_4}$</td>
<td>$\frac{k_2}{1 + k_3 / k_4 + k_5 / k_6}$</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Parameters in 2TCM</th>
<th>Expressed by parameters of the 3TCM</th>
</tr>
</thead>
<tbody>
<tr>
<td>$K_1$</td>
<td>$K_1$</td>
</tr>
<tr>
<td>$k_2$</td>
<td>$\frac{k_2}{1 + k_3 / k_6}$</td>
</tr>
<tr>
<td>$k_3$</td>
<td>$\frac{k_3}{1 + k_5 / k_6}$</td>
</tr>
<tr>
<td>$k_4$</td>
<td>$k_4$</td>
</tr>
</tbody>
</table>

The term $1/(1+k_6/k_6)$ represents the free fraction of tracer in the non-displaceable tissue compartment which is available for transfer back to blood or for specific binding, and is often called $f_2$. 

PMOD Kinetic Modeling (PKIN)  
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A good summary on the configuration and interpretation of linear models for receptor tracers is given by Koeppe et al. [15]. Starting from the 3-tissue model it is nicely shown how the parameters of simplified models are related to the true rate constants. PKIN can take these relations into account when switching from a more complex model to a simpler one. This behavior can be switched on by the model conversion check on the Extras panel.

**Relation of K1 with Perfusion**

As mentioned earlier, $K_1$ is related to tissue perfusion $F$. With a capillary model, the relation

$$K_1 = EF$$

can be derived. $E$ represents the unidirectional first-pass extraction fraction (i.e., the fraction of tracer that penetrates the capillary wall and is extracted into the tissue during the first pass of tracer through the capillary). The reverse transport is not regarded in the calculation of $E$. Renkin [40] and C. Crone [41] calculated the extraction as

$$E = 1 - e^{-PS/F}$$

$P$ denotes permeability, and $S$, the surface of the endothelium by which the capillary exchanges with the tissue. The relation demonstrates that the extraction depends on the properties of the endothelium with respect to the tracer ($PS$), as well as the perfusion $F$. The higher the perfusion, the smaller the extraction, because the average time spent close to the capillary wall decreases. However, total mass transport in general still increases with flow, as the reduced extraction is more than compensated by the higher abundance of tracer ($F$ in the term $EF$). For tracers with very high permeability, the extraction is virtually independent of perfusion and $E$ approaches 1 (such as for $^{15}$O-labeled water). In this case, $K_1$ equals perfusion.

**Volumes of Distribution for Receptor Tracers**

For receptor tracers, it is important that the extraction from plasma to tissue is sufficient, such that the kinetics is not limited by the extraction step. Additionally, the relation of the transfer coefficients for specific and nonspecific binding determines whether these different types of binding can be distinguished.

Naturally, the main interest is in specific binding. Besides the individual rate constants $k_3$ and $k_4$, which are often difficult to estimate precisely, the distribution volume of specific binding $V_S$ is a measure which is often used to quantify specific binding. $V_S$ is defined by the ratio of specific binding concentration to total parent at equilibrium (which may not occur during the experiment). However, it can also be calculated from the rate constants of the 3-tissue compartment model as follows.

$$V_S = \frac{K_1}{k_2} \frac{k_3}{k_4}$$

3-Tissue Compartment Model

Under the assumption of a rapid equilibration between the free and non-specific compartment the same expression with the rate constants from the 2-tissue compartment model is often used as an approximation of $V_S$.

If the assumption of rapid equilibration is not justified one must resort to considering the entire tracer concentration in tissue as an indicator of specific binding. The measure used in these cases is called the "total distribution volume" $V_T$. In general it is defined as the ratio of the tracer concentration in tissue to total parent in plasma at equilibrium. A $V_T$ equal to 20 hence means that the tracer is being concentrated in tissue by 20:1. $V_T$ can be easily computed from the rate constants of the compartments model.

$$V_T = \frac{K_1}{k_2}$$

1-Tissue Compartment Model

$$V_T = \frac{K_1}{k_2} (1+k_3/k_4)$$

2-Tissue Compartment Model

$K_1/k_2$ can be interpreted as the distribution volume of the first tissue compartment $C_1$. In receptor tracer experiments, this compartment is usually called the non-displaceable compartment, and $K_1/k_2$ the distribution volume $V_{ND}$ of non-displaceable uptake.

$$V_{ND} = \frac{K_1}{k_2}$$

2-Tissue Compartment Model
This naming indicates that the concentration of free and non-specifically bound tracer in tissue is not reduced when a non-labeled ligand is brought into the tissue, competing with binding to the same receptor as the tracer. The rational for this assumption is that the potential for non-specific binding to various cell components is practically unlimited as compared to the number of specific binding sites.

**Rate Constants and Binding Potential for Receptor Tracers**

- $k_2$ represents the fractional rate constant for the washout of free tracer from tissue to plasma. It is a composite rate constant and is related to flow in the same way as $K_1$. Consequently it is often reasonable to assume that the ratio $K_1/k_2$ is insensitive to flow and regionally constant. This property can be exploited as a restriction in a model fit, hereby reducing the number of fitted parameters.

- $k_3$ is regarded as the rate of association of the ligand with the specific binding sites. It is a function of the concentrations of the free radioligand, the available binding sites and intrinsic second order association rate constants. Consequently it is a pseudo first order rate constant incorporating the concentration of available binding sites.

- $k_4$ is considered as the dissociation rate constant of the receptor-ligand complex; often it is assumed to be invariant, what is still a matter of debate.

The binding potential (BP) quantifies the equilibrium concentration of specific binding as a ratio to some other reference concentration. Particularly, the ratio at equilibrium of specifically bound radioligand to that of non-displaceable radioligand in tissue is referring to as $BP_{ND}$ (unitless) and equals the ratio $k_3/k_4$.

### 2.1.2 Input Curve

With rare exceptions only a fraction of the tracer in blood is exchangeable with tissue and thus needs to be determined from blood aliquots which are withdrawn from an artery during the whole study. This process is highly crucial for the quantification outcome. Please refer to our application note for a thorough discussion of all required processing steps and potential pitfalls.

A whole blood sample comprises tracer in the red blood cells and different tracer categories in the plasma: free unchanged tracer (parent), tracer bound to plasma proteins, tracer chemically changed by metabolite processes (metabolites). Depending on the tracer ligand, exchange processes are continuously going on between the different tracer pools in blood.

**Blood Sample Analytics**

The following blood analysis steps are required in quantitative PET studies to determine the concentration of unchanged tracer in plasma as the Arterial Input Function (AIF).

1. Separation of the plasma from the red blood cells by centrifugation.

Plasma represents about 60% of the total blood volume.
2. Further separation of tracer in plasma into tracer metabolites and unchanged tracer by filtration or HPLC (High Performance Liquid Chromatography).

3. Determination of the free fraction of unchanged tracer in plasma ($f_p$), i.e. the fraction of tracer not bound to plasma proteins. Often $f_p$ is difficult to measure and thus not explicitly used. Hereby, $f_p = 1$ is assumed, and the unchanged tracer concentration is regarded as the input function.

After the different tracer fractions in blood have been separated, it is necessary to calculate their activity concentrations. This is done by measuring them in a radioactivity counter and dividing the activity by the fractional volumes.

**Fitting of Blood Curves by Models**

Due to the experimental procedures involved the blood activity measurements are typically noisy and contribute a considerable amount of uncertainty to the analysis. Therefore it is often reasonable to fit analytical functions to the blood-related measurements, which is also applied for the interpolation between the samples, as well as for the extrapolation at late times when blood samples might be lacking.

The example below illustrates the blood model approach.

- The black triangles represent the activity of tracer (free, bound to proteins, metabolites) in the arterial plasma during the study. A tri-exponential function which was fitted to these measurements and shown as the black line.
- The green circle represents the measurements of the fraction "Unchanged tracer in plasma" to "Total tracer in plasma". A model was fitted to these fraction values and shown as the green line.
- The arterial input function (AIF) is obtained by the multiplication of the two model functions and represented by the red curve, implicitly assuming $f_p = 1$.

Note the sharp concentration changes at the early times after injection. To capture these changes adequately, rapid blood sampling is required during the first minutes after tracer injection.
2.1.3 Calculation of the Model Curve

The signal of a dynamic PET measurement represents the averaged activity in the image pixels at acquisition times \( t \) starting at tracer injection. It is described by the operational equation:

\[
C_{\text{Model}}(t) = (1 - \nu B) C_{\text{Tissue}}(t) + \nu B C_{\text{Blood}}(t) \equiv C_{\text{PET}}(t)
\]

This equation means that the activity concentration \( C_{\text{PET}} \) measured by PET in a certain tissue volume is composed of two contributions:

- tracer which has been extracted into tissue, where it has an instantaneous concentration of \( C_{\text{Tissue}}(t) \).
- tracer which is circulating within the blood, with a concentration \( C_{\text{Blood}}(t) \).

Hereby it is assumed, that the fractional volume \( \nu B \) is composed of small capillaries (2%-5%) filled with whole blood, and that the fraction \((1-\nu B)\) represents tissue. The blood activity \( C_{\text{Blood}} \) must be measured during the acquisition. In tissue, the tracer may be present in different spaces or forms (eg. free, specifically bound, non-specifically bound), which are described by the compartments of the model. All compartments contribute to the tissue signal, so that it is modeled by the expression

\[
C_{\text{Tissue}}(t_k) = \sum \frac{1}{t_{k_{\text{end}}} - t_{k_{\text{begin}}}} \int_{t_{k_{\text{begin}}} \to t_{k_{\text{end}}}} C_i(t) \, dt
\]

whereby

- \( C_{\text{Tissue}}(t_k) \) represents the average concentration of tracer in tissue during the acquisition \( k \) which starts at \( t_{k_{\text{begin}}} \) and lasts until \( t_{k_{\text{end}}} \)
- \( C_i(t) \) represents tracer concentration in compartment \( i \) at time \( t \). These expected concentrations are calculated from the differential equations using the current model parameters and the plasma input curve(s).

The operational equations used for other than compartment models are specified in the PKIN Model Reference section.

2.1.4 Fitting and Residual Weighting

Least Squares Optimization

The fitting methods available optimize the agreement between the measurements and the model curve. Effectively, they minimize the difference between them, whereby the difference is described by the Chi Squares criterion (cost function) below:
This expression implies that the squared residuals (measured value minus estimated model value) are multiplied by weights. To satisfy the requirements of least squares fitting, the weights $w_i$ should be related to the variance $\sigma_i^2$ of the measurements by

$$w_i = \frac{1}{\sigma_i^2}$$

In this case, and provided that the distributions of the measurement error are normal, the estimate obtained is the maximum likelihood estimate. Under the same premise it is also possible to obtain standard errors of the model parameters as the square root of the diagonal elements in the covariance matrix.

**Weighting of PET Data**

The variance of reconstructed PET data is dependent on many factors, including the duration of the acquisition, the time since the scan start which needs to be compensated by a decay correction, scatter and random correction, the sampling volume, the reconstruction method, etc. Therefore, PET variance models used in weighting of the residuals during fitting are always approximations.

In his presentation *Parameter Estimation: Least squares and why it gives you fits* (Handouts of Pharmakokinetic Course 2009) Richard E. Carson gives the following hints on using weights:

- Uniform weights (ordinary least squares fitting): Even if the data does not have uniform variance the estimates should be unbiased, but the parameter standard errors will be higher than they could be.
- Wrong weights: The greater the error in the weights, the larger the loss of precision in the parameter estimates (more important for $^{11}$C than $^{18}$F).

### 2.2 Reference Tissue Models

The measurement and analysis of the blood samples for kinetic modeling is an invasive and demanding procedure. Therefore, methods have been developed to obviate the need for invasive blood sampling. The solutions found replace the arterial input curve by an indirect input curve, namely the time activity curve of some reference tissue. Therefore they are called reference methods. Reference methods are not able to provide a full kinetic analysis. However, assuming certain relations between the kinetics of the tissue of interest and the reference tissue, they can provide valuable measures of interest.

Most of these reference methods are dedicated to reversibly binding neuroreceptor tracers. A reference tissue must be found which is devoid of receptors, and then it is assumed that the distribution volume of the non-displaceable compartment (free tracer in tissue and non-specific binding) is the same among the tissues. Under these assumptions a measure of the receptor concentration called binding potential can be calculated from the two time-activity curves.

**Model Structure**

The compartment models are usually based upon the following configuration:
In the model equations \( C'(t) \) represents the TAC from the reference region (\( k_3 = 0 \) in the 2-tissue compartment model), and \( C(t) \) the TAC from a receptor-rich region (\( k_3 > 0 \)).

However, the various reference methods differ in their mathematical approaches, and they show substantial differences with regard to noise sensitivity and processing speed. They are described in the PKIN Model Reference section.

### 2.3 Spectral Analysis (SA)

The closed-form solutions of compartment models involve the convolution of the input function with decaying exponentials. Based on this observation, a generalized technique called spectral analysis (SA) was introduced by Cunningham and Jones [66]. The operational equation of SA is given by

\[
C_{\text{Tissue}}(t) = \sum_{i=1}^{N} a_i C_p(t) \otimes e^{-\beta_i t} = \sum_{i=1}^{N} a_i B_i(t)
\]

that is, tissue uptake is modeled as a sum of \( N \) possible tissue responses. Due to the constraint of first order tracer kinetics, the coefficients \( a_i \) and the decay constants \( \beta_i \) must be non-negative. In practice, a discrete set of the decay constants \( \beta_i \) is selected which covers the physiologically reasonable range, typically logarithmically spaced in the range \([10^{-5}, 1] \text{sec}^{-1}\). The corresponding tissue responses

\[
B_i(t) = C_p(t) \otimes e^{-\beta_i t}
\]

are the basis functions of spectral analysis.

When fitting the operational equation above to a tissue TAC, the only unknowns are the coefficients \( a_i \), because only a pre-defined set of discrete \( \beta_i \) values is considered. Therefore, the problem is that of a non-negative linear least squares estimation (NNLS) with the constraint of non-negative coefficients. There is a well-known NNLS algorithm available [67], which allows readily calculating the optimal set of \( a_i \) coefficients. They allow calculating the model function and visualizing it together with the tissue TAC in the same way as for the compartment models.

An advantage of SA is the fact that no particular compartment structure is imposed. Rather, its result can be used to estimate how many kinetic tissue compartments can be resolved by PET. To this end, the results are plotted as a spectrum with the selected decay constants \( \beta_i \) along the x-axis (as the "frequencies") and the estimated coefficients \( a_i \) along the y-axis (as the "amplitudes"). Because of the large range, \( \log(\beta_i) \) is used in spectrum plotting rather than \( \beta_i \). The number of peaks in this spectrum corresponds to the number of distinct compartments. A peak appearing to the far left (low frequency, slow component) indicates irreversible trapping. A peak to the far right (high frequency, fast component) corresponds to kinetics indistinguishable from the input curve, thus to vascular contributions. Intermediate peaks represent compartments which exchange reversibly with plasma or with other tissue compartments [68].
The examples below used synthetic data generated using compartment models without noise and blood contributions. In the first example the simulated tissue TAC of a 1-tissue compartment model was fitted by SA with 500 basis functions. It is evident that, because of the discrete nature of the basis, the compartment was split into two neighboring frequencies. The adjacent amplitudes are usually summed to provide the combined peak height for a compartment.

In the next example an irreversibly trapping compartment was added (k3>0, k4=0) to the same first compartment. The corresponding peak is clearly seen to the left of the spectrum.

When changing kinetics to become reversible (k4>0), the second compartment appears as a second peak in the inner of the spectrum.
In general it is not possible to calculate the compartment rate constants from the spectral analysis outcome. However, an estimate of $K_1$ can be obtained as the sum of the peak amplitudes

$$K_1 = \sum_{i=1}^{N} a_i$$

and an estimate of the distribution volume $V_T$ as

$$V_T = \sum_{i=1}^{N} \frac{a_i}{\beta_i}$$

An additional information which can be calculated is the impulse response function

$$IRF(t) = \sum_{i=1}^{N} a_i e^{-\beta_i t}$$

The impulse response function completely describes the system, and the expected tissue TAC for any given input function and can simply be calculated by convolution with the IRF:

$$C_{PET}(t) = C_p(t) \otimes IRF(t)$$

In the practical application of spectral analysis potential problems were found: Caution should be applied when interpreting the number of peaks [66], and the error properties of the estimates are difficult to assess [69]. Several authors have proposed variants to overcome these problems [70,71], but in current practice spectral analysis is not yet frequently applied.
3 PKIN Configuration

There are two places where the functionality of PKIN tool is configured: the available models and their order in the list in the global PMOD configuration, and the other settings in the tool-specific configuration.

3.1 Global Configuration

Use the button on the PMOD ToolBox to open the general configuration dialog window. The PKIN models are available on the PKIN Models sub-panel of the USERS configuration.

Note that there are four different types of models, for Tissue, Blood|Plasma, Plasma/Whole-blood Ratio and Parent Fraction, each with a dedicated panel. The example above shows the Tissue panel which consists of the list of configured models, and tools to change the list. All of these panels behave in the same way.

A list entry shows in bold the model name as it appears in the PKIN interface, and in parentheses the actual file name of the model plug-in. The order how the models appear in the list and consequently in the PKIN application can be changed by selecting an entry and moving using the arrows to the right, or apply alphabetical sorting. The ? button opens the documentation with a short model explanation. To remove unneeded models from the model list just select an entry in the list and activate Remove. Conversely, the Add new button will show a dialog window listing the models currently not configured, so that they can be added again.

3.2 Local PKIN Configuration

The configuration of PKIN can be adjusted by the configuration button from the status line.
It opens the following dialog with the current definitions, which can be modified. The **Paths** tab allows setting the initial data paths of file-based loading as well as the file extension filters.

The **Display** pane gives access to the settings of the initial PKIN appearance and the appearance of the report page.

The **PKIN** pane shows the current models of tissue, whole blood, plasma and the parent fraction with their respective default parameters. Changing the parameters here and saving will establish new default parameters.

The next section allows specifying the time and value units of data loaded from text files. If the header lines in the data file do not contain valid units, these units are applied to the imported data.

The lower section serves for modifying the layout:

- **Curve controls position** allows displaying the curve controls to the **Right**, **Left** or at the **Bottom**. Their purpose is to optimize the layout depending on the display aspect ratio. **Left** and **Right** are recommended for wide screen systems, otherwise **Bottom**.

- **Center model parameters display**: If this box is enabled the parameters area is centered vertically within the available space.
- **HD curve model display**: If this box is checked the model curves are calculated for display purposes at many intermediate points, not just at the measurement times. The result is a smooth appearance of the model curves. However, when exporting the model curves, these interpolated points may be disturbing. Note that for the fitting only the original acquisition times are used.
4 P kin Data Processing

4.1 General Assumptions in PKIN

The following is generally assumed:

- All loaded data has been decay corrected to the same time point. This means that the PET scanner and the blood sampling times must have been synchronized. An exception is the processing of dosimetry data, which is preferably not decay corrected.
- The acquisition time intervals may have gaps, but must not overlap.
- The whole blood curve \( C_{\text{Blood}}(t) \) represents the activity concentration of the tracer and all its metabolites in whole blood samples.
- Most PET models require the input curve \( C_{\text{pl}}(t) \), namely the authentic tracer in arterial plasma which can exchange with tissue. Depending on how the plasma-related information is prepared and loaded, the input curve needs to be calculated differently.
- In compartment models the whole-blood signal contribution can optionally be considered in the operational equation by
  \[
  C_{\text{Model}} = vB \cdot C_{\text{Blood}} + (1-vB) \cdot C_{\text{Tissue}}
  \]
  where \( v_B \) represents the fractional volume of the blood space in the VOI, \( C_{\text{Blood}}(t) \) the whole blood concentration, and \( C_{\text{Tissue}}(t) \) the summed concentrations in all tissue compartments.
- The results of non-cardiac flow models are returned per cm³ tissue. To convert to flow per g tissue the values must be divided by the tissue density.

Exceptions to these rules are specified in the description of the individual models.

4.2 Processing Overview

Data processing of studies with PET or SPECT tracers typically consists of the following parts:

| 1. Data Loading | ▪ Create a new workspace (tab) in the PKIN tool.  
  |                | ▪ Load the whole-blood curve (total activity concentration of tracer in the blood samples). 
  |                | ▪ Load the plasma activity curve. Depending on the preparation of the blood data the plasma activity curve can represent (1) the unchanged tracer (parent) in plasma, or (2) total tracer in the plasma of the blood samples including the metabolites. In the latter case a metabolite correction has to be performed in PKIN. 
  |                | ▪ An alternative to loading the plasma activity curve is loading the plasma fraction curve. In this case the total plasma activity is calculated by multiplying whole blood activity with the plasma fraction. 
  |                | ▪ Load the parent fraction curve (fraction of unchanged tracer in plasma). This is only necessary if the loaded plasma curve represents total tracer activity in plasma including metabolites. 
<table>
<thead>
<tr>
<th></th>
<th>▪ Load the average time-activity curves of one or multiple tissue regions.</th>
</tr>
</thead>
</table>
| 2. Input Curve Configuration | ▪ Define interpolation functions (also called models) for the different types of blood data and fit them to the data. The default is linear interpolation between measurements, and a parent fraction identical to 1. 
<p>|                | ▪ The input curve is obtained as the multiplication of the plasma model with the parent fraction model. |
| 3. Kinetic Model Configuration | ▪ Select a region with a &quot;typical&quot; time-activity curve (TAC) from the Region list. |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>4. Kinetic Model Fitting</td>
<td>• Select a simple model from the <strong>Model</strong> list.</td>
</tr>
<tr>
<td></td>
<td>• Enable the parameters to be fitted by checking their boxes; unchecked parameters remain at the initial values entered.</td>
</tr>
<tr>
<td></td>
<td>• Select a weighting scheme of the residuals; variable or constant (default) weighting is available.</td>
</tr>
<tr>
<td></td>
<td>• Fit the model parameters by activating the <strong>Fit model, current region</strong> button.</td>
</tr>
<tr>
<td></td>
<td>• Consult the residuals to check whether the model is adequate; there should ideally be no bias in the residuals, just random noise.</td>
</tr>
<tr>
<td></td>
<td>• If the model fit is fine, configure <strong>Copy the model to all regions</strong> to <strong>Model &amp; Par</strong> and activate the button to establish the same initial model configuration for all TACs. Then activate <strong>Fit model, all regions</strong> to adjust the parameters to the regional measurements.</td>
</tr>
<tr>
<td></td>
<td>• Check the fit result for all the TACs by stepping through the regions. Note that mouse scrolling is supported when placing the cursor on top of the <strong>Region</strong> list.</td>
</tr>
<tr>
<td></td>
<td>• If the model is not yet fine, test more complex models.</td>
</tr>
<tr>
<td>5. Kinetic Model Comparison</td>
<td>• Switch between compartment models of different complexity and fit. The parameters are either maintained for each model type, or converted, according to the <strong>Parameter conversion</strong> setting in the <strong>Extras Panel</strong> panel.</td>
</tr>
<tr>
<td></td>
<td>• The results of the different fits are stored in the <strong>Model History</strong> which can be used for inspecting the results per region.</td>
</tr>
<tr>
<td></td>
<td>• Check the residuals for judging model adequacy.</td>
</tr>
<tr>
<td></td>
<td>• Check the different criteria on the <strong>Details</strong> tab (Schwartz Criterion SC, Akaike Information Criterion AIC, Model Selection Criterion MSC) to decide whether a more complex model is supported by the data.</td>
</tr>
<tr>
<td></td>
<td>• Check for parameter identifiability. As an indicator of the parameter identifiability standard errors (%SE) are returned from the fit. They should remain limited for all relevant parameters. Additionally, <strong>Monte Carlo</strong> simulations can be performed to obtain distribution statistics of the parameter estimates.</td>
</tr>
<tr>
<td></td>
<td>• If justified by physiology, try to improve the stability of parameter estimation by enforcing common parameters among regions in a <strong>Coupled fitting</strong> procedure.</td>
</tr>
<tr>
<td></td>
<td>• Compare the outcome of compartment models with that of other models, such as reference models or graphical plots.</td>
</tr>
<tr>
<td>6. Saving</td>
<td>• Save all model information together with the data in a composite text .km file which allows restoring a session including the fitting history.</td>
</tr>
<tr>
<td></td>
<td>• Save a summary of all regional parameters in an EXCEL-ready text file .kinPar.</td>
</tr>
<tr>
<td>7. Batch Processing</td>
<td>• Lengthy calculations such as coupled fits with many regional TACs or Monte Carlo simulations can be run in a <strong>batch job</strong>. The results are saved both as .km files and in a .kinPar text file.</td>
</tr>
</tbody>
</table>
4.3 Starting the Kinetic Modeling Tool

The kinetic modeling tool is started with the Kinetic button from the PMOD ToolBox or by directly dragging kinetic modeling (.km) files onto the above button.

4.3.1 User Interface

The user interface is organized as illustrated below. The left part of the display visualizes the data, the model and the fit. The green squares represent the tissue measurement values (tissue time-activity curve, TAC), the red circles the input curve, the yellow circles the blood spillover curve (whole blood time-activity curve), and the blue line the calculated model curve with the current model configuration shown to the right. All curves can be shown or hidden using the check boxes in the control area. Per default, the blood curves are switched off as they usually peak on a much higher value.

The lower curve display shows the weighted residuals, i.e. the difference between the measurement and the model curve multiplied by a weighting factor. Unweighted residuals can also be shown.

The right part gives access to the different models and operations which are described in the following sections.

The Taskbar to the right provides shortcuts to loading and saving, as well as the configuration of frequently used model filters and fitting options.

4.3.2 PKIN Menu

The Kinetic menu is located in the lower left corner and is mainly used for data operations such as loading, saving or closing. Additionally
Loaded data sets can be added into new workspaces, so multiple data sets can be available simultaneously and selected for processing using named tabs.

<table>
<thead>
<tr>
<th>Feature</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Acceptance Test</strong></td>
<td>Self test of the program testing 26 model configurations. It is executed automatically the first time PKIN is started.</td>
</tr>
<tr>
<td><strong>Batch Mode</strong></td>
<td>Open interface for <strong>batch processing</strong>.</td>
</tr>
<tr>
<td><strong>Aggregate Parameters</strong></td>
<td><strong>Aggregate</strong> the results from different KM Parameter Files (.kinPar) in a single table for statistical analysis.</td>
</tr>
<tr>
<td><strong>Study Compare Dialog</strong></td>
<td>Open a PKIN dialog window with the same data so that two model configurations can be examined side by side in separate windows.</td>
</tr>
<tr>
<td><strong>Coupled Studies Dialog</strong></td>
<td><strong>Fit multiple data sets</strong> together, keeping selected parameters synchronized.</td>
</tr>
<tr>
<td><strong>Create Synthetic PXMOD Study</strong></td>
<td>Generate a <strong>synthetic image series</strong> which has a different combination of model parameters in each pixel.</td>
</tr>
<tr>
<td><strong>Create Synthetic KM Study</strong></td>
<td>Generate a <strong>synthetic modeling data set</strong> from the current data with ideal tissue TACs and potentially modified timing.</td>
</tr>
<tr>
<td><strong>Create Parametric Maps</strong></td>
<td>If the TACs have been transferred from PVIEW to PKIN and the images are still open in PVIEW, open a <strong>dialog window</strong> for assembling the fitted parameters into images. If average TACs were transferred, the VOIs will be homogeneous showing the regional result. If pixel-wise TACs were transferred, the VOIs will be filled by parametric maps.</td>
</tr>
<tr>
<td><strong>Clear all KM Data</strong></td>
<td>Close by all workspaces.</td>
</tr>
</tbody>
</table>
Close Current Study | Close currently selected workspace.
New Study | Open a new, empty workspace.
BIDS | WIP implementation for creating data sets according to the emerging Bioimaging Data Structure.
Save all Model Curves | Save the model curves of all regions in a tab-delimited text file.
Save all Tissue TACs | Save the tissue TACs of all regions in a tab-delimited text file.
Load Tissue TACs | Load the tissue TACs from a tab-delimited text file.
Load Authentic Fraction | Load a fraction curve representing the ratio of parent (aka authentic, intact, unchanged) tracer to total tracer in plasma.
Load Plasma/WB Ratio | Load a fraction curve representing the ratio of total tracer in plasma to whole-blood.
Load Plasma Activity | Load a curve representing total tracer in plasma.
Load Whole Blood Activity | Load a curve representing tracer in whole-blood samples.
Save KM Parameters File | Save the parameters of all regional tissue models in a tab-delimited text file (.kinPar).
Save KM File | Save all data of the current workspace with all model configurations in the history as a proprietary KM text file (.km).
Load KM File(s) | Load a number of selected KM files into separate workspaces, clearing already loaded data.
Add KM File(s) | Load a number of selected KM files into additional workspaces, keeping existing data.

4.3.3 Curve Layout

The curve display supports the visualization of the different loaded measurements as well as the calculated model curves. In PKIN the curves available for display depend on the selected tab (Tissue, Whole Blood, Plasma, Metabolites).

Curves with selected Tissue Tab

After loading all data, the time-activity information of the current region is shown in the curve display in a default layout. Note that the blood curves are initially switched off as they often have a much higher dynamic range and compress the tissue curve which is of main interest.

The curves available for display in the upper plotting area are:

- the green TAC values representing the tissue TAC of the selected region, in the example the Whole Brain TAC;
- the red **Parent input curve model** which is used as the input curve for the model calculations;
- the yellow **Whole blood model** used for blood spillover corrections;
- the blue **Model Curve** which results from evaluating the current model configuration with the input curve.

Per default, only the **TAC** and the **Model Curve** are enabled for display by the check box in the control area. Depending on the context, there are additional curves available for display but hidden (box not checked). In the example below the compartment concentrations $C_1$ and $C_2$ of the 2-Tissue compartment model as well as a finely resolved **Model Curve HD** could also be shown by checking their boxes.

![Data and fit to tissue curve](image)

The smaller curve display at the bottom visualizes the difference between the measured TAC and the model curve. Note that for screens with limited height it may be convenient switching the configuration to the layout with the curve controls to the right instead of below.

The general manipulations available in the PMOD curve display are explained in the next section.

### 4.3.4 General Curve Display Functionality

A common curve display object is used in all PMOD tools. It consists of a curve area and a controls area underneath.
In some situations the control area may initially be hidden. In this case, please open the context menu by right clicking into the plot area, and select Hide/Show curve controls.

Curve Area
The curve area shows the curves which are enabled for display. There is always an active curve, which is shown in bold. A curve can be made active by holding down the CTRL key and clicking at one of its points, or by pushing its button in the controls area as illustrated with the Model Curve above.

The definition of the active curve is relevant for the tools which interrogate the curve values:
- There are two small handles at the top of the curve area: a little rectangle to the left, and a line to the right. These are handles which can be moved left/right using the mouse, and the gray vertical lines move with them. The values at the top center of the curve area represent the interpolated (x/y) values of the active curve at the location of the handles. To get the measurements of a different curve just CTRL+Click at that curve to get the values updated.
- Only in some curve displays: When the cursor is brought close to a point of the active curve, its x/y value pair is shown at the upper left of the curve area.

To zoom into an area of the curve just click the left mouse button to the corner of the area of interest and drag a rectangle. After releasing the mouse button the display is zoomed into the defined rectangle. An alternative is to define the axes Range in the context menu. A single mouse click into the curve area is sufficient to reset the zoom.

Context Menu
By clicking the right mouse button into the curve area a context menu with some additional options can be opened.
The functions are:

**Use restricted value range**
Set the range of the x- and y-axis by entering a numeric value.

If the **Use restricted value range** box is checked, the range is maintained until the configuration is opened again and the check removed. Otherwise, a single click resets the range to the default.

**Reset interactive zoom**
To reset the curve range to the default full range. It is grayed if the display is not zoomed or the range is fixed.

Mouse operation: single click into curve area.

**Set closest curve active**
Selects the curve nearest to the point clicked with the right mouse button when opening the context menu.

Alternative mouse operation: CTRL+Click at a curve.

**ON/OFF closest point**
Disable a measurement of the active curve. This is reflected by setting the symbol to gray.

In the context of model fitting measurements marked in gray are regarded as outliers and not considered when evaluating the cost function.
Alternative mouse operation: select the relevant curve to active, then **SHIFT+Click** at measurement.

**Switch ON/OFF all points in visible area**

In combination with zooming this option allows to quickly disable/enable a section of the active curve.

**Hide/Show curve controls**

Allows hiding the controls if the curve display area is small, and to show them again.

**Show/Hide curve names**

Controls, whether a box with the visible curve names is shown in the upper right. Note that if the available vertical space does not allow listing all names, the box is not shown.

**Show/Hide active curve max value**

**Hide/Show grid lines**

Controls the display of the grid lines in the plot.

**Hide/Show vertical rulers**

Controls whether the rulers are shown which allow measurements on the active curve.

**Plot properties**

With this entry a configuration dialog is opened for setting the annotation **Font size**, for enabling **Curve antialiasing** (smooth curve appearance), and for excluding the horizontal and vertical zero lines.
Show in separate window
With this option, the curve display can be opened in a separate, large window to closely examine the plot.

Save multiple curves
Allows saving the numeric data of the curves in a single or separate text files.

Value table of visible curves
Opens a dialog window which shows the numeric values of all visible curves in a dialog window. The window contents can be copied to the Clipboard and pasted to a different application.

Switch all curves ON/OFF
To quickly change the visibility of all curves. When switching all off, the active curve is still shown.

Curve Control Area
The control area lists the curves which are available for display. There are several elements to modify the curve appearance:

- **Show/Hide**
  To show/hide a curve check/uncheck the visible box.

- **Active curve**
  To set a curve to active click at the active button, or directly CTRL+Click on the curve itself. The line/symbols get bold.

- **Style**
  The list selection can be used to change the style of a curve:

  - Normal line
  - Bold line
  - Dots only
  - Dots and line
  - Filled area

Further useful interface elements:
- **Linear**
  Changes the shape of the lines defined by the measurements.
  Note that calculations are not based on the display representation of a curve.

  - Cubic
  - B-Spline

- **Save**
  Saves the numeric data of the curve as a text file with two columns. These files obtain a .crv suffix and can easily be opened in Excel or a text editor.

When working with a database the curve can be attached to a particular image using the Attach to Patient (Serie) in the appearing dialog.

If this button is enabled, each curve is normalized to its own maximum and shown as percent values. This mode is helpful for
compare the areas of the curve when the dynamic range of the curves is very different.

Creates a capture of the curve area. The captured image can be saved as a JPEG, TIFF, or DICOM file. It can also be copied to the Clipboard to paste it into some office application.

4.4 Data Loading/Saving

4.4.1 File-based Data Import

There are slightly different formats for the time-activity data of blood and tissue. With blood data it is assumed that instantaneous sampling was performed at well-defined time points. With tissue data it is assumed that the measurements represent the average tracer concentration during a certain observation period, the PET frame duration defined by start-time and end-time.

4.4.1.1 Data Units

Timing Units

The following timing units are supported.

- [milliseconds]
- [seconds]
- [minutes]
- [hours]

During data loading, a conversion of the timing information to seconds is performed.

Value Units

The unit [1/1] is generic. In this case the user has to take care that all data used for modeling have been translated into compatible values.

PET data typically arises in concentration units, which are a requirement for modeling. The following activity concentration units are supported.

- [Bq/ml], [Bq/cc], [kBq/cc], [MBq/cc]
- [uCi/cc], [nCi/cc]
- [kcps/cc]

Note: During PET data loading, the unit specification will be used to convert the specified activity concentration values into the internal representation of kBq/cc. All activity concentration data will be displayed in kBq/cc in the user interface.

The formats for CT and MRI data are [HU] and [1/1], respectively.

The unit string in square brackets should immediately follow the column headers as described in the following sections. If units are not defined or not recognized, it is assumed that the units specified in the PKIN configuration apply.

4.4.1.2 Import of Blood Data

PET tracers are usually applied by intravenous injection and then brought to the tissue of interest by circulation. Often, a part of the tracer may be bound to red blood cells. Another part may be processed in organs and end up as labeled metabolites circulating in blood plasma. The remaining
category of tracer in plasma is the unchanged (authentic) tracer (parent) which is available for exchange with tissue and represents the input curve relevant for modeling.

In PKIN, the following four types of blood data are supported to model the contributions of the different forms of tracer in blood to the expected PET signal:

1. The concentration of all forms of tracer in Whole Blood samples: As the average signal from a VOI always contains a fraction of signal from blood vessels and capillaries, the concentration in whole blood is needed for modeling the blood content in the expected PET signal.

2. The concentration of all forms of tracer in plasma (Plasma Activity): Although the unchanged tracer in plasma is required as the input curve of the models, it is common practice to measure the tracer activity of the whole plasma sample.

3. The fraction of unchanged ligand in plasma (Parent Fraction): As a result of the metabolite analysis of plasma samples, the relative concentrations of the unchanged tracer and the metabolites are known. This information is represented by the ratio of unchanged tracer to total tracer in plasma. The parent fraction is expected to start with a value of 1 at the time of injection (all tracer is unchanged), and gradually decreases as the metabolites build up. Since plasma analysis is often experimentally complex and therefore error-prone, it is advisable to fit a smooth curve to the measured parent fraction.

4. The Plasma Fraction: An alternative to using the measured plasma activity is the use of the ratio of plasma activity to whole blood activity, called the "Plasma Fraction". If the plasma fraction is known it can be multiplied with the whole blood activity to obtain the plasma activity. For tracers without metabolites this obviates the need for the actual blood analysis.

**Whole Blood and Plasma Activity**

The whole blood and plasma activity concentrations must be prepared in text files, and can be loaded using Load Whole Blood Activity and Load Plasma Activity from the Kinetic menu. Such text files with the blood data can be prepared for example in MS Excel and then saved as tab-delimited or csv separated text files. There are two variants of the format:

1. Separate files for whole blood and plasma activity. In this case separate files are prepared with a header line, the sample time in the first column and the sample value in the second column as illustrated below:

   ![Whole Blood Activity Example](image1)

2. A composite file containing both whole blood and plasma activity. Note that in this case the keywords sample-time, plasma and whole-blood are required to define the meaning of the
Plasma and Parent Fractions

The plasma and parent fractions need to be prepared in a similar file and loaded using **Load Plasma/WB Ratio** or **Load Authentic Fraction** from the **Kinetic** menu.

**Notes**

1. The header line is mandatory in the format shown above with valid units in square brackets after the column headers.

2. If no whole blood data is loaded into PKIN, the plasma concentration is used for blood spillover correction.

3. If the activity of unchanged tracer in plasma is loaded instead of the total plasma activity, no correction with the parent fraction is required. No further action is required in this case, because per default it is assumed that the parent fraction equals the constant of 1.

4. There are a few models which require two input curves, and in principle models with up to 10 input curves can be handled in PKIN. For these models the loading sub-menus contain appropriately labeled entries.

### 4.4.1.3 Import of Tissue Time-Activity Data

Tissue time-activity curves are similarly organized in tab-delimited text files. The frame start and end times are arranged in the first two columns. Next come columns with the measurements from different tissues.

**Note:** The header line is mandatory for providing the units and the region names. The TAC value units are specified after the heading of the second column.
Tissue time-activity curves are loaded using **Load Tissue TACs** from the **Kinetic** menu.

**Note:** This data import assumes that all curves have the same timing. If it is differing, the timing has to be edited after loading using the **Edit Data** facility, or the data need to be arranged in the **Composite Format** which supports individual timing.

### 4.4.1.4 Composite Import Format for all Data (.kmData)

The composite format allows including all types of blood and tissue data into a single tab-delimited text file. The advantage is that all data can be prepared externally and loaded at once, and that such files can directly be used for batch processing.

#### File Start

The file starts with the two lines

```
# APPLICATION
KINETIC MODELLING DATA
```

It is followed by a sequence of sections for the data and other related information.

#### Demographic and Study Data

The demographic information and study-related data can be defined in a section headed by the `# PATIENT` tag. Each line starts with a tag in capital letters which is followed by a tab, and then the corresponding value.

<table>
<thead>
<tr>
<th>Tag</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>PATIENT_NAME</td>
<td>PKIN</td>
</tr>
<tr>
<td>PATIENT_ID</td>
<td>Dyn. CPPFX bolus &amp; MRI</td>
</tr>
<tr>
<td>PATIENT_BIRTH_DATE</td>
<td>01.01.1981</td>
</tr>
<tr>
<td>PATIENT_SEX</td>
<td>M</td>
</tr>
<tr>
<td>PATIENT_SIZE</td>
<td>1.79</td>
</tr>
<tr>
<td>PATIENT_WEIGHT</td>
<td>75</td>
</tr>
<tr>
<td>PATIENT_REFERING_PHYSICIAN</td>
<td>Dr. B</td>
</tr>
<tr>
<td>PATIENT_INSTITUTION</td>
<td></td>
</tr>
<tr>
<td>PATIENT_ACQUISITION_DATE</td>
<td>200608021301226</td>
</tr>
<tr>
<td>PATIENT_SERIES_DATE</td>
<td>200608031301226</td>
</tr>
<tr>
<td>PATIENT_STUDY_DATE</td>
<td>200608011301226</td>
</tr>
<tr>
<td>PATIENT_STUDY_ID</td>
<td>CPPFX Bolus</td>
</tr>
<tr>
<td>PATIENT_ACCESSION_NO</td>
<td></td>
</tr>
<tr>
<td>PATIENT_STUDY_DESC</td>
<td>CPPFX Bolus</td>
</tr>
<tr>
<td>PATIENT_SERIES_DESC</td>
<td>Dynamic PET</td>
</tr>
<tr>
<td>PATIENT_BODY_PART</td>
<td>BRAIN</td>
</tr>
<tr>
<td>PATIENT_LATERALITY</td>
<td></td>
</tr>
<tr>
<td>PATIENT_RADIOINUCLID</td>
<td>C-111Al/5RT/18°Fluorine/</td>
</tr>
<tr>
<td>PATIENT_HT_TIME</td>
<td>6586.2</td>
</tr>
<tr>
<td>PATIENT_UNIT</td>
<td>kBq/cc</td>
</tr>
<tr>
<td>PATIENT_ACTIVITY_PRE</td>
<td>280</td>
</tr>
<tr>
<td>PATIENT_ACTIVITY_POST</td>
<td>0</td>
</tr>
</tbody>
</table>

#### Comments
Comments can be included for describing the data and recording important facts. The # DESCRIPTION section allows specifying text which will be imported with the data and can be shown in the user interface. The three elements to enter in a line are the Date of the comment, the User who edited it, and the actual comment text as the Description.

Further comments can be added as lines preceded by a // in a # NON_PMOD_COMMENTS section. They are only for documentation purposes of the raw data and not interpreted by PKIN.

Model Specification
A tissue model can optionally be specified in a # MODELS section.

The Model entry corresponds to the class name, and the Name entry to the string shown in the model list. Both can be determined in the configuration utility as illustrated below.

For reference models the reference region can be specified in by an additional line with the REFERENCE_REGION tag.

Note that the region number (starting from 0) is relevant, not the region name.

Data Sections
Each data part to be included is specified in a section starting with a line # DATA followed by a header line of the actual data. For blood data the following format is supported. Note the data units
which are specified in the brackets directly after the column names in the header line.

```
<table>
<thead>
<tr>
<th># DATA</th>
<th>sample-time[minutes]</th>
<th>whole-blood[MBq/cc]</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.75</td>
<td>1.67E-05</td>
<td>4.818734894</td>
</tr>
<tr>
<td>1.5</td>
<td>2.25</td>
<td>21.04629028</td>
</tr>
<tr>
<td>3.02</td>
<td>3.75</td>
<td>27.0131493</td>
</tr>
<tr>
<td>4.52</td>
<td>5.25</td>
<td>12.99452924</td>
</tr>
<tr>
<td>7.98</td>
<td>6.1</td>
<td>8.360559731</td>
</tr>
<tr>
<td>10.07</td>
<td>4.48897549</td>
<td>3.660262888</td>
</tr>
</tbody>
</table>
```

For tissue data, it is assumed that the first two columns define the acquisition start and end times, whereas the following columns contain the data for the different tissue regions.

```
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>60</td>
<td>2.2</td>
<td>0.457304936</td>
<td>2.2</td>
<td>0.49</td>
<td>2.29</td>
</tr>
<tr>
<td>60</td>
<td>120</td>
<td>2.2</td>
<td>1.393019209</td>
<td>2.2</td>
<td>1.69</td>
<td>2.29</td>
</tr>
<tr>
<td>120</td>
<td>180</td>
<td>2.2</td>
<td>1.674894667</td>
<td>2.2</td>
<td>1.99</td>
<td>2.29</td>
</tr>
<tr>
<td>180</td>
<td>240</td>
<td>2.2</td>
<td>2.120504157</td>
<td>2.2</td>
<td>2.19</td>
<td>2.29</td>
</tr>
<tr>
<td>240</td>
<td>300</td>
<td>2.2</td>
<td>2.372558724</td>
<td>2.2</td>
<td>2.39</td>
<td>2.29</td>
</tr>
<tr>
<td>300</td>
<td>420</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
```

Each region is represented by a column with header tac.name, where "name" will be shown as the region name in PKIN. Two regions will result from the example above, with names `first-region` and `second-region`. Optionally, the volumes corresponding to the TAC can be defined in additional columns, indicated by the header `volume.name`. The volume unit is cc, so there is no need to specify it.

Multiple blood and tissue data sections can be included in a composite data file, for instance to handle TACs with different timing. The following key words are parsed in the header lines.

<table>
<thead>
<tr>
<th>Key Word</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>sample-time</td>
<td>Timing of blood samples.</td>
</tr>
<tr>
<td>whole-blood</td>
<td>Activity of whole blood samples, to be used for spillover correction. May also be used for calculating the plasma activity if a plasma fraction is loaded.</td>
</tr>
<tr>
<td>plasma.1</td>
<td>Plasma activity of the first input curve, the second, etc. Plasma activity may represent the activity of parent, if the metabolite correction has been done outside of PKIN, otherwise the activity of parent and metabolites. In the latter case a parent fraction has to be loaded for the metabolite correction within PKIN.</td>
</tr>
<tr>
<td>plasma.2</td>
<td>Plasma activity of the first input curve, the second, etc. Plasma activity may represent the activity of parent, if the metabolite correction has been done outside of PKIN, otherwise the activity of parent and metabolites. In the latter case a parent fraction has to be loaded for the metabolite correction within PKIN.</td>
</tr>
<tr>
<td>plasma-fraction.1</td>
<td>Plasma fraction of the first input curve, the second, etc. These fractions will be multiplied with <code>whole-blood</code> for the calculation of <code>plasma.1</code>, <code>plasma.2</code> etc. Supported units: [1/1], [%].</td>
</tr>
<tr>
<td>plasma-fraction.2</td>
<td>Plasma fraction of the first input curve, the second, etc. These fractions will be multiplied with <code>whole-blood</code> for the calculation of <code>plasma.1</code>, <code>plasma.2</code> etc. Supported units: [1/1], [%].</td>
</tr>
<tr>
<td>parent-fraction.1</td>
<td>Parent fraction of the first input curve, the second, etc. These fractions will be multiplied with <code>plasma.1</code>, <code>plasma.2</code> etc. for the calculation of the input curves. Supported units: [1/1], [%].</td>
</tr>
<tr>
<td>parent-fraction.2</td>
<td>Parent fraction of the first input curve, the second, etc. These fractions will be multiplied with <code>plasma.1</code>, <code>plasma.2</code> etc. for the calculation of the input curves. Supported units: [1/1], [%].</td>
</tr>
<tr>
<td>start</td>
<td>Start time of the PET/SPECT acquisition frames. The time units of the tissue TACs are specified after <code>begin</code>, eg. <code>end[seconds]</code>.</td>
</tr>
</tbody>
</table>
| end            | End time of the PET/SPECT acquisition frames. Note: if the values of the tissue TACs have common units they can be
File Format and Loading

The information described above can be saved in a tab-delimited text file (.kmData) or an XLS (Excel 2003, notxlsx) file and loaded with the Load KM File entry from the Kinetic menu. Alternatively, the file can be dragged and dropped onto the PKIN window or the PMOD ToolBox.

4.4.2 Data Import from VOIs

An important usage of the VOI analysis is the generation of time-activity curves (TAC) for subsequent kinetic modeling. This can easily be achieved in PVIEW by the following steps:

**VOI Definition**

The image data is loaded as a dynamic series with the correct acquisition times and the correct input units. This is important, because otherwise the acquisition start/end times in kinetic modeling will be wrong, and the TACs may be different in magnitude with respect to the blood data. Such problems result in erroneous model parameters.

In dynamic image series there is generally not enough anatomical information to delineate VOIs. Often, averaging of a subset of the acquisition frames resolves the problem. The VOIs are then delineated in the summed images, transferred to the dynamic images, and optionally saved to a file.

**TAC generation**

Switch the tool to the dynamic study, and activate the button

A dialog window appears which is organized in three panels:

1. The left area allows defining the proper type (TISSUE, WHOLE BLOOD) of the calculated TACs and selecting the regions to be sent to the Kinetic modeling tool.
2. The central area shows the TACs.
3. The right area controls the displayed TACs. Additionally, the +- buttons allow for simple arithmetic operations with the curve values before transfer to PKIN.
Average TACs
The standard and default procedure is to use the signal average in the VOIs, corresponding to the radio button **Averaged VOI TAC**. With this setting, not only the average is calculated and transferred, but also the standard deviation which may be used for weighted fits in PKIN.

Voxel-wise TACs
When the transfer mode is set to **TAC of each VOI voxel**, the individual voxel-wise TACs are transferred. There is a maximum number of TACs which will be considered for the transfer, per default set to **10'000**. The number of voxels in the selected VOIs is shown in brackets in the control section. Naturally, the standard deviation is zero in this case.

TAC Value Operations
The calculated TAC values can be modified in two ways.

1. Every individual curve has a ± button which allows multiplying the values with a factor, dividing, adding or taking the logarithm.

2. All generated curves can be scaled with a factor defined by the **Global scale factor to kBq/cc** option.

TAC Time Shifting
If image data acquisition requires multiple table positions (e.g. dynamic whole-body PET), the timing between VOIs may differ. The timing relative to the injection is crucial for modeling and also for dosimetry. Therefore, the **Delay by VOI location** option is supported.
with two alternatives:

- **Stop & Go** corresponds to the standard acquisition mode with the table halting in adjacent, overlapping field of views. As there is usually not enough information in the image headers to know for each slice the exact timing, the timing of the different table positions is obtained by dividing the acquisition duration by their number.

- **Continuous bed movement** assumes that the patient is scanned while the table is continuously moving. Here the axial scanner field of view is required for approximating the time when a VOI is scanned.

Note that the timing derived from these settings can only be approximate and may need correction. The resulting timing of the TAC curves is updated in the curve window whenever one of the settings is changed. Depending on the scanning, the time shifts may be applied in the wrong direction. For instance, the bladder curve may start before the brain curve although the data was acquired in head-in position. To rectify such a case there is a switch **Time increases with slice number/**Time decreases with slice number.**

**Dosimetry Model**

If the flag **Dosimetry model** is enabled, the **OLINDA Residence Times** model will be set after the TAC data have been transferred to the PKIN tool together with the **Total body mass** of the patient. This information is extracted from the demographic information in the image, and can be edited if it is not present.

**Decay Correction**

If the **Decay correction to Injection time** flag allows correcting the TACs to an injection time which may be before the first acquisition. If the **Source data is decay corrected**, only a scaling factor is applied. Otherwise, a time-dependent correction is used.

**Data Transfer**

The **Send** buttons initiate the transfer of the activity curve data to the PKIN tool. Selecting **Send[SET]** transfers the TACs to the currently selected tab in an open PKIN tool. With the **Append TAC Data** box checked, the curves are appended as new regions to the existing data, otherwise the current data is over-written. **Send[ADD]** first creates a new tab in PKIN, to which the data is added. If PKIN is not running, the tool is first started and the data added.

Both the average value and the standard deviation within the VOIs are transferred, as well as patient and study related information. The standard deviation may be used for weighted fits in PKIN.

**4.4.3 Data Saving**

Data can be saved in various formats using the following **Kinetic** menu entries:

<table>
<thead>
<tr>
<th>Save KM File</th>
<th>Saves all data and the configuration in a comprehensive text file with suffix .km. Loading a .km file restores the state of the previous processing with the exception of the display settings.</th>
</tr>
</thead>
</table>

Note that the **model fitting history** is also saved/retrieved.
Save KM Parameters File
Generates and saves a summary of the model parameters with their standard error in all regions.

**Note:** The results of the latest Fit in the model fitting history are exported, rather than the currently viewed parameter set which may be manually changed.

Result is a tab-delimited text file with extension .kinPar, which is readable with any text editor and with numerical programs such as Excel. There is a Save and an Append sub-menu, the latter for combining results of several studies in a single file. kinPar files can be aggregated for performing statistical analyses. Saved model parameters can be inspected using the View Parameters button from the taskbar to the right.

Save all Tissue TACs
Saves just the regional time-activity data in a multi-column text file with .tac extension. This option may be helpful to export TAC data for visualization in a different program.

Another useful application of Save/Load Tissue TACs is appending a TAC from a different tissue: first save the current TACs, add the TAC of an additional region as a new column in Excel, then load the .tac file again.

Save all Model Curves
Exports the tissue model curves of all regions into a text file. Note that these curves are not interpolated between the frame mid-times. To obtain smoother curves please use the Create Synthetic KM Study menu item.

**Note:** Units of the saved data TACs are always [kBq/cc] and [seconds].

### 4.5 Preparation of Blood Data

**Types of Blood Information**
PKIN supports four types of blood data which together allow computing the input curve.

1. The tracer activity concentration in whole blood. This information is used for the spillover term in the compartment models as well as in combination with the plasma fraction.
2. The total activity concentration of tracer in the plasma part of the blood.
3. The plasma fraction (concentration ratio of tracer in plasma to tracer in whole blood). The plasma fraction is multiplied with the whole blood concentration to calculate the total concentration of tracer in plasma.
4. The parent fraction (concentration ratio of unchanged tracer to total tracer in plasma). The parent fraction is multiplied with the total plasma concentration to calculate the input curve.

Note that the plasma options 2. and 3. above are excluding each other. If plasma activity has been loaded, the plasma fraction option cannot be used. On the other hand, if the plasma fraction has been loaded, the plasma activity option cannot be used.

**Input Curve Calculation**
The input curve is defined as the activity concentration of parent tracer in the arterial plasma. Depending on the imported data, the input curve needs to be calculated differently. Some typical scenarios are:

- **External metabolite correction**: In this case the loaded plasma activity concentration already equals the input curve.
- **Use of parent fraction**: In this case the plasma activity concentration (which should equal total activity in plasma) is multiplied with parent fraction for calculating the input curve.
- **Use of plasma fraction**: In this case the whole blood activity is multiplied with the plasma fraction for calculating the plasma activity concentration. If the plasma fraction includes the metabolite correction part, the plasma concentration already represents the input curve. Otherwise, a parent fraction needs to be loaded, and the input curve results from multiplying the whole blood curve with the plasma fraction and then the parent fraction.

In practice, PKIN behaves as follows:

- As soon as a whole blood activity concentration curve is loaded and no other information is available, it is also used as the input curve. To this end the plasma mode is set to "fraction", and a Fix plasma fraction model is applied with a factor of 1.
- As soon as a plasma activity concentration curve is loaded, any plasma fraction data is discarded, and the plasma mode set to "activity". The activity interpolation defaults to Lin. Interpolation. The setting of the parent fraction is not changed.
- As soon as a plasma fraction curve is loaded, any plasma activity concentration data is discarded, and the plasma mode set to "fraction". The fraction interpolation defaults to Lin. Interpolation. The setting of the parent fraction is not changed.
- A parent fraction equal to 1 is applied as long as no parent fraction curve is loaded, or no analytical parent fraction model selected. Correspondingly, the input curve corresponds to the plasma activity concentration.

**Organization of the Blood-related User Interface**

The different blood-related information parts are readily available for configuration via the radio buttons **Whole blood, Plasma and Metabolites** on the **Blood** tab as illustrated below.

4.5.1 **Interpolation of Blood Data**

The tracer activity in blood is often only sampled at a few time points during the acquisition. However, when calculating the operational equations (eg. by numerically integrating a system of
differential equations), the blood activities must be available at any arbitrary time point during the acquisition period. This means that the blood curves must be interpolated according to some underlying model function. The interpolation models available for the different blood curves are described in the sections below, and the model fitting always works in the same way as explained for the tissue models.

4.5.2 Whole Blood

It is assumed that the time-course of the tracer activity in whole blood has been loaded with Kinetic/Load Whole Blood Activity. To configure the interpolation model of whole blood please select the Blood tab, and set the Whole blood radio button. A list of model functions is available which can be shown with the arrow button indicated below.

Default is Lin. Interpolation, which just represents linear interpolation between sample times. To replace linear interpolation by a smoother function select an appropriate definition from the Whole Blood model list. The model functions are described in the reference section. As soon as a model function is selected, the parameters are updated in the Standard pane, and a corresponding model curve is shown in the curve window as Whole blood model.

In the example above the model has not yet been fitted so that the distinction between the measurement and the model (sum of exponentials) is clearly visible. The parameters of the blood model can be configured for fitting purposes by enabling the check boxes. Activating the Fit whole blood button starts a fitting process which adjusts the model parameters such that the interpolation curve comes into optimal agreement with the measurements. Fitting works exactly in the same way as is explained for the tissue models here.

Replace activity data will replace the actual data samples by the current model values at the original sampling times. This functionality is mostly useful for replacing a blood curve by its dispersion-correction as described in a separate section.

The Blood Delay Parameter

All standard blood models have a Delay parameter to correct for a timing error between tissue and blood data. Positive delays represent delayed blood information and hence shift the blood curves to earlier times (to the left). This parameter is only relevant for fitting of the tissue model.
Therefore, when fitting the shape of the blood curve with Fit whole blood, it is automatically disabled. When fitting the kinetic tissue model, however, it can be allowed for fitting.

4.5.3 Plasma

Plasma Activity
Here it is assumed that the time-course of the tracer activity in the arterial plasma has been loaded with Kinetic/Load Plasma Activity. The configuration and fitting of the interpolation model works in exactly the same way as for whole blood.

Please select the Blood tab, and set the Plasma radio button. Note the label Plasma activity label which indicates that the current working mode uses measured activities, not a derivation from whole blood activity. The same list of models as for the plasma activity of whole blood is available. They are described in the reference section.

To the right of the Fit plasma activity button, there is an option button for changing the current data.

- **Replace data**: This function replaces the data samples by the values of the current model function at the original sampling times.
- **Resample input curve**: This function creates an explicit input curve from the current blood definitions. It resamples the input curve at the sampling times of the plasma curve and replaces the plasma measurements. Additionally, it sets the parent fraction to one, as any metabolite fraction is now included in the input curve. This approach may have advantages, if the plasma activity and/or the parent fraction can't be reasonably fitted, but their combination has a fitable shape.
- **Clear activity data**: This function discards the loaded plasma data and switches the mode to "plasma fraction". In that case the assumption is that an analytical plasma fraction function will be applied to the whole blood activity curve to derive the plasma activity.

Plasma/Whole blood Fraction
Here it is assumed that the time-course of the plasma fraction has been loaded with Kinetic/Load Plasma/WB Ratio. To configure the interpolation model please select the Blood tab, and set the Plasma radio button.

Note the label Plasma fraction which indicates that the current working mode uses a function to derive plasma activity from whole blood activity. There is a list of models which can be fitted to the plasma fraction data. As soon as a model is selected, its parameters are shown in the Standard...
pane, and a corresponding model curve is displayed in the curve window as Plasma fraction model. The available models are described in the corresponding reference section.

To the right of the Fit plasma fraction button, there is an option button for changing the current data.

- **Replace data**: This function replaces the data samples by the values of the current model function at the original sampling times.
- **Resample input curve**: This function creates an explicit input curve from the current blood definitions. It resamples the input curve at the sampling times of the plasma curve and replaces the plasma measurements. It then switches back to "plasma activity mode" and sets the parent fraction to one, as any metabolite fraction is now included in the input curve. This approach may have advantages, if the plasma fraction and/or the parent fraction can't be reasonably fitted, but the input curve has a fitable shape.
- **Clear fraction data**: This function discards the loaded plasma fraction data. In that case the assumption is that an entirely analytical plasma fraction function will be applied to the whole blood activity curve in order to derive the plasma activity.

### 4.5.4 Parent/Plasma Fraction

The transformation of the plasma activity curve into the input curve is governed by the configuration on the Metabolites tab. Two cases are to be distinguished.

**Measured Parent Fraction Data is available**

If a parent fraction curve has been loaded, it is shown in the curve area. Initially, the parent fraction model is set to Lin. Interpolation so that linear interpolation between the samples is used to calculated intermediate parent fractions. The model selection choice offers smoother models, and they can be fitted to the measurements using the Fit plasma fraction button as illustrated below.
To the right of the Fit fraction button, there is an option button for changing the current data.

- **Replace fraction data:** This function replaces the data samples by the values of the current model function at the original sampling times.
- **Clear fraction data:** This function discards the loaded parent fraction data. In that case the assumption is that an entirely analytical parent fraction function will be applied to the plasma activity curve in order to derive the parent activity.

**No Parent Fraction Data available**

If no data parent fraction data have been loaded, the parent fraction model is set to **Fix** with a free parent fraction $f_p = 1$. This is adequate for all tracers which do not require metabolite correction, and for cases when the metabolite correction has already been applied to the plasma data prior to loading them with **Load Plasma Activity**.

In the case of a tracer which shows a characteristic buildup of the metabolites it may be possible to derive an average parent fraction curve and fit a model to it. To apply a metabolite correction the user can then simply select the appropriate model function and enter the demographic parameter values, as illustrated below. To apply a demographic correction, it must be ensured that time zero of plasma data always represents the time of tracer injection. Naturally, no fit will be possible in this situation.
The available models are described in the corresponding reference section.

**Note:** The metabolite correction is applied to the plasma curve using the original plasma sampling times. The assumption is that the parent fraction and the plasma curve have a common time scale. Only after the metabolite correction has been applied is the input curve shifted by the blood sampling time delay.

### 4.5.5 Default Model Configurations and Configuration Sets

All interpolation models allow the customization of the model default parameters, and the saving of specific configuration sets for certain tasks. The mechanism is the same as described for the tissue models.

### 4.6 Configuration and Fitting of Kinetic Tissue Models

As soon as a kinetic model has been selected, a model curve is calculated and shown in the curve display. The model curve represents the activity concentration values PET would measure if the model correctly described the observed physical processes. Fitting assumes that the model configuration is adequate and tries to find the set of model parameters for which the model curve is in optimal agreement with the measurement.

The following sections explain how a model is configured, fitted to data, and assessed for goodness-of-fit.

#### 4.6.1 Kinetic Model Configuration

The main functionality for the definition and fitting of kinetic tissue models is available on the Tissue pane. The upper part contains data and model selections, while the lower part gives access to the model details and fitting results.

![Tissue pane screenshot](image)

The main elements for the model configuration on the Tissue pane are:

<table>
<thead>
<tr>
<th><strong>Region</strong></th>
<th>When several regional TACs have been defined and imported, the current regional TAC can be set using this list selection. The left and right arrows allow stepping forwards and backwards through the TACs. It is possible to scroll the selected region with the mouse wheel if the cursor hovers on top of the list.</th>
</tr>
</thead>
</table>
After region switching, the display is updated with the configured model and the curves of the selected region.

**Reference**

Selection of the regional TAC serving as reference in models which have the input curve replaced by a tissue TAC. Note that *Edit Data* allows creating a TAC averaged from several loaded TACs, which may be useful as a reference.

**Model**

The PKIN tool includes a comprehensive list of standard kinetic models and some auxiliary calculations. This list allows selecting among all currently configured models. As soon as the model is changed, the pane in the lower section is updated to show the parameters of this model.

Note that the models in the list can be rearranged, added or deleted in the *Global Configuration* utility. Furthermore, the list can be focused on the current analysis by filtering.

**Button for showing a quick information about the current model.**

---

### 4.6.1 Reference Models

When a reference *Model* is selected which uses the activity in a reference region as an implicit input curve, the blood-related items become disabled and the *Reference* region selection becomes active.

PKIN will pre-select a region containing "Cerebellum" or "Reference" in the name as the reference. If the selection is not adequate, please select the proper region from the *Reference* list.

---

### 4.6.2 Model Parameters

The *Standard* sub-tab illustrated above shows the configuration of the currently selected model. It encompasses different types of parameters:

- Input parameters which are needed to specify information used for model calculations.
- Fitable parameters which require a starting value and show the updated results after model fitting. A parameter is only optimized if its fit box is checked, otherwise it remains fixed.
- Macroparameters are calculated from model parameters in the upper part to instantly provide composite information such as the distribution volume or the flux.
The fitable and derived parameters have four columns of information:

- The **Parameter** name. In order to see a brief explanation of the parameter place the mouse pointer over the name. The explanation text is then displayed in the status line at the bottom.
- The **Current value**, which can be edited to set an initial value, and which is updated as the result of a fit.
- The **Unit** of the parameter value.
- The standard error **%SE** of the parameter expressed as a percentage of the parameter value itself. It is only available, if a fit has been performed using the Marquardt-Levenberg method.

### 4.6.1.3 Default Model Configurations and Configuration Sets

The parameters of the current model can be saved or retrieved using the buttons illustrated below.

**Model Default Configuration**

Each model has a set of default parameter values. Initially, they are factory settings, but they can be re-defined by the user if he wants to establish a default configuration which is more adequate from him.

<table>
<thead>
<tr>
<th>Settings as default</th>
<th>Save the current parameter configuration as the new default for the particular model type. Included are the values, the fit flags, and the restrictions of all parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Default Settings</td>
<td>Resets the configuration of the current model to the saved defaults.</td>
</tr>
</tbody>
</table>

**Model Configuration File**

A model configuration can also be saved as an explicit definition file (.kmModel). It includes the tissue model as well as all blood models used. Included are the initial parameters, the fit flags, the restrictions and the residual weighting scheme (except for the prescribed weighting with the explicit weighting factors).

| Settings to file | Save the current model and parameter configuration in a file.                                                                                                                                  |
Default from file | Retrieve a model configuration. Note, if the model in the file is different from the current model, the model is switched accordingly.

Configuration files make it simple to establish defaults for particular data processing tasks: just load the appropriate .kmModel file and set it as the default. Another application is batch processing. There, multiple models can be fitted to the data by simply selecting the corresponding .kmModel files.

### 4.6.1.4 Visualization of Parameter Effect

The **Increment** sub-tab is mainly used for visualization purposes. For each fitable parameter the current value is shown next to an increment value and an increment slider. When the arrow buttons are selected, the parameter value is incremented or decremented accordingly. The slider allows to increment/decrement a parameter smoothly. The minimal and maximal values for the slider as well as the initial increment are obtained from the values on the **Restrictions** tab.

After every parameter change the curve display is updated, so that the user can observe its effect on the global uptake (model curve) and on the uptake in the different compartments (C(t), C(t)).

For instance, as shown below, it can be nicely demonstrated that a change in K1 simply scales the model curve, but does not change the curve shape: Set the curve display to relative scaling by selecting the % in the upper right and then change K1. Nothing changes in the % display, provided that the blood volume fraction is set to 0. Any modification of the other parameters changes the curve shapes notably.

### 4.6.1.5 Parameter Restriction

The **Restrictions** sub-tab is relevant for fitting. For each fitable parameter there is a box for enabling the restriction which is defined by the **Lower bound** and the **Upper bound** value. The purpose of parameter restrictions is to ensure that the resulting parameters remain within a reasonable physiologic range. Without restrictions, there is a higher chance to end up with a meaningless solution. Note that these bounds are also used for the sliders on the **Increment** pane.

The **Search** value is used as an initial step-size for iterative fitting. It is recommended to set this value to about the same size as the parameter value itself, as it is automatically decreased if needed. This approach is not successful if the value is very small from the beginning.
4.6.1.6 Model Parameter Propagation/Conversion during Model Switching

The behavior of PKIN when switching between models is governed by the Parameter conversion check in the Extras pane, and the corresponding Conv option in the taskbar.

Parameter Conversion Disabled

The first time a model is selected for a particular region, the model default values are used. Thereafter, the most recent model parameter set as used for the same region is applied.

Parameter Conversion Enabled

Parameter conversion can be used to propagate parameter values from one model to another which includes the same parameter, or to convert compartment parameter values due to a change of model complexity. 

- **Parameter propagation**: Parameter propagation allows to transfer the exact value of certain parameters from one model to another. This feature can for instance be used to fit a parameter using a robust method and propagate it to a more complex model where it is fixed during fitting. Examples of important propagated parameters are \( v_B \), \( t^* \), Flux, \( k_{2} \).

- **Parameter conversion**: When switching between models with the same number of compartments, the conversion is exact. If the number of compartments is decreased, the lumped parameters of the simplified model are calculated. If the number of compartments is increased, the same values are used for corresponding parameters, and the additional parameters set to the default values. The consequence of this functionality is that when switching the model from 1-tissue to 2-tissue and then back, the \( K_1 \) and \( k_2 \) values will be different due to this conversion process.

4.6.1.7 Model Filtering

The basic model configuration should include all models which may be relevant for a user. Consequently, the model list in the PKIN user interface can still be quite long. Model filtering can be used to quickly tailor it to the current task.

Model Filter Dialog Window
The filter configuration window is opened with the 🎉 button in the taskbar.

The Model filter criteria area has five classifiers which can be enabled:

<table>
<thead>
<tr>
<th>Method</th>
<th>Selections: Classical PET, Compartment Model, Graphical Plot, Spectral, Multilinear, Utility.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Domain</td>
<td>Selections: Tissue Perfusion, Receptor Binding, Heart Perfusion.</td>
</tr>
<tr>
<td>Result</td>
<td>Selections: Vt, BPnd, Flux, Bmax.</td>
</tr>
</tbody>
</table>

After selecting one or more filter classifiers the model list in the window is reduced accordingly.

The Adjust models to filter flag implements a prior condition on the selected models. Currently, only the Trapping has an impact. It causes setting $k_4=0$ in the compartment models. Show only selected models allows reducing the models further to the ones currently selected in the list.
Filter Activation

When closing the dialog window with Filter, the entries in the Model list are reduced to the entries (selected or non-selected) in the Model List Filter dialog. Note that in the example below the filter shortcuts in the taskbar are greyed out, as they don’t correspond to the current selection. The model list can be reset in the Model List Filter dialog or by the shortcut button in the taskbar.

Filter Shortcuts in Taskbar

Note the set of radio buttons in the taskbar to the right for changing the model filter settings quickly.
While All shows the unfiltered list, Bld, NoB, Ref and Irr restrict the list to blood-based, all but blood-based, reference tissue and irreversible models, respectively.

### 4.6.2 Kinetic Model Fitting

The main elements on the Tissue pane for model fitting are highlighted below. The left button applies fitting to the current region, whereas the right button applies fitting in all regions.

<table>
<thead>
<tr>
<th>Current Region</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Fit model, current region</strong></td>
<td>Starts a process which fits the selected Model to the measurements of the selected Region. For compartment models the fit requires an iterative optimization, while for some other models only a single calculation may be sufficient. Note: in the case of a reference model the reference region itself is not possible to fit.</td>
</tr>
<tr>
<td><strong>Macro fit, current region</strong></td>
<td>Starts a process which fits the models in the selected Macro to the measurements of the selected Region. The fitting results are stored in the model history, and the fit of the last model is shown.</td>
</tr>
<tr>
<td><strong>Fit model w. blood delay</strong></td>
<td>With this option, the model parameters are fitted simultaneously with the delay of the model curves, see Blood Delay Fitting.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>All Regions</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Fit model, all regions</strong></td>
<td>Starts a process which fits the configured Model in each Region to the measurements. Note that the model may differ among regions.</td>
</tr>
<tr>
<td><strong>Macro fit, all regions</strong></td>
<td>Starts a process which sequentially fits the models in the selected Macro to the measurements of all Regions.</td>
</tr>
</tbody>
</table>

**Model Propagation and Fitting**

When prototyping the data analysis the approach described in the Processing Overview is recommended:
1. Select a representative tissue region; it may also be helpful to first average a few curves using the Edit Data facility.

2. Select the model to be tested.

3. Fit the data.

4. Propagate the model together with all related configurations (fit box state, restrictions) to all regions.

5. Fit the model in all regions.

6. Open the Parameter Explorer and check the regional summary on the Last Revision pane.

4.6.2.1 Iterative Fitting Principle

For most models, minimization of the Chi square expression requires an iterative approach. The procedure starts with a set of initial parameter values, generates the corresponding model curve, calculates the Chi square criterion, and adjusts the model parameters such that Chi square is expected to be reduced. The cycle

- chi square calculation
- parameter adjustment

is then repeated until a further reduction of Chi square is not any more possible. The final set of parameters upon termination is regarded as the best-fit result. It is not analytically exact and may depend on the initial parameters, on the optimization procedure, and on the termination criterion of the iterations.
### 4.6.2.2 Fitting Options on Extras Panel

The **Extras** pane contains some options which have an impact on model fitting.

<table>
<thead>
<tr>
<th>Option</th>
<th>Description</th>
</tr>
</thead>
</table>
| Linearized initialization | This configuration is only related to the compartment models and determines which initial values are used for the iterative fitting:  
  - Disabled: the parameters in the user interface are used.  
  - Enabled: an automatic procedure determines initial parameters which should be reasonably close to the solution. This is done by the linear least squares method. Note that the initial values are not shown to the user. However, to see the values, please switch to the linear least squares method and fit. |
| Parameter conversion    | Behavior of the parameters when switching between models.                                                                                                                                                       |
| Float blood parameters   | If this box is checked on, all parameters of the Input curves which have the fit box checked are also included into the set of fitted parameters as described separately. This feature has two uses:  
  - Fitting the relative time delay between the tissue and the blood data simultaneously with the kinetic rate constants. For this use, there is a dedicated fitting shortcut.  
  - (Rarely) Fitting a shape parameter such as the half-time of a metabolite build-up together with the rate constants. Note: When fitting all regions, the blood parameters will not be fitted, even if Float blood parameters is enabled. |
| Single fitting           | Default mode of performing one single fit.                                                                                                                                                                     |
| Random fitting           | This configuration is only related to models which require iterative fitting:  
  - Multiple fits are performed, and the result parameters of the fit with minimal chi squared returned. The fits are started with randomized sets of initial parameters using a uniform distribution with ±100% range. Note: If Parameters initialization is enabled, randomization uses the automatically determined parameters for the random number generation rather than the parameters visible in the user interface.  
  - In case randomized fitting is enabled, the labels of the fitting buttons are adjusted correspondingly. Note: The purpose of randomized fitting is to avoid local minima, but there is still a chance that the returned solution may contain unphysiological parameter combinations. |
| Grid fitting             | This configuration is only related to models which require iterative fitting:                                                                                                                                   |
Multiple fits are performed, and the result parameters of the fit with minimal chi squared returned. Initial parameter sets are generated by stepping each parameter in increments between a lower value and an upper value. The number (n) specified equals the number of distinct values used, so the number of increments is n-1. The interval of the parameter range to be covered is defined on the Restrictions panel. It is necessary to adjust these ranges to physiologically reasonable values, because the default ranges are very wide. In case grid fitting is enabled, the labels of the fitting buttons are adjusted correspondingly.

Multiple fits are performed, and the result parameters of the fit with minimal chi squared returned. Initial parameter sets are generated by stepping each parameter in increments between a lower value and an upper value. The number (n) specified equals the number of distinct values used, so the number of increments is n-1. The interval of the parameter range to be covered is defined on the Restrictions panel. It is necessary to adjust these ranges to physiologically reasonable values, because the default ranges are very wide. In case grid fitting is enabled, the labels of the fitting buttons are adjusted correspondingly.

Data shortening to

Enables a sequence of fits with a successively shorter data segment to study the parameter dependence on acquisition duration.

Fitting method

Selection of the numerical optimization method applied for iterative fitting:

- **Marquardt**: Marquardt-Levenberg algorithm [59]. Usually works faster and has the additional advantage that it accumulates information about the covariance matrix. Therefore, estimates of the standard error can be calculated for the fitted parameters. They are shown as \%SE, i.e. as a percentage of the parameter value.
- **Powell**: Conjugate direction set method [59].

The iterative optimizations terminate, when no substantial improvement can be found any more, or when the Max iterations restriction is hit.

Note: Several models are not fitted iteratively because they have closed form solutions or use a basis function approach. In these cases the Fitting method settings are not relevant.

Max. iterations

Maximal number of iterations which are allowed in the iterative Marquardt and Powell fitting.

**Taskbar Shortcuts**

Note the set of radio buttons in the taskbar to the right for enabling/disabling some of the options quickly.

The Init and Conv boxes correspond to Linearized initialization and Parameter conversion, respectively. The Sng, Rnd and Grd radio buttons switch between Single, Random or Grid fitting.

**4.6.2.3 Residual Weighting**

Weighting is used to increase or decrease the influence of certain data samples on the fitting. For instance, initial samples with short acquisition duration and little uptake may be considered less reliable than samples with longer durations in the presence of high uptake. The weights are multiplied with the squared residuals when calculating the Chi squares criterion.

In PKIN, several different weight definitions are supported. They can be specified on the Weighting pane using the list selection with the three entries Constant weighting, Prescribed weighting and Calculated weighting.

**Constant Weighting**
Note that the weights are calculated from the definition of a standard deviation \( \sigma_i \) or the variance \( \sigma_i^2 \) by

\[
w_i = \frac{1}{\sigma_i^2}
\]

With constant weighting the same weight is applied to all residuals. Its size can be adjusted by the Scale factor and the radio button setting. In the case of TAC mean \( \sigma_i \) is obtained as the average of all values in the TAC multiplied by the Scale factor. Similarly, with TAC max \( \sigma_i \) is obtained as the maximal TAC value multiplied by the Scale value. With All weights = 1 the weights are all set equal to 1 independent of the Scale \( (\sigma_i = \sigma_i^2 = w_i = 1) \).

Note that the scale factor doesn’t have an impact on the fit result. It is only relevant when the weighting definition is used for specifying the noise generated in Monte Carlo simulations.

**Prescribed Weighting**

The default behavior of this method is to use the standard deviation of the VOI pixel values for calculating the TAC for weighting. This information is automatically available if the TACs have been transferred from the PMOD viewing tool to PKIN, and can be visualized using the button Show measured stdv as illustrated below.

This definition can be overwitten by user-defined weights, see below.

**Loading Externally Defined Weights**
Some users might want to implement their own residual weighting schemes. To this end they should calculate the standard deviations $\sigma_i$ for the different acquisition frames and arrange them in a tab-delimited text file of the following form:

<table>
<thead>
<tr>
<th>t[seconds]</th>
<th>standard-dev[kBq/cc]</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>0.10300076</td>
</tr>
<tr>
<td>30</td>
<td>0.01717448</td>
</tr>
<tr>
<td>50</td>
<td>0.01195735</td>
</tr>
<tr>
<td>70</td>
<td>0.01010257</td>
</tr>
<tr>
<td>etc</td>
<td>...</td>
</tr>
</tbody>
</table>

The header line is required. The first column represents the frame mid-times (although the values are not interpreted), and the second column the standard deviations in appropriate units. The number of entries must be equal to the number of acquisitions, and the columns separated by spaces or tabs. For example, such a file can be prepared in MS Excel and then saved as a text file with tab delineation.

This definition can be loaded by the **Load stdv from file** button available with **Prescribed weighting**.

**Calculated Weighting**

With this selection the variance is calculated for each TAC value from an equation which may take into account the measured uptake, the radioactive decay, and the duration of the acquisition. The following equation is used:

$$\sigma_i^2 = \alpha^2 \frac{C_{\text{PET}}(t_i)}{\Delta t_i} e^{-\lambda t_i}$$

It includes a **Scale** factor $\alpha$, the decay-corrected **Uptake** $C_{\text{PET}}(t_i)$ at frame mid-time $t_i$, the frame **Duration** $\Delta t_i$, and the **Decay** constant $\lambda$ which is obtained from the half-life $T_{1/2}$ of the isotope. If the half-life has not yet been defined, it can be set on the **SUV** panel with the **Edit Patient** facility.

The settings of the radio buttons have the following effect on the equation above for calculating $\sigma_i^2$:

**Uptake**:
- **Not considered**: $C_{\text{PET}}(t_i)$ is replaced by 1.
- **Measured value**: $C_{\text{PET}}(t_i)$ is used, corresponding to **Poisson weighting**. The effect is that more emphasis is given on low-uptake values than on high-uptake values. This may be helpful for example if the fit seems to not account enough for the tail of a decreasing function such as the input curve.
**Measured value squared**: $C_{\text{PET}}^2(t_i)$ is replaced by $C_{\text{PET}}^2(t_i)$, corresponding to *Relative weighting*. With this option the influence of the uptake is further increased. It should be used with care because by squaring small (initial) uptake values ($<<1$) very small variances are generated which transform into heavy weighting. As a consequence, a few small uptake values may have excessive impact on the fit.

**Duration:**
- **Not considered**: $\Delta t_i$ is replaced by 1.
- **Measured value**: The individual frame durations $\Delta t_i$ are used, so that longer acquisitions are considered more reliable.

**Decay:**
- **Not considered**: $e^{-\lambda t_i}$ is replaced by 1.
- **Frame mid-time**: $e^{-\lambda t_i}$ is used, so that earlier acquisitions are considered more reliable.

**Note**: As a corrective measure, very small standard deviations of the first 1/3 data points (likely to have low uptake) are adapted to provide reasonable weights as follows: If the stdv is smaller than the average stdv and smaller than 50% of its successor, it is replaced by the stdv of the successor.

**Weights Visualization**
Whenever the weighting configuration is changed, the standard deviation $\sigma_i$ is calculated and plotted as an error bar around the measurements. This gives the user a visual feedback as a help in judging the adequacy of the specified weighting.

**Display of the Residuals**
The curve area with the residuals per default shows the weighted residuals, that is the difference between the measurement and the model curve divided by the standard deviation.

$$t_i = \frac{C_{\text{PET}}(t_i) - C_{\text{Model}}(t_i)}{\sigma_i}$$

However, the raw residuals are also available. The context menu as illustrated below (right mouse click into the curve area) can be employed for showing/hiding the curve controls, which allow changing the displayed information.
Use of the Weights in Monte Carlo Simulations

Monte Carlo simulations require the generation of noise which is defined by a distribution type as well as its deviation characteristics. Assuming that the weights specification corresponds to the standard deviation of the measurement noise, the regional weighting specification is used in the Monte Carlo noise generation together with the distribution type. Note that the Scale factor has no impact on fitting, but that it is highly relevant in Monte Carlo simulations.

4.6.2.4 Model Fitting History

The system maintains a history of model configurations for each region. All operations, which change the model configuration result in a new entry in the history list. The history is saved within a .km file and will be continued after reloading the data.

The history functionality is accessible by the user interface elements right to the model configuration:

- Switches to the previous model configuration in the history list.
- Appends the current model configuration to the list.
- Switches to the following model configuration in the list.
- Shows the model history window.

**CAUTION:** This history mechanism does not account for changes in the configuration of the blood-related models, except for the blood delay. However, it includes the definition of the residual weighting, the valid points, and the tissue TAC values.

Model History Window

The model history list is shown in the Regional Fitting History Explorer window as illustrated below. It has the same organization as the window for comparing all regional results, except that only the information of the current region is available.
The upper part of the **History** panel lists the model configurations in rows. The initial columns are descriptive, followed by the actual parameters. Note that the parameters are sorted into appropriate columns even across different model types. For instance, $V_t$ resulting from all compartment models as well as the Logan plot appears in the same column so that the values can easily be compared. The user can change the order of the columns by dragging the column headers left or right. The smallest value per column is encoded in green, the largest in red. Columns having a constant value are encoded in blue. The **No** column refers to the chronology of list entry. **Revision** serves for grouping results from a certain task such as the parameter sensitivity analysis. **Type** indicates the process which added the configuration entries. The list can be flexibly sorted by clicking on the column headers.

To restore the situation of a model configuration in the list just double-click the entry or mark it **Select**. Model configurations can be removed by selecting them in the list and activating **Delete**. The **Clear Region** button removes all list entries, and **Clear all Regions** empties the histories of all regions.

**Optional Elements**

The window has some configuration **Settings** which can be opened as illustrated above. The elements to be changed are:

<table>
<thead>
<tr>
<th>Optional Element</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Include standard error (%SE) in table</td>
<td>Include a column with the % standard error for each parameter.</td>
</tr>
<tr>
<td>Include distribution statistics at table bottom</td>
<td>Enable the lower area showing the min, average, median, max and stdv per column.</td>
</tr>
<tr>
<td>Use last region fits for table</td>
<td>Not relevant for the fitting history of a single region.</td>
</tr>
<tr>
<td>Display tab with parameter plots</td>
<td>Shows an additional panel for the visualizing the parameters of the selected History rows as curves.</td>
</tr>
<tr>
<td>Display tab with relative parameter plots</td>
<td>As above, but with normalization to the last value.</td>
</tr>
</tbody>
</table>
Display tab with model curves

Shows an additional panel for concurrently visualizing the model curves of the rows selected on the History tab.

Region filter
Not applicable here.

Model filter
For filtering the entries listed. With "Comp", for instance, only compartment model results will be listed.

Saving Options
There are different types for saving the parameters from the fitting history using selection in the lower left.

Copy to Clipboard copies the entire window contents, so that it can be pasted into another application for further analysis. Save Current State saves the current parameters from all regions into a .kinPar file. Save Selected Revision saves the fits from all regions with the currently selected Revision number to a .kinPar file and can be used for several purposes like exporting the results from a macro fit or a duration sensitivity analysis.

4.6.2.5 Fitting Problems

In the case of a failed fit it is worthwhile to check the console output. For example, the output text reproduced below clearly indicates that the 4th fit parameter caused a problem.

Singular Matrix-2

covar[1][1] = 2.1652603147908747E-6
covar[1][2] = -5.120936302955895E-4
covar[1][3] = 2.862241687078937E-7
covar[1][4] = 0.0
covar[2][1] = -5.120936302955895E-4
covar[2][2] = 0.3918087002588389
covar[2][3] = -2.0406232308544742E-4
covar[2][4] = 0.0
covar[3][1] = 2.862241687078937E-7
covar[3][2] = -2.0406232308544742E-4
covar[3][3] = 1.4044964781403112E-7
covar[3][4] = 0.0
covar[4][1] = 0.0
Such problems might be caused by initial values very close to parameter restrictions.

### 4.6.3 Macro Fitting

Macros are a convenience for fitting multiple models to the data with a single click and exporting the results in a single operation.

#### 4.6.3.1 Macro Creation

The **New** button in the **Macro** line opens the editor illustrated below for creating new macros.

The available models are listed to the left. This list can be reduced by the filtering options to the right explained in the [Model Filtering](#) section. The list can be reordered by selecting an entry and shifting it with the arrow buttons indicated above.

**Models in the Macro**

Once the model order is proper, select the models to form the macro in the list. When activating **Show only selected models** the list is condensed and the macro content can be inspected. Note that the models are fitted sequentially in list order, and the results are stored chronologically in the model history. Consequently, at the end of macro execution, the result of the last model will be shown.

**Fitting Options**

Macros store the **fitting options** in the lower section. During macro execution they will be applied for all models as applicable. Valuable uses of **Parameter conversion** include propagation of a common $vB$, $k_2^*$ or $t^*$ as detailed [here](#).

**Macro Saving**

Once the models are in correct order and the fitting options specified, the macro can be saved in two ways:
1. **Save as default:** This will save the macro in the `resources/pkin/macros` directory

![Macro Editor screenshot](Image)

and as a result the macro will immediately appear in the **Macro** list

![Confirmation dialog](Image)

Note: For cleanup of the list please delete unnecessary `.kmMacro` files from the `resources/pkin/macros` directory in the PMOD installation.

2. **Save to DB:** This allows saving the macro in a PMOD database, or via **Save to File System** anywhere on a disk.

### 4.6.3.2 Macro Execution

Existing macros can be executed to fit the measurements of the current region or all regions. First select the appropriate **Macro** from the list (**Vt calculation** in the example below). To start the macro switch the fitting button to **Macro fit, current region**, or **Macro fit, all regions** as illustrated below.

![Macro Execution screenshot](Image)

Note that if the **Dialog** box is checked, the macro contents will be shown in a dialog window as below
This allows adjusting the preset fitting options and specifying a .kinPar file to which the parameters resulting from the fits will be written.

**External Macro**

Macros which are not stored in the default location can be loaded with the **Load external macro** entry. In the appearing dialog window select the macro from a database, or use **Load from File System** to browse for a .kmMacro file on the disk. The selected macro will be listed below the **Load external macro** entry.

Note that only one external macro is supported. If a different macro is selected, the previously listed external macro will be replaced.

**Initial Parameters**

The initial parameters of the models applied in the macro are retrieved from the corresponding **Default Configurations** of the model. This means, if specific initial parameters are to be applied, these model configurations should be stored beforehand and retrieved before macro execution with **Settings from file** as illustrated below.

An alternative is to prepare dedicated users in the global PMOD configuration for different types of data analysis tasks, because the set of model default configurations is serialized per user between PMOD sessions.

### 4.6.3.3 Macro Results Inspection

The results of the sequential fitting of all macro models are recorded in the **Model Fitting History**. They are labeled by the same **Revision** number, and the name of the macro is indicated in the **Type** column.
Single Region Macro Results
The regional macro results can be inspected using the Model Fitting History as illustrated below. All fits of a macro have the same Revision number and include the macro name in the Type field.

All Regions Macro Results
If the macro has been applied to all regions, the results are best inspected and exported using the Parameters Explorer. If it is opened directly after macro execution, the pane Last revision assembles the result parameters of all models in the macro across all regions. These values can then be exported conveniently via Copy to Clipboard.

4.7 Coupled Fitting for Improving Parameter Estimates
In some situations the introduction of prior (physiologic) knowledge can help improving the stability of model fitting.

For instance, in receptor experiments it may be adequate to assume that the distribution volume \((K_1/k_2)\) of the non-displaceable compartment and/or the dissociation rate from specific binding sites \(k_4\) are equal in certain tissues of the brain. Hence, a simultaneous fit of selected regional TACs of a scan can be performed which finds separate \(K_1\) and \(k_3\) values per region, but which delivers a \(K_1/k_2\) and \(k_4\) which is common for all TACs. As an example, Sanabria-Bohorquez et al [1] have used a coupled \(k_4\) for solving the problem of unstable fits. In PKIN this type of fitting is called Coupled Fitting.

Another valuable application of Coupled Fitting is determination of \(k_2'\) for Reference Tissue models. \(k_2'\) denotes clearance of the tracer from the reference tissue. Hence, it is natural to enforce a common \(k_2'\) as physiologic restriction when fitting the TACs from different target tissues as explained below.

The concept can be extended to the fitting of data from different scans of the same subject, for instance test/retest scans which are replicate measurements under the same conditions and where the parameters should be very similar if the regions are consistently outlined. Plisson et al. [2] have applied simultaneous fitting with the data of pigs which were studied in a baseline...
condition and with different degrees of receptor blocking by cold compound. By applying the simultaneous fitting, they were able to get stable estimates, which was not possible with independent fitting. In PKIN this type of fitting is called **Coupled Studies Fitting** and distinguishes two variants of coupling. **GLOBAL** coupling of a parameter results in single value which is common for all fitted TACs. **REGIONAL** coupling of a parameter results in common value per region. For instance, if $k_3$ is defined as a parameter with **REGIONAL** coupling and the coupled regions from two scans are caudate, putamen and frontal, a different $k_3$ estimate will result for each of these three regions, but each of them is common across the TACs of the two scans.

The requirements for coupled fitting in PKIN are:

- The same model applies to the TACs of several tissue regions (Note: not all models may be coupled, only models with iterative fitting).
- There exists one or more parameters in the model configuration which can be assumed to have a common value for different TACs.

**References**


### 4.7.1 Common Parameters across Regional TACs of a Single Study

**Coupled Fitting to Estimate and Apply a Common $k_2'$**

The following sequence of steps is recommended for coupled fitting. The example illustrates fitting of a common $k_2'$ using the **SRTM2** model. It is equivalent to the **SRTM** model, but due to a different arrangement of the equations $k_2'$ appears as a fitting parameter which allows using coupled fitting.

1. Select a regional TAC with good signal.
2. Switch to the model (**SRTM2**) to be applied in coupled fitting.
3. Fit the TAC with **Fit model, current region** and propagate the model to all other regions with the button **Model & Par**. Use **Fit model, all regions** to establish model and reasonable starting
values for all TACs.

4. On the **Coupled** panel, enable the parameter(s) which are to be treated as common, and enable all target regions which should be fitted together. Note that the non-coupled parameters will be estimated per region.

5. Start processing with **Fit coupled**. Coupled fitting may take significant processing time. Thus the **Batch Mode** facility may be a helpful option to run such jobs at less busy times.
6. On the **Tissue** panel, switch to one of the coupled regions. The $k_2'$ shown there is the common estimate. This can be verified by opening the parameter explorer as below.

![Parameter Explorer](image1)

7. To use the common parameters for the fitting in all regions select one of the coupled regions. The **Standard** panel shows the result of the coupled fit including the estimated values of the common parameters. Note their standard error %SE which should be decreased compared with individual fits. Disable the fit box of the common parameter(s) and propagate this configuration to all regions with the **Model & Par** button. Apply **Fit model, all regions** to re-fit the model with the fixed common $k_2'$ parameter which is now enforced in all regions.

![Model and Parameter](image2)

**Coupled Fitting to Improve Robustness of 2-Tissue Compartment Model (2TCM) Fitting**

It is often difficult to get robust estimates for $k_3$ and $k_4$ with 2TCM fitting. In order to support parameter coupling, an alternative formulation of the 2TCM is supported which uses $K_1/k_2$ (distribution volume of non-displaceable compartment) as fitting parameter instead of $k_2$.

To perform 2TCM coupled fitting proceed similarly as described above:

1. Select a regional TAC with good signal.

2. Switch to the model (**2 Tissue Compartments, K1/k2**) to be applied in coupled fitting. Fit the TAC with **Fit model, current region** and propagate the model to all other regions with **Model & Par**.
**Par. Fit model, all regions** to establish reasonable starting values for all TACs.

3. On the **Coupled** panel, enable the parameter(s) which are to be treated as common, and enable all target regions which should be fitted together. Note that the disabled parameters will be estimated per region.

4. Start processing with **Fit coupled**.

5. To use the common parameters for fitting all regions please proceed as described for the common $k_2'$ example above.

**Coupled Fitting Procedure**

The coupled fitting process works as follows:
- A global target curve is created by stacking the TACs of the checked regions (valid points only).
- The corresponding weights are calculated according to the weighting configuration for each TAC.
- A table of fitted parameters is formed by entering the common parameters once and adding all the other fit enabled parameters in the coupled regions. The initial values of common parameters are taken from the first checked region, for the other parameters from the individual regional models. If **Float blood parameters** is enabled on the **Extras** pane, the fit-enabled parameters of the blood curves are also taken into account. As a convenience there is a button **Fit coupled and delay** to estimate a blood delay which considers the information from all coupled regions.
- The optimizer calculates the model curve in all checked regions, creates the global result curve corresponding to the target curve, weighs all the residuals and forms the global sum of squares as the figure of merit. The parameters are adjusted until the cost function has been minimized, and the resulting parameters are copied back to the individual models.

### 4.7.2 Common Parameters across Regional TACs from Different Studies

**Preparations**

For a simultaneous fit of TACs from multiple studies the user first has to

- load all of the data sets so that they are available on different workspace tabs;
- ensure that all of the TACs to be coupled have the same compartment model configuration and the same residual weighting;
- ensure that all model parameters have reasonable initial values by either fitting the models, or propagating a model configuration.

The coupling is performed using a separate dialog window which can be opened from the **Kinetic** menu with the **Coupled Studies Dialog** entry.

**Study and Model Selection**

The dialog window opens on the first tab **Study/Model selection** containing three sections.
The **Study selection** lists all open data sets. As soon as one of them is selected in the list, the **Model selection** list is updated, showing the different models which are used for the TACs of the study (only models used in same region across studies). In the example shown above one of the regions uses a **1 Tissue Compartment** model, whereas the regions intended for coupling use the **2 Tissue Compartments, K1/k2** model.

After selecting the appropriate model, the **Parameters selection** list is updated showing all parameters of the model together with a **Coupling** selection. A parameter can have one out of three configurations:

- **NOT COUPLED**: The parameter will be individually fitted in the different regional models or kept fix depending on the state of the individual fit box.
- **REGIONAL**: The parameter will be coupled among all regions with the same name. For instance, the parameter will have a common value for all *caudate* regions, a different common one for all *putamen* regions, etc. Note that with this configuration the state of the individual parameter fit boxes is not relevant, the parameter will be modified in all coupled regions.
- **GLOBAL**: The parameter will be coupled among all coupled regions independent of their name. For instance, the parameter will have the same common value among all *caudate, putamen*, etc regions. Note that with this configuration the state of the individual parameter fit boxes is not relevant, the parameter will be modified in all coupled regions.

**Regions Selection and Fitting**

The next step is defining the TACs included in the coupled fit on the **Regions selection** pane as illustrated below.
The list contains all regions which are defined in the coupled studies with exactly the same names. The regions are selected for inclusion by the Include checkbox. Initially, all regions are included. When working with a large number of regions (e.g. from atlas definitions) the context menu may be helpful which can be opened by right clicking into the list. It contains Select all and Select none for quickly selecting or de-selecting all regions, respectively.

Coupled fitting is started using the Fit coupled button and proceeds as follows:

- A global target curve is created by stacking the TACs of all coupled regions from the selected studies (valid points only).
- The corresponding weights are calculated according to the weighting configuration for each TAC.
- A table of fitted parameters is formed by entering the GLOBAL parameters once, the REGIONAL parameters once per region, and adding all the other fit enabled parameters in the coupled regions. The initial values of common parameters are taken from the first study, for the other parameters from the individual regional models. Note that Fit blood parameters has no impact on fitting, all the parameters of the blood curves remain fixed.

The optimizer calculates all model curves, creates the global result curve corresponding to the target curve, weighs all the residuals and forms the global sum of squares as the figure of merit. The parameters are adjusted until the cost function has been minimized. If the box Random Fits is checked, the fitting will be repeated as many times as specified with randomly changed starting parameters, see Fitting Options. Grid fitting works similarly stepping the initial parameters within intervals.

Result Parameters
After the coupled fit has completed, the resulting parameters in the coupled regions are listed on the Result parameters pane which is immediately shown.
Note in the example above that the **REGIONAL** parameter $K_1/k_2$ has the same value in all regions with the same name, whereas the **GLOBAL** parameter $k_4$ has the same value in all regions.

The results can be prepared for use in another program such as MS Excel with the **Copy to Clipboard** button. The **Use results** button transfers the result parameters together with their standard errors to the individual models and closes the dialog window, whereas the results are discarded by the **Close** button.

### 4.8 Assessing Fit Results

The physiologic behavior of PET tracers is usually quite complex. A comprehensive model accurately describing its kinetics therefore requires many compartments and exchange parameters. However, as the PET signal is limited in quality and information, the model must be simplified to a degree that only a few parameters remain. When estimating a model with many parameters, the variance of the parameter estimates tends to be very high, so that a reliable interpretation of the results becomes impossible. A simplified model with fewer parameters will provide more precise results, but these parameters may be biased. Therefore, the optimal trade-off between bias and uncertainty has to be sought by testing models of decreasing complexity.

After every model fit detailed information about the parameter estimates and the goodness-of-fit is available on the **Details** tab.
Parameter Standard Error

Nonlinear regression using the Levenberg-Marquardt optimization reports a standard error (SE) for each fitted parameter value. The SE is calculated by multiplying the diagonal elements of the covariance matrix obtained by the fit with Chi Square and taking the square root. Note that SE is displayed as % of the actual parameter value.

Calculation of the macroparameter SE involves the covariance matrix and the partial derivatives at the solution. Example of the SE calculation for Vs of the 2-Tissue compartment model (Vs=K1k3/k2k4):

\[
\text{if } (k2k4 > 0) \{ \quad \text{// Calculation of the partial derivatives (pder) of Vs at the solution} \\
pder[K1\_IND] = k3 / k2k4; \\
pder[K2\_IND] = -K1 * k3 / (k2 * k2k4); \\
pder[K3\_IND] = K1 / k2k4; \\
pder[K4\_IND] = -K1 * k3 / (k4 * k2k4); \\
\text{variance} = \text{varianceFromModelCovar}(pder, covar); \quad \text{// matrix multiplication:} \\
\text{SE} = \sqrt{\text{variance} * \text{chiSqr}}; \\
\}
\]

The reason why the SE of a macroparameter can be lower than that of the parameters involved in the calculation is that the parameters are correlated.

Parameter Confidence Intervals

If the inherent fitting assumptions are true, a 95% confidence interval can be approximated by the result parameter plus/minus two standard errors. This confidence interval is displayed for each fitted parameter ([Conf.low, Conf.high]).

There will be a 95% chance that the confidence interval contains the true parameter value. A sufficiently narrow confidence interval indicates that the parameter could be determined with a reasonable certainty, whereas a wide interval makes it necessary to revise the configuration of the used model, or look for a more appropriate model.

Goodness-of-Fit

The following measures are defined which allow a direct or indirect assessment of the goodness-of-fit:

<table>
<thead>
<tr>
<th>DOF</th>
<th>Degrees of freedom</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>defined as the number of valid measurements minus the number of fitted parameters.</td>
</tr>
</tbody>
</table>
### 4.8.1 Parameter Comparison across Regions (PKIN Parameters Explorer)

A convenient overview of the models and their parameters can be obtained using the **Study-wide parameter explorer** illustrated below. It shows a similar dialog window as the **Fitting History** window, with the **Region** list entries corresponding to the different regional TACs of the data set.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>SumSquared</td>
<td>Sum of squared (unweighted) residuals.</td>
</tr>
<tr>
<td>ChiSquare</td>
<td><strong>Reduced Chi square.</strong> Sum of squared, weighted residuals, divided by the degrees of freedom. The reduced chi-square provides a useful measure of goodness-of-fit. If the model describes the measured data, the reduced chi-square will mostly represent the variance of the data and will be close to 1.0 (when weighting is appropriate).</td>
</tr>
<tr>
<td>AIC</td>
<td><strong>Akaike Information Criterion.</strong> The AIC methodology attempts to find the model that best explains the data with a minimum of free parameters. The AIC is calculated with the second order correction for small sample size (&lt;40). The preferred model is the one with the lowest AIC value.</td>
</tr>
<tr>
<td>SC</td>
<td><strong>Schwartz Criterion.</strong> Also called Bayesian Information Criterion (BIC). The preferred model is the one with the lowest SC value.</td>
</tr>
<tr>
<td>MSC</td>
<td>Another criterion used in the <strong>Scientist</strong> Software (MicroMath, Saint Louis, Missouri USA) is the <strong>Model Selection Criterion.</strong> The preferred model is the one with the highest MSC value.</td>
</tr>
<tr>
<td>R2</td>
<td>There is also a measure of the goodness-of-fit, the <strong>coefficient of determination</strong> R2, a number between 0 and 1. A value of 0 means that the fit is not better than a horizontal line through the mean of all measurements, whereas a value of 1 means that all measurements lie exactly on the curve. High R2 values indicate that the model curve is close to the measurement.</td>
</tr>
<tr>
<td>Sy.x</td>
<td>Another information about the residuals is provided by the <strong>root mean square value Sy,x.</strong> It is defined as the standard deviation of the residuals and can be used to generate synthetic measurements in Monte Carlo simulations, provided all measurements have the same variability.</td>
</tr>
<tr>
<td>Runs test p</td>
<td>The runs test is a statistical test to decide whether the model curve deviates systematically from the data. It is based on the number of runs resulting from the fit. A run is a set of consecutive measurements which are above (positive residuals) or below (negative residuals) the measurement. Given the assumption that the residuals are randomly distributed, the probability p of the occurrence of a number of runs can be calculated. If p is small (eg. p&lt;0.05) the measurements systematically deviate from the model curve. Such a finding signals that most likely an inadequate model was fitted and further investigations of the result are not sensible.</td>
</tr>
<tr>
<td>AUC</td>
<td>Area under the model curve.</td>
</tr>
</tbody>
</table>

**Note:** The results shown on the **Regions** tab depend on the configuration settings. After changing the configuration, the window has to be closed and re-opened.
The column maximum is highlighted in red, the minimum in green. Values can be sorted by clicking into the column headers, and the parameter order changed by dragging the column headers left/right. Data export is facilitated via Copy to Clipboard.

The Last revision panel shows the entries of the last fitting operation. It is most useful for Macro Results Inspection. It will only show a single line after fitting an individual region, and the same information as the Regions panel after fitting all regions.

**Configuration Settings**

The configuration button opens a dialog for defining the information to be displayed.

<table>
<thead>
<tr>
<th>Include standard error (% SE) in table</th>
<th>For each model parameter add a column with the % standard error.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Include distribution statistic at table bottom</td>
<td>Show the lower section with min/average/median/max/standv.</td>
</tr>
<tr>
<td>Use last region fits for table</td>
<td>Show the last successful fit from the history for each region. Otherwise the current regional model configuration will be listed, whether fitted or not.</td>
</tr>
<tr>
<td>Display tab with parameter plots</td>
<td>Visualization of the parameter values as a curve. Select the regions of interest on the Parameters panel.</td>
</tr>
</tbody>
</table>
The parameter plots are then shown on the Display parameters panel.

**Display tab with relative parameter plots**
Visualization of the normalized parameter values as a curve. Select the regions of interest on the Parameters panel. The Display relative change curve panel then shows the plot of each parameter normalized to the value of the last region.

**Display tab with model curves**
Visualization of multiple regional TACs together with their model curve. Select the regions of interest on the Parameters panel. The Display model curve panel then shows the data and the models.
4.8.2 Comparison of Model Curves for a Region

In addition to comparing the same data in different PKIN windows, model curves from different model fits can be plotted in a single curve display. The facility is accessible via the fitting history mechanism, after enabling Display tab with model curves.

Now the model fits of interest can be selected on the History tab, and the corresponding model curves together with their targets inspected in the Model curves panel.
4.8.3 Comparison using two PKIN Windows

PKIN allows the user to examine different models and compare their numerical output using the History mechanism. Sometimes it is also helpful to process the same data with different model configurations in parallel. This can be achieved using the Study Compare Dialog entry in the Kinetic menu.

A new dialog window is opened with a clone of the current data set. It supports all features of PKIN except for the functions related to data management. So the user can arrange the original PKIN window and the dialog window next to each other and process/visualize the same data in parallel.

4.8.4 Statistical Criteria for Comparing Models

The following criteria can be used for comparing different models which use the same weighting and are fitted to the same data. The notation is:

- $n$: Number of independent measurements considered in the fit
- $p$: Number of fitted parameters
- $w_i$: Weight applied to residual of acquisition $i$

**Akaike Information Criterion (AIC)**

The Akaike Information Criterion [1] is defined by the formula

$$AIC = n \ln \left( \sum_i w_i [y(t_i) - \hat{y}(t_i)]^2 / n \right) + 2p$$

If applicable, PKIN uses an adjusted procedure with a second order correction for small sample size ($<40$) [2]

$$AIC = n \ln \left( \sum_i w_i [y(t_i) - \hat{y}(t_i)]^2 / n \right) + 2p + \frac{2p(p+1)}{n-p-1}$$

The more appropriate model is the one with the smaller AIC value.

**Schwartz Criterion (SC)**

The Schwartz Criterion is defined by the formula

$$SC = n \ln \left( \sum_i w_i [y(t_i) - \hat{y}(t_i)]^2 / n \right) + p \ln(n)$$

The more appropriate model is the one with the smaller SC value.

**Model Selection Criterion**
Another modified version of the AIC criterion used in the Scientist Software (MicroMath, Saint Louis, Missouri USA) is the Model Selection Criterion.

\[
MSC = \ln \left( \frac{\sum_{t} w_t [y(t_i) - \hat{y}(t_i)]^2}{\sum_{t} w_t [y(t_i) - \hat{y}(t_i)]^2} \right) - 2p/n
\]

This criterion has the advantage that it is independent of the magnitude of the \( y \). Opposed to the AIC and the SC the more appropriate model is that with the larger MSC.

**F-Test**

Two nested models can be compared by an F-test of their sum of squared residuals [3]. The idea is to compare the total sum of squares into a component removed by the simpler model and into a component additionally removed by the more complex model. For each component, the mean square (sum of squares per degree of freedom) is calculated. The residual mean square is an estimate of the variance of the original data. The ratio of the two mean squares is the F-statistic used to test for significance of the variance reduction by the additional parameters, as follows:

\[
F = \frac{(Q_1 - Q_2)/(p_2 - p_1)}{Q_2/(n - p_2)}
\]

where

- \( Q_1 \) represent the weighted residual sum of squares or the simple model with \( p_1 \) parameters,
- \( Q_2 \) the weighted residual sum of squares of the more complex model with \( p_2 \) parameters,
- \( p_1 < p_2 \)

The F-statistics has \((p_2-p_1, n-p_2)\) degrees of freedom. If the calculated F is larger than the tabulated value at a specified \( p \) value, the reduction of the residual variation by the addition of the \((p_2-p_1)\) extra parameters of the more complex model is statistically significant. Usually, significance \( p=0.05 \).

Note: in MS Excel the \( F_{p_2-p_1, n-p_2} \) can be calculated by the function FINV\((p;p_2-p_1;n-p_2)\).

**References**


**4.8.5 Analysis of Parameter Sensitivity and Identifiability**

The following two approaches can be employed to investigate whether a selected model is too complex for the data. The implementation in PKIN is based on the handout *Kinetic model evaluation with sensitivity functions and correlation matrices* by Dr. M.M. Graham, University of Washington at the Technical Exhibition of the Society of Nuclear Medicine Annual Meeting 1995 (JNM vol 36, no 5, P1209). As to our knowledge, there is no follow-up paper about this subject.

**Sensitivity**

If a parameter value is changed there should be some visual change in the model output. Some parameters will cause more change than other. The model is more sensitive to a parameter that causes more change in the output.
The sensitivity functions $\text{Sens}(k_i)$ for the different parameters $k_i$ are obtained by calculating the model function twice: with the initial parameter set, and after changing the parameter $k_i$ by 1%. Then, the expression

$$\text{Sens}(k_i, t) = 100\% \frac{\text{Model}(k_i \times 1.01, t) - \text{Model}(k_i, t)}{\text{Model}(k_i, t)}$$

is calculated over time. When the 1% change of $k_i$ has changed the output by 1% at a certain time the sensitivity function is equal to 1. In PKIN, the sensitivity functions are calculated for all model parameters, whenever the Sensitivity pane is selected. The example below shows the sensitivity functions of a FDG data set.

Examination of the sensitivity curves shows at what time a parameter has significant sensitivity and how sensitive a parameter is compared to another. If the sensitivity curves of two parameters have the same shape it is unlikely that one can tell which parameter has changed (i.e. they are not identifiable).

In the example above the model is quite insensitive to $v_B$. Furthermore, the $k_3$ and $k_4$ curves have a quite similar (but opposite) shape throughout the acquisition duration, indicating that there is a substantial correlation between these parameters.

**Identifiability**

The model output must change in a unique way when changing a model parameter. If the change in the model output is the same when changing either of two parameters they can not be independently identified. In this situation the model must be simplified further, or one of the parameters can be fixed at a physiologic value. The latter approach is better since simplification usually means essentially setting one of the parameters equal to zero or infinity.

A more quantitative way to look at parameter identifiability is to generate a correlation matrix. It shows how the various parameters tend to correlate with each other. A low value, close to 0, means they do not correlate and that they are identifiable from one another. A high value, close to 1, means they are highly correlated and not independently identifiable.

The correlation matrix is calculated by:

1. generating the sensitivity matrix $SM$ given from the sensitivity functions by $SM(i,j) = \text{Integral}(\text{Sens}(k_i, t) \times \text{Sens}(k_j, t))$;
2. inverting the sensitivity matrix $SM(i,j)$, resulting in the covariance matrix;
3. normalizing the covariance matrix by dividing each element by the square root of the product of the corresponding diagonal elements; result is the correlation matrix.

The correlation matrix is shown in the Sensitivity pane. Using the Save button the correlation matrix and the sensitivity functions can be save into a text file.
4.9 Special Processing Types

4.9.1 Parameter Sensitivity to Acquisition Duration

A standard task when analyzing kinetic data of a new tracer is to assess the dependence of the parameters on the length of the available data. This type of analysis can easily be performed using the Data shortening option on the Extras panel.

As soon as this option is enabled with a minimal data duration, the fit buttons are labeled accordingly

When activating Fit model, current region, shorten the following happens:

1. The set of initial parameters is created.
2. A fit is performed with the minimal data segment (e.g. the first 20 min), corresponding to a small number of Degrees Of Freedom (e.g. DOF=16). All other fit options on the Extras panel are also active.
3. An additional sample is added to the data segment and the fit repeated with the same initial parameters (DOF=17).
   This step is repeated until the full data segment has been fitted.

Results

The results are stored in the model fitting history and can be opened with the button. All sensitivity fits have the same Revision number. The data segment length is represented by the DOF column. In the curve plot the unused samples at the end are indicated in grey.
The results of the sensitivity analysis are most conveniently saved by selecting an entry with the appropriate Revision and using the Save Selected Revision function illustrated below. It saves the fitting results from all regions into a .kinPar file.

Sensitivity Visualization
A visual impression can be obtained by stepping through the fitting history with the forward and backward buttons and observing the shape of the model curve as the number of included samples changes.

For a visualization of the parameters the results can be exported via Copy to Clipboard, but the trend can also directly be visualized on the two panels Parameter plots and Relative parameter plots. The selection in the history list defines the data to be visualized:

- Selection of a single line as illustrated above: all entries with the same Revision number are used.
- Explicit selection of a subset of lines: only the selected data is used.

When selecting the Parameter plots panel the absolute parameter values are visualized. Each parameter corresponds to a curve and can be enabled/disabled with the checkboxes in the curve control area. The x-axis value corresponds to the number in the No column, so the length of the data segment increases to the right.
When selecting the Relative parameter plots panel the same curves are shown, but the values are normalized to the last point (which in the case of the sensitivity fitting can be considered as the most reliable one).

**Leave-one-Out**

The operation behaves differently, if a negative Data shortening to time is specified. In this case, not the whole end segment at the end is disabled, but only a single point, as indicated below. Using the model history, the user can easily step through the different fits with a single measurement disabled and see the impact on the result parameters.

### 4.9.2 Monte Carlo Simulations

The purpose of performing Monte Carlo simulations in kinetic modeling is to get information about the standard error of the parameter values found in a fit. There are two different kinds of sources which contribute to the uncertainty of the parameters:

1. All the input data are measured and hence prone to measurement errors.
2. The model may contain more parameters than supported by the data, so that the effect of one parameter may be counter-balanced by another parameter.
The paradigm used for performing Monte Carlo simulations is described by Flannery et al. [11]. The basic idea is to simulate a series of measurements and statistically analyze the results when modeling these synthetic "measurements".

In PKIN Monte Carlo simulations can be performed for models applied to a single tissue and models coupled across several tissues by selecting the respective option button on the **Mt-Carlo** panel.

### Noise Specification

Three types of "noise" can be defined. For each "noise" the distribution type as well as the variance has to be configured. After selecting the distribution a dialog window appears for defining the variance as described in the Residual weighting for model fitting.

1. **Add noise to tissue** TAC. Note that here only the noise distribution needs to be selected. The noise variance is adopted from the **Weighting** tab on the **Tissue** pane. If Add noise to measurement is not selected, noise is added to the tissue model curve, otherwise to the measured TAC. The latter e.g. allows studying the effects when fitting a simplified model to the data.

2. **Add noise to blood** allows simulating noisy input curves. This option is only supported for blood models of type Lin. Interpolation.

3. **Variation of initial parameters** is disturbing the starting conditions from the ideal solution. For instance 20% noise can be added to the ideal parameter values.

### Noise Distributions

Three different distributions are available for noise generation, Gaussian, Poisson, and Uniform. They are applied for each individual measurement with the prescribed variance.

For the tissue noise simulations there is also the **Bootstrap** method available. With the **Bootstrap** approach, the weighted residuals \( w_r \) of the initial fit are used [73]. For simulating a noisy curve, these residuals are randomly selected and added to the model curve. Specifically, the noise for a frame \( j \) is constructed from a randomly selected \( w_{r_i} \) of frame \( i \) by dividing it by the weight of frame \( j \). As a consequence, it will have the same effect in the fit as it originally had in frame \( i \).

### Monte Carlo Process

When the **Start Monte Carlo for current region** button is selected the following processing starts:

1. An initial fit is performed. From then on it is assumed that the result parameters represent the "true" parameters.

2. The number of **Simulations** times a noisy data set is prepared according to the definitions and then fitted. Noise is added to the model (or tissue) curve, and optionally the input curve. The "true" parameters, optionally disturbed by some noise, are used as starting values, and the fit to the noisy data performed. If **Record results in fitting history** is checked, the result is
saved to the fitting history together with the noisy TAC. Otherwise, just the resulting parameter values are recorded.

Note: For non-compartment models the noise is added to the measured values, as the target may represent transformed data such as a graphical plot. For reference tissue models noise is only added to the target TAC.

3. After all runs have completed, the distribution of the result parameters is analyzed resulting in a mean and a standard deviation value for each fitted parameter. These values are shown on the Standard tab of the Tissue page, and the model curve calculated with the mean parameters is shown. Note that often it does not follow the measurement as closely as with fitted parameters. The Monte Carlo results should now be saved, because as soon as a parameter is changed or a new fit is initiated, the results get overwritten.

```
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Current value</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>K1</td>
<td>0.681873</td>
<td>ml/cm/min</td>
</tr>
<tr>
<td>k2</td>
<td>0.127204</td>
<td>1/min</td>
</tr>
<tr>
<td>k3</td>
<td>0.027436</td>
<td>1/min</td>
</tr>
<tr>
<td>k4</td>
<td>0.048202</td>
<td>1/min</td>
</tr>
<tr>
<td>Vd</td>
<td>0.305002</td>
<td>ml/cm</td>
</tr>
<tr>
<td>k1</td>
<td>0.841053</td>
<td>ml/cm</td>
</tr>
<tr>
<td>k2</td>
<td>0.053501</td>
<td>ml/cm</td>
</tr>
<tr>
<td>k3</td>
<td>0.051324</td>
<td>1/min</td>
</tr>
<tr>
<td>LambdaK</td>
<td>0.014865</td>
<td>ml/cm/min</td>
</tr>
<tr>
<td>ChiSquare</td>
<td>0.93545</td>
<td></td>
</tr>
</tbody>
</table>
```

Note: Fitting uses all settings defined on the Extras panel.

It is possible to visualize a summary of the fit results during the Monte Carlo runs by switching back from the Tissue to the Mt-Carlo pane. In the curve display, a new curve Monte Carlo Data Min/Max appears with vertical bars. The end points of these bars mark the minimal and the maximal value that any of the resulting model curves reached at a particular time point.

Fitting History

The value of using the fitting history is illustrated below after a Monte Carlo simulation with enabled Record results in fitting history. Each entry from the simulation has a common Revision number. The Monte Carlo Initial Fit line contains the result of the first fit to the original data. If Add noise to measurement is not selected, the model curve shown is the curve to which the generated noise is added. The following entries Monte Carlo Simulation are the individual fits to the generated noisy TACs. The final line Monte Carlo Result represents distribution of all individual fits: the parameters are calculated as the average across the individual fits, and the standard error as their standard deviation.
Double-click into the history to show the noisy TAC and the fit result of a simulation. Use the arrows to conveniently step through the simulations.

Note that the results of the simulations can also be visually presented by selecting all simulations in the History panel and selecting the other panels. The Parameter plots and Relative parameter plots show the individual parameters across the simulations as absolute and relative values, respectively. Model curves shows all noisy TACs with the respective fitted model curves.

The individual fit results can also be saved for dedicated statistics. To this end select an entry with the Revision of the Monte Carlo simulation, then use Save Selected Revision as illustrated below.

Note: When saving a .km file, the history is included. Therefore the file created can become large when including extended Monte Carlo simulations.
Several Tissues Coupled
The same principle as for the single tissue Monte Carlo simulation applies to a coupled Monte Carlo simulation. It can be started by Start Monte Carlo for coupled regions and uses the configuration prepared on the coupled panel. Model noise uses the weighting individually defined for each coupled region. The fitting history of all coupled regions uses the same Revision number, so that all results can be exported at once.

4.9.3 Batch Mode

The Batch Mode serves for the sequential processing of multiple data sets. It can be used for fitting with a range of models, Monte Carlo simulations and to aggregate the parameters from existing KM files.

Batch processing is started from the Kinetic menu and opens a dialog window for setting up the batch queue as illustrated below. There are two batch types:

1. **Batch based on Macros** to sequentially fit models to the data.
2. **Batch based on Model Definitions** either in the data file itself or via loaded .kmModel definitions. Here a variety of processing types are supported.

Data Files
Two types of data files can be processed:

- **KM files**: A .km file saved from PKIN includes a complete definition of the data and the models for the different types of information (blood interpolation model, the tissue model, weighting etc) for each region. Therefore, multiple processing types are supported including coupled fitting and Monte Carlo simulations. Additionally, during fitting, the **options** on the **Extras** pane such as parameter initialization and randomized fitting will be effective. A new KM file will be created with the fitting results in the **fitting history**.

- **Composite .kmData files**: A .kmData file contains only the **data definition**, except for the global specification of a model. Therefore, only the individual tissue model fitting is supported. The result will be saved as a new .km file with the same name as the input file, and with the results in the history.

Use the **Add data** button for opening a data selection window and select the data sets to be processed from a database or the file system. Further data can incrementally be added, and selected list entries removed.

**Starting the Batch**

The **Run** button starts processing, and PMOD will be blocked until processing completes.

**Batch Results**

The **New file** and the **Append to file** buttons can be used to specify a .kinPar file as a container for saving the result parameters of the batch run. These results can be transferred to the R console for performing statistics.

For allowing visual inspection of the results each original data file is replaced by new file. It contains in the **history** all model fits from the batch (as illustrated below), so that a user can simply load the file and step through the fits using the history buttons.

**4.9.3.1 Batch based on Macros**

**Macros** are a convenient way of fitting multiple models to the data. They already include information about fitting options, so the batch configuration interface has a simplified appearance.
Add macro allows selecting .kmMacro definitions to set up the analysis queue. Note that the default macros can be loaded by using Load from File System in the appearing dialog window and navigating to the PMOD subdirectory resources/pkin/macros.

**4.9.3.2 Batch based on Model Definitions**

If .km files have been selected it is assumed that appropriate configurations are included for the following Operations:

<table>
<thead>
<tr>
<th>Operation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fit each individual region per file</td>
<td>All regional TACs are fitted with the respective models as defined in the .km file.</td>
</tr>
<tr>
<td>Perform one coupled fit per file</td>
<td>Performs coupled fitting with the model and coupling as defined in the .km file.</td>
</tr>
<tr>
<td>Perform a Monte Carlo simulation for the active region per file</td>
<td>Using the noise and model definitions in the file a Monte Carlo simulation of the current TAC is performed.</td>
</tr>
<tr>
<td>Perform a Monte Carlo simulation for each region</td>
<td>As above, but applied to all regional TACs. Can for instance be used to evaluate the sensitivity at different parameter combinations.</td>
</tr>
<tr>
<td>Perform one coupled Monte Carlo simulation per file</td>
<td>Monte Carlo simulation using the coupled fitting and Monte Carlo settings defined in the file.</td>
</tr>
<tr>
<td>Copy model parameters from KM to kinPar file</td>
<td>A convenience tool for aggregating the parameters of number of KM files into one tabular text file. Note that a Parameter Results file must be defined with the New file button, or the results can be appended to an existing file selected with the Append to file button.</td>
</tr>
</tbody>
</table>

**Batch with Model Configuration Files**

The batch mode also allows fitting multiple models to each data file. To do so, corresponding .kmModel files can be selected with the Add model button in the Kinetic.
In this situation, only the **Fit each individual region** option is available. The results of the different model fits are saved in the regional model histories.

**Fitting Options**

The **Model fit options** give access to the mechanisms for improving objectivity and reliability of the results discussed in **Fitting Options on Extras Panel** with one exception. The **First fit blood models** option is intended for cases, where interpolation functions are defined for the different blood components but not yet fitted.

### 4.9.4 Blood Delay Fitting

The timing of the tissue and the blood data should be synchronized to a common clock. Nevertheless, there may be an inherent shift of the blood activity curves relative to the tissue activity curves because of the arrival time difference of the blood in the target tissue and at the sampling location. For brain PET scans with radial artery blood sampling the tracer will appear later in the blood samples than in the brain. This delay is even prolonged if the blood samples are not taken directly with a syringe, but via a catheter which adds additional pathlength as required by online blood sampling devices.

Such a relative delay of the blood samples is accounted for by the **Delay** parameter of the **blood models**. Positive delays represent delayed blood information and hence shift the blood curves to earlier times (to the left). While both the whole-blood and the plasma model include a separate **Delay** parameter, it is reasonable to assume a common delay since the plasma activity is derived from whole-blood. Such a common delay can be fitted together with the kinetic parameters of the tissue model as described below.

**Delay Fitting with Single Tissue TAC**

The **Tissue tab** features a dedicated **Fit region and blood delay** button. It is hidden underneath the **Fit current region** button and can be activated as illustrated below.

When it is activated, the following actions are performed:

- The state of the fit flags of the **Whole blood** and **Plasma** models is saved.
- The fit flag of the **Delay** is enabled, and all other fit flags disabled.
- A fit of the tissue model is performed, whereby the **Delay** (common to plasma and whole blood) is also optimized.
- The state of the fit flags of the **Whole blood** and **Plasma** models is restored.
The estimated Delay can be inspected on the Blood tab or in the model history. The illustration below shows a history of the outcome when fitting a 1- and 2-Tissue compartment model to the WB TAC including the delay.

<table>
<thead>
<tr>
<th>No</th>
<th>Type</th>
<th>Region</th>
<th>Model</th>
<th>Blood delay</th>
<th>K0</th>
<th>K1</th>
<th>K2</th>
<th>K3</th>
<th>K4</th>
<th>V1</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Fit</td>
<td>WB</td>
<td>1 Tissue Compartment</td>
<td>0.0, 0.06</td>
<td>0.61412</td>
<td>0.08398</td>
<td>0.720</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Fit</td>
<td>WB</td>
<td>1 Tissue Compartment</td>
<td>0.128192</td>
<td>0.05</td>
<td>0.05203</td>
<td>0.05247</td>
<td>0.7276</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Fit</td>
<td>WB</td>
<td>2 Tissue Compartment</td>
<td>5.412845</td>
<td>0.05</td>
<td>0.050505</td>
<td>0.134906</td>
<td>0.038107</td>
<td>0.058012</td>
<td>0.6160</td>
</tr>
</tbody>
</table>

It is notable that the optimal delay depends on the fitted model, and it will also depend on the selected regional TAC. When using a single TAC it is recommended using the signal from a relatively large VOI, for instance the whole brain, and then keep the delay fixed.

**Note:** A change of the blood Delay affects the model curve of all regions and therefore requires re-fitting all regional models. If the Delay value is changed manually, it has to be set to the same value for Blood and Plasma.

When using Fit model, all regions, the blood parameters will not be fitted separately for each region, if Fit blood is enabled on the Extras panel. For example, with the default Lin. Interpolation model for Blood and Plasma with the Delay parameter fit enabled, the blood delay will be estimated regionally as illustrated below.

**Delay Fitting with Multiple Tissue TACs**

An alternative to using a single representative tissue is the use of coupled fitting for the determination of a blood delay which is optimal with respect to tracer appearance in several regions. In the example below 8 regions are selected on the Coupled pane all of which have a 2-tissue compartment model configured. Note that at least one tissue parameters needs to be coupled in this case.

Fitting of the common delay is started using the Fit coupled and blood delay button which is hidden under the Fit coupled button. The procedure will fit all 8 compartment models while at the same time optimizing the blood timing.
4.9.5 Fitting Blood Shape Parameters and Tissue Parameters together

The PKIN tool allows the simultaneous fitting of the shape parameters of the blood models and the kinetic parameters of the tissue model. This fitting mode can be activated by checking the Fit blood parameters box on the Extras pane. It is active in the fitting of the single tissue TAC as well as in coupled fitting.

With Fit blood on, ALL parameters of the Whole blood and the Plasma model (but not the plasma fraction or parent fraction parameters) which have the fit check on will be modified during the fit of the kinetic tissue model. In the situation illustrated below 9 parameters (plus additional on the Whole blood panel) will be fitted at the same time.

Note: Only the tissue model curve is fitted to the measurement. The blood models are not fitted to the actual blood measurements. It is only the impact of the blood curves on the calculated tissue model curve which results in the modification of the blood model parameters.

4.9.6 Dispersion Correction for Automated Blood Samples

Automated Blood Sampling Setup

Automated blood sampling is the preferred method for measuring the radioactivity concentration in whole blood. In such a setup, a catheter is inserted into an artery, and blood continuously run through a detector which measures the radioactivity from the time of tracer injection until the end of the PET acquisition. If needed, automated blood sampling can be stopped when the blood concentration changes have become slow, and some manual blood samples be withdrawn at later times.
A typical blood sampling setup is illustrated below with a catheter in the radial artery, which is run through a coincidence detector (twilite, www.swisstrace.com) and a pump which ensures constant blood drawing speed. Three-way valves serve for occasionally taking blood assays, which are needed for measuring the metabolite build-up over time.

Blood Dispersion

Because blood needs to travel an extra distance outside the body until it reaches the detector, the blood measurements are delayed. Furthermore, the time course of the concentration is altered, because fast concentration changes are smoothed in the catheter, an effect called "dispersion". In kinetic modeling, results can be biased if dispersed blood information is used, particularly for short measurements such as $^{15}$O-water bolus studies for quantifying tissue perfusion.

The sections below describe the methodology and implementation of a dispersion correction methodology developed by Munk et al. [1].

4.9.6.1 O. Munk Dispersion Correction Methodology

Dispersion Correction

Several dispersion correction methods have been developed. Most of them describe dispersion as a convolution of the true concentration curve with a dispersion function, and correction amounts to a numerical deconvolution which produces results suffering from excessive noise. Munk et al. [1] have developed an alternative approach which circumvents numerical deconvolution. It describes transport of blood through a catheter by a "transmission-dispersion" model which includes two components: molecules which travel undisturbed in the inner of the catheter (convective flow), and molecules close to the catheter wall to which sticking occurs (stagnant flow).

The true (corrected) blood activity concentration $C_B(t)$ can be calculated from the measured concentration $C_{mb}(t)$ as follows.

$$C_B(t) = \frac{C_{mb}(t+T)}{1-\alpha} - \frac{\alpha k}{(1-\alpha)^2} \int_0^{\infty} C_{mb}(\tau+T)e^{-k(\tau-t)/(1-\alpha)} d\tau$$

There are three parameters in the equation which are characteristic for the experimental acquisition setup:

- the transit time $T$,
- a parameter $k[1/\text{min}]$ which refers to the stickiness of the catheter for a tracer, and
the stagnant fraction $\alpha$. If $\alpha=0$, no dispersion occurs, just a delay. $\alpha=1$ is not allowed in the equation, so $0 \leq \alpha < 1$.

The effect of $\alpha$ and $k$ on the corrected curve shape is illustrated below with two plots from [1]. The dashed line represents the dispersed, measured concentration, and the solid lines the true concentration recovered by correction.

![Plot 1](image1.png)

**Calibration Measurement**

The parameters $k$ and $\alpha$ have to be obtained from a calibration measurement with exactly the same conditions as the live experiment, i.e. with same catheter length, withdrawal speed and tracer as described in [1]. Basically, two beakers are prepared: one beaker with blood only, the other with blood and tracer (taken 1 min after tracer infusion). A three-way tap is used with one catheter leading to the blood sampling system, and the other two into the beakers. Blood sampling is started with connection of the blood-only line to the detector for measuring the baseline with no radioactivity. Then the tap is switched to the catheter with blood and tracer. Correspondingly, the measured radioactivity is rising up to a constant height. Sometime later the tap is switched back to the blood-only line, and the radioactivity is decreasing back to the baseline. Instead of the true rectangular shape of the concentration at the tap a concentration shape similar to the dispersed rectangle below [1] is measured.

![Plot 2](image2.png)
Formula can be derived [1] describing the step-up as well as the step-down edge as a function of the dispersion parameters $k$ and $\alpha$, as well as the transit time $T$:

**Step-up:**

$$C_u(t) = \begin{cases} 0 & t < T \\ C - \alpha C e^{-\frac{t}{k}} & t \geq T \end{cases}$$

**Step-down:**

$$C_d(t) = \begin{cases} 0 & t < T \\ \alpha C e^{\frac{t}{k}} & t \geq T \end{cases}$$

By fitting these functions to the measurement, the parameters can be determined and subsequently used in the correction of any live measurement using the same setup.

**Reference**


### 4.9.6.2 O. Munk Dispersion Calibration in PKIN

**Calibration Measurement**

A calibration measurement should be performed as described above [1], measuring the response in the blood counter to a rectangular input activity. Note that the measured data have to be decay corrected and calibrated to kBq/cc.

**Fitting of the Dispersion Parameters**

In PKIN, please start a new workspace and load the measurement with the menu item **Load Whole Blood Activity**. On the Blood tab, switch the **Whole Blood** model to **Calibration: Step-up** as illustrated below.
Mask all samples except for a baseline, rising edge and the following plateau as follows:

1. Right click into the curve area. In the context menu select **Switch OFF all Points in visible area**. Consequently all samples appear gray, meaning they are not considered in fitting operations.

2. Open the context menu again and select **Use restricted value range**. Enter the start/end times of the relevant section and close with **Ok**.

3. Open the context menu again and select **Switch ON all Points in visible area**.

Next comes the fitting of the step-up model. Enter the time where the signal rises as a starting value for the $T$ parameter, the known concentration value as the $C$ parameter, and enable the fitting flags as illustrated below.
With **Fit Whole Blood** the model curve can be fitted to the section.

A similar fitting can be done with the decreasing edge and the model **Correction: Step-down**. Fitting the different parts of the calibration experiment provides several sets of the dispersion parameters, which allows determining their average values. Those values of the \( \alpha \) and \( k \) parameters can be applied for the correction of the actual blood curve of live experiments.

**Recommendations by Methodology Author, Ole Munk**

\( \alpha \) has an impact on the noise in the dispersion-corrected input functions. If \( \alpha \) is allowed to get close to 1, then the noise increases (too) much (Fig.8 in Munk et al.). Therefore, in practice, \( \alpha \) can be fixed at a reasonable value, eg. 0.75, and only \( k \) fitted along with the two "known" parameters \( T \) and \( C \).

**Notes:**

1. The parameters can be tracer-dependent. It takes less than an hour to do the calibration measurement and the parameters can be re-used for all future measurements using the same blood sampling setup.

2. Dispersion correction is only important when modeling fast processes such as blood flow. When estimating binding potentials and other "slow processes", dispersion correction is considered less important.

**4.9.6.3 O. Munk Dispersion Correction in PKIN**

**Correction of Whole-Blood Curve**

In PKIN, please start a new workspace and load the measurement with the menu item **Load Whole Blood Activity**. On the **Blood** tab, switch the **Whole Blood** model to **Calibration: Correction** as illustrated below.
Next, enter the values for alpha and \( k \) determined with the calibration experiment. The dispersion corrected blood activity is shown as the Whole blood model curve. The Delay parameter represents the delay of the corrected curve relative to the blood arrival at the target tissue, as usual.

From now on, the corrected whole blood activity will be used in calculations such as spillover correction, or calculation of the plasma activity using a plasma fraction function.

**Fitting of Smooth Function to Corrected Curve**

In case the correction introduced substantial noise, a smooth function may be fitted by applying one of the usual whole blood models as follows:

1. **Replace activity data**: Use this button to replace the original (uncorrected) data by the corrected curve.
2. Select an appropriate Whole Blood model, in the example below Gamma + 2 Exponentials.
3. Use Fit whole blood to adjust the model to the dispersion corrected whole blood curve.

### 4.9.6.4 Exponential Deconvolution

In literature, dispersion has often been modeled as convolution of the temporal activity concentration with an exponential kernel

\[
d(t) = \frac{1}{\tau} e^{-t/\tau}
\]

whereby the parameter \( \tau \) is called Dispersion (eg. Meyer et al. [1]).

PKIN offers a facility to perform a numerical deconvolution by such a kernel. The feature can be used by loading the data with Load Whole Blood Activity and selecting Deconvolve whole blood as illustrated below.
A dialog window appears as illustrated below. Note that the **blood model** curve will be deconvolved, not the blood measurement. The reason is that the deconvolution process amplifies noise in the data, so it is recommended first fitting a smooth function to the data.

**Dispersion** is the main parameter, which has to be known and entered. The deconvolved curve can be shifted left by a **Delay**. The other parameters are related to the deconvolution process which performs the following operations when **Deconvolve** is activated:

1. Regular resampling of the blood model function with the specified **Resampling interval**. The signal is extended past the measurement using the model function to reduce the cutoff effect (if the function is decreasing). **Extend past end x** is a factor which is multiplied with the acquisition duration.

2. Calculation of the dispersion kernel for the same sampling times.

3. Forward Fourier transform of the model and the exponential after some tapering and applying a frequency cutoff corresponding to the **Smooth** parameter and a tapering Hanning filter.

4. Divide the transformed model by the transformed exponential.

5. Inverse Fourier transform of the result, and resampling the real part of the result.

6. Convolve the **Result: Deconvolved blood** with the exponential kernel to get **Confirmation: Result convolved with exp**.

Ideally, **Result** and **Confirmation** would be identical. In practice, this will never be the case due to the numerical procedures. The result depends on the curve shape, the noise, and the **Smooth** and **Dispersion** parameters. The user will have to find parameter combination which provides a reasonable result. If this is possible, the result can be used for further processing with **Replace original input**.
Reference

4.10 Convenience Tools

In the lower right part of PKIN there is an ensemble of user interface elements supporting some auxiliary tasks.

4.10.1 Changing Display Types

The Display type selection allows choosing which information is shown in the main curve area.

- **Standard** shows all curves related to the current region.
- **Blood** shows only the blood activity curves.
- **TACs** the measured tissue data from all regions.
- **Targets** all regional target curves to be fitted (in most cases identical to the TACs).
- **Models** all fitted model curves.
- **Residuals** all of the regional residuals.
4.10.2 Copy Operations

This multi-function button can be used to propagate a model from the current region to all regions in the study.

- **Model & Par.** copies the model with the current parameters (which can then be used as starting values for Fit model, all regions).
- **Model** copies the model structure to all regions. If the Conversion is enabled, it performs a parameter conversion operation to maintain $v_B$, $t^*$ and $k_2$.
- Otherwise, the current default parameters of the model will be established.
- **Weighting** copies the current definition how the residuals are weighted to all regions.
- **Valid** copies the definition of the currently valid points to all regions.

### Copy to Blood Curves

This multi-function button allows transferring activity curves as follows:

- **Region to Plasma** copies the current region TAC to the input curve.
- **Region to Whole Blood** copies the current region TAC to the whole blood curve.
- **Plasma to Whole Blood** copies the plasma curve to the whole blood curve.
- **Whole Blood to Plasma** copies the whole blood curve to the plasma curve.

4.10.3 Auxiliary Tools

The **Tools** multi-function button offers the following functionality.

**Edit Data**

**Edit Data** allows editing/processing the data values and generating new TACs, for instance by averaging TACs or adding noise, see below.

**Edit Patient**

**Edit Patient** allows changing the demographic patient data as well as the radionuclide, see below.

is particularly helpful, if the PKIN tool is opened as an intermediate dialog and hence has no menu available. In this case, **Save KM, Save Par, Append Par** just allow to save the data as a .km file, as a parameter summary text file, and to append to a latter file.
4.10.4 Parameters Aggregation

The Kinetic menu contains an entry Aggregate Parameters. Its purpose is to concentrate the parameter values which have been saved in KM Parameters Files (*.kinPar) for statistical analysis. The general concept and the details of data aggregation are explained in the PMOD Base Functionality Guide.

Component Selection

Aggregate Parameters opens a dialog window with two tabs as illustrated below. The Select components pane allows defining the .kinPar files from which the parameters are extracted.

The files may be available in a database as in the example above, or in a disk directory. In the latter case use Load from File System to define the directory were the files reside. All appropriate .kinPar data sets are listed. Select all data sets to be used for the aggregation and then switch to the Set Criteria pane.

Content Selection

The upper part Reorder Selected Components lists the data sets which have been selected for aggregation. They can be reordered if needed.
The Parameters section shows the information contained in the .kinPar files. The left part lists the available VOIs, the right part the kinetic model parameters. A yes label in the Common columns indicates that all of the kinPar files include the particular element. Otherwise, a no label is shown. Note that entries which are not available in a data set will result in a NaN value in the aggregate.

Data Aggregation
Select the VOIs and parameters to be included so that they are highlighted, and activate Aggregate and Save. Specify the name of a database table, or a file name if the database is not used. When opening the saved aggregate with the View aggregated button, the statistics window appears with a selection for the Statistics set.

Aggregated parameters can also be opened again using the View Aggregated button from the taskbar to the right.
Application Cases

A typical example is the aggregation of the results of a test study in a test aggregate, and the results of the same population in a retest study in a retest aggregate. The two aggregates can then directly be used for a test-retest analysis using the PMOD R console.

Another example is the comparison of two analysis methods. In this case the results of the different methods would be aggregated in corresponding aggregates, which can then be compared with a Bland-Altman analysis.

4.11 Editing Facilities

4.11.1 Disable Measurements for Removing Outliers

Occasionally there might be the need to disregard sample points when evaluating the matching criterion, for example to exclude an outlier.

In PKIN this can be easily achieved by (temporarily) disabling samples. First select the Measured curve by selecting it in the controls area or by CTRL+Click at a sample point. The green measurement points must appear highlighted in the display. Then individual points can be disabled by SHIFT+Click on them.

For entire ranges it is advised to zoom into the region (drag left mouse button), then activate the context menu (right mouse button) and Switch OFF all Points in visible area. This operation is illustrated below.
resulting in a "shortened" acquisition for fitting purposes:

Disabled measurements appear in grey. They can again be enabled by a second **SHIFT+Click**, or by **Switch ON all Points in visible area** in the context menu.

Note that there is a dedicated functionality for assessing Parameter Sensitivity to Acquisition Duration.

### 4.11.2 Edit Data and Create New Curves

The **Edit Data** button in the **Tools** list brings up a dialog window which allows the user to examine and manipulate the TAC data of the current data set.
The upper section lists the different curves. The blood-related curves are at the top, then come the regional tissue TACs. The arrows to the right can be used to change the TAC ordering, and a TAC can be removed by the x button.

The lower section lists the data of the selected curve. Shown for each sample are the acquisition start and end times, the measurement value (usually the VOI average), its standard deviation (the VOI stdv, if available), and the VOI volume. Note that the volume may vary if the VOI definition changes throughout the time course. The values of the tissue TACs can be interactively edited by clicking into a cell and changing the value.

The configuration button between the lists houses the following operations to modify TACs and generate new TACs, which will be appended to the list.

<table>
<thead>
<tr>
<th>Average Weighted by Volume</th>
<th>When several curves are selected, this button allows calculating the volume-weighted average TAC.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average</td>
<td>TAC averaging with equal weights.</td>
</tr>
<tr>
<td>Add noise</td>
<td>Creates a copy of the selected TACs and adds noise according to the definition defined in a dialog window.</td>
</tr>
<tr>
<td>Basic operations</td>
<td>Applies simple arithmetic operations to the values of the currently selected curves, and allows multiplying a curve by the values of an exponential function.</td>
</tr>
<tr>
<td>Curve by Curve operations</td>
<td>Arithmetic operations between the values of two selected TACs.</td>
</tr>
</tbody>
</table>
### PKIN Data Processing

**Rename**
For renaming of the currently selected curve.

**Edit volume**
Change the volume of the selected TAC.

**Trim times**
Check for acquisition time overlaps of the selected TACs and correct.

**Correct decay**
Apply a decay correction to the selected TACs using the isotope half-life $t_{1/2}$.

**Uncorrect decay**
Undo the effect of decay correction for the selected TACs using the isotope half-life $t_{1/2}$.

**Edit curve**
Edit the data of the selected curve in a text editor as described below.

**Append TAC from file**
Opens a facility for appending regional TACs stored in a file. After selecting the file with **Load TAC**, the regions are listed so that the user may select a subset. With **Ok**, the regional curves are appended to the existing ones.

![Append TAC from file](image)

**Dosimetry IDAC Export**
This entry is applicable only for the **Cumulated Activity (OLINDA, IDAC)** dosimetry model. It creates an input file to the IDAC dosimetry program.

**Dosimetry OLINDA/EXM Export**
This entry is applicable only for the **Cumulated Activity (OLINDA, IDAC)** dosimetry model. It creates an input file to the OLINDA/EXM dosimetry program.

**Dosimetry OLINDA 2 Export**
This entry is applicable only for the **Cumulated Activity (OLINDA, IDAC)** dosimetry model. It creates an input file to the OLINDA 2 dosimetry program.

### Edit Curve or Add New Curve

The **Edit Curve** tool brings up a dialog showing the data of the selected curve. The structure depends on the curve type selected, blood or tissue.
The data can directly be edited in this simple text editor, exported with Copy to Clipboard, edited e.g. in Excel, and brought back by Paste from Clipboard. Closing with Ok will replace the original curve data. If Append as new curve is checked, the user can change the curve name, and the curve will be appended to the list.

### 4.11.3 Edit Subject/Study Info and Change Isotope

The Edit Subject button in the Tools list brings up a dialog window which allows you to examine and change the demographic information of the current data set.

In case the TAC data was transferred from PVIEW to PKIN, this information is automatically filled in from the image data (if available in the data format). However, the contents can also be edited and saved if required. If the subject is already in the database, the IMPORT SUBJECT INFO
FROM facility can be used. With the DATABASE format a selection window opens which allows choosing an image series from the database. The required information will be retrieved and filled into the text fields.

Note the SUV PARAMETERS tab which gives access to the dose and isotope definition.

![SUV Parameters Tab](image)

This information is relevant for some models as well as calculated residual weighting\(^\text{[58]}\).

Note the BIDS PARAMETERS tab represents an upcoming format which is not yet fully specified and usable.

### 4.11.4 Comments

As a data set is processed, intermediate results build up in the Model Fitting History\(^\text{[57]}\) which will be saved with the data. In order to keep track of the efforts spent and insights gained, comments can be added to the data via the Comments panel. Note that the upper part lists the comment titles, while the lower part shows the contents of the selected comment.

![Comment Panel](image)

**Comment Creation**

Start by the Add button. A new entry is created in the comment list. The greyed Edit mode indicates that now the comment title and the body can be edited. After entering the information, the comment can be finalized with the Save button.
Comment Editing

To change an existing comment select it in the list and activate View mode. Consequently it changes to Edit mode. Now change the title and body text as needed, then activate Save to store the changes and return to the viewing mode.

Note that you have to use Save KM File to include updated comments permanently into the data set.

4.12 Synthetic Data Generation

The compartment models in PKIN can be employed for creating data sets representing a perfect PET measurement with a certain acquisition protocol of the tissue and blood activities.

4.12.1 Synthetic Modeling Study for PKIN

PKIN always shows the model curve given the loaded input curve(s) and the configured regional kinetic model with the timing of the current study. The user may modify any of the defining parameters and observe the effect on the model curve, which represents the expected PET measurement. With the Create Synthetic KM Study facility well-defined and smooth TACs and blood curves can be generated which represent an ideal acquisition.

Synthetic Modeling Study Generation
1. In each region select a compartment model and define an appropriate set of parameter values.

2. For all blood components, configure a representative model function. This is most easily achieved by fitting one of the available smooth functions to actual data.
3. Select the entry **Synthetic KM Study** in the **Kinetic** menu. A dialog appears showing the blood models and the model curve of the current region.

4. Initially all curves have the original sampling times. To modify sampling, use the buttons **Set Blood Sampling** and **Set Tac Sampling**. For instance, very fine sampling allows to generate smooth curves for publication or simulation purposes.

5. When **Save Synthetic Study** is activated, all regional TACs are recalculated with the current sampling, the old measurements are replaced by the values of the model curve, the original blood values are replaced by the values calculated with the current blood models, and the data is saved in a file specified by the user. Then, the original data are shown again. As a convenience for adding the generated data set there is the button **Save & Append as new Study** available underneath:

6. Finally, to see and work with the synthetic study, load the created file. The synthetic model curves are now the measured TACs, and the resampled blood curves have the **Lin. Interpolation** function associated. So any underlying model of the blood curves (which have been used for synthetic model calculation) get lost. This may cause small deviations between the calculated model curve and the saved synthetic TAC, depending on the density of blood sampling.

Note that after loading well-defined noise could be added to the synthetic curves using the **Edit Data** facility as illustrated below. Select the curves to be modified, activate the **Add noise** operation, configure the noise, and quit with **Ok**. The created curves are then appended to the existing data and can be saved.
4.12.2 Synthetic Imaging Study for PXMOD Tests

Create Synthetic PXMOD Study is an extension of the Create Synthetic KM Study approach. The idea is that synthetic model curves can be generated for many different "tissues", i.e. combinations of model parameters, at once. These curves are compiled into images, so that a synthetic study is generated which has a different combination of model parameters in each pixel.

For instance, TACs can be simulated by loading a .km file, selecting the 2-tissue compartment model and using the parameter ranges:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>From</th>
<th>To</th>
</tr>
</thead>
<tbody>
<tr>
<td>$K_1$</td>
<td>0.05</td>
<td>0.1</td>
</tr>
<tr>
<td>$k_2$</td>
<td>0.05</td>
<td>0.25</td>
</tr>
<tr>
<td>$k_3$</td>
<td>0.02</td>
<td>0.04</td>
</tr>
<tr>
<td>$k_4$</td>
<td>fixed to 0.05</td>
<td></td>
</tr>
</tbody>
</table>

To enter these parameter ranges select Create Synthetic PXMOD Study from the File menu. The following dialog appears showing the parameters of the current model.
To define the synthetic study the following configurations must be set:

- Which parameters are varied along the X, Y, and Z axis (Axis selection),
- the value ranges of the varied parameters (Start from, Finish at),
- the number of parameter increments to cover the ranges (X, Y, Z samples),
- the acquisition times; initially the times of the loaded .km study are used, but they can be changed using the Set time points.
- Add noise allows defining a noise generator for disturbing the ideal curves. In the example above, if 10x10x10 curves with the same parameter combination are created, the distribution of the estimated parameters can be analyzed to study their variance in the presence of noise.

**Synthetic Data**

With Generate the model curves of all parameter combinations are generated and the data arranged as a 4-dimensional study. This synthetic study, which is suitable for pixel-wise modeling, is shown in the upper image display. In the example above the interpolation has been disabled to show the 10x10 layout of parameter combinations in plane. Note that the data inspector can be used for inspecting the curve in each pixel of the synthetic data set.
Use the **Save Synthetic Results** button to save the study in any supported output format.

**Gold Standard**

The lower image display contains images of the true parameter used in each pixel. These generation parameters can be saved using the **Save True Results** button. They can serve as the reference for the analysis of the outcome from a pixel-wise analysis of the synthetic study.

### 4.13 Modeling in Batch Pipeline

The PKIN models can be applied as part of a processing pipeline. For instance, a dynamic PET series can be motion corrected, an atlas fitted to the data and the regional TACs calculated which are then fitted by a model. Please refer to the **Pipeline Processing** section in the *PMOD Base Functionality User Guide* for a detailed example.

**Data Preparation by the Association**

For blood-based models blood information related to the image data (input curve, whole blood curve, parent) must be specified. This requires that the data resides in a PMOD database and uses the "association" concept which allows defining relationships between data elements.

For instance, the whole-blood curve is associated as follows in the **PVIEW** tool. In the **Subjects** list of the database browser select the subject, and in the **Series** list the dynamic PET. Next, activate the association list as illustrated below and select the **Associate Whole blood** entry.
A component browser appears which is used to select the whole-blood curve and Set selected.

The same procedure is repeated for the plasma curve with the Associate Plasma entry, and if applicable with the Associate Parent fraction. From then on the roles of the blood curves is established, and selection of the dynamic PET is sufficient for modeling purposes.
5  **PKIN for Regional Parametric Mapping (option)**

PXMOD is the main PMOD tool for parametric mapping and provides a streamlined workflow for the data analysis. However, the modeling incorporated in PXMOD features a limited set of models and is not aimed at detailed interactive modeling of TACs from individual pixels. PKIN offers an alternative for generating parametric maps of limited image sub-volumes, leveraging interactive modeling with the full model set as described below. For instance, the heterogeneity of a tumor might be investigated using the PKIN-based parametric mapping approach.

**Note:** The use of this functionality requires that the PXMOD Tool has been licensed in addition to PKIN.

5.1  **Data Transfer for Parametric Mapping in PKIN**

In order to use parametric mapping in PKIN, the dynamic data have to be loaded in the PVIEW tool, VOIs outlined, and then the transfer to PKIN started as illustrated below. As PKIN can only handle a limited number of TACs, the default **Maximal number of TACs** is set to 10000, but can be changed.

Please select only the relevant VOIs in the list to the left. The VOI average TACs are shown in the curve area, with the number of included voxels indicated in brackets in the list to the right. Activate **TAC of each VOI voxel**, and then transfer the data with **Send TACS to PKIN [New workspace]** to create a new PKIN workspace, or **Send TACS to PKIN [Current workspace]** to use the selected PKIN workspace, in case PKIN is already running.

If no blood information is available for modeling, an average TAC from a reference VOI will be needed. It is recommended transferring it separately as an average TAC by calling the transfer window again, this time only selecting the reference VOI, enabling **Averaged VOI TACs**, enabling **Append TACs to region list in PKIN** and finally activating **Send TACS to PKIN [Current workspace]**.

**Important:** The PVIEW tool needs to remain open with the images loaded during parametric mapping in PKIN.
5.2 Voxel-wise TAC modeling in PKIN

After the transfer from the PVIEW tool the voxel-wise TACs are arranged in the Region list labeled with the VOI name followed by the voxel coordinate in the image. As for average TACs, the blood and plasma data have to be loaded, or a reference approach has to be used.

The whole PKIN functionality is then available for modeling. For instance, a few TACs can be averaged, a 1-tissue compartment model fitted as illustrated, propagated to all TACs using the Copy to All Regions entry Model & Par, and fitted with Fit model, all regions. Another possibility is to transfer the averaged TACs also from PVIEW, use them for setting up the proper model, propagate and fit.

5.3 Parametric Mapping Interface

After all pixelwise TACs have been fitted, and the average TACs removed, select the entry Create Parametric Maps in the Kinetic menu in order to generate parametric maps from model parameters. Note that this entry is only active when a voxel-wise TAC transfer from PVIEW has been performed.
The dialog window offers three methods on the **Setup** panel.

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Create maps from PKIN model fits</strong></td>
<td>This method uses the results of the existing model fits in the PKIN workspace to generate the parametric maps.</td>
</tr>
<tr>
<td><strong>Fit PKIN TACs to create maps</strong></td>
<td>This method is the same as above, but first a model fit is performed in all regions using the model configured in the selected region.</td>
</tr>
<tr>
<td><strong>Fit original image TACs to create map</strong></td>
<td>This method is independent from the TAC contents in PKIN. It will use the model configured for the active region and fit it to the TACs of all voxels within the selected regions if <strong>Use only selected regions</strong> is enabled, and in the whole image volume otherwise. Note that the time required depends on the number of voxels to be fitted.</td>
</tr>
<tr>
<td><strong>Additional parameters maps: Macro</strong></td>
<td>Supports the creation of parametric maps from all the macro parameters of the model.</td>
</tr>
<tr>
<td><strong>Additional measures maps: Quality</strong></td>
<td>Supports the creation of parametric maps from all quality measures of the fit.</td>
</tr>
</tbody>
</table>

The **Generate** button starts processing with the selected method, resulting in images on the tabs **Parametric Maps**, **Error Maps** and **Model Curve Image**. Note the synchronization between the images on the various panels and the **Region** selected in the PKIN interface as described above.

**Note:** If average TACs were transferred, **Create Parametric Maps** can also be executed. All the settings described are applicable, but the parametric maps will consist of a single value in each region, the regional result.
5.3.1 Original Image Panel

The Original Image tab shows the image that was used for transferring the voxel-wise TACs, together with the VOIs. Besides providing reference for existing maps, it supports an interactive mode of fitting the TACs in voxels of interest.

Once the Transfer and fit pixel TACs option has been enabled, the following happens when clicking into the image:

- The TAC of the voxel is transferred to PKIN and a new entry in the Region list is created,
- the model of the previously active region is fitted to the TAC,
- and the resulting parameters are added in the image on the Parametric Maps panel.

The example below shows the Vt map after clicking at 3 voxels in the frontal area.

5.3.2 Parametric Maps Panel

The Parametric Maps panel shows the results from the voxel-wise fits as parametric images, with the selection between the different model parameters in the upper right.
Save Parametric Maps allows saving the currently selected map, or All at once. Show fusion of selected parameter opens a dialog window showing the parametric map on top of the dynamic images.

Fitting Inspection
If the parametric map has been produced from curves in the PKIN interface, the fit in the individual voxels can be inspected in detail. When clicking at voxels in a parametric map the Region in the main PKIN interface is switched to the corresponding entry, and the curve panel shows the TAC of the voxel with the model curve.
5.3.3 SE Maps Panel

The **Error Maps** panel shows the % standard error of the corresponding parameters on the **Parametric Maps** panel.

5.3.4 Model based Image Panel

The last panel contains a dynamic image series which is constructed from the fitted model curves. It can be saved via **Save Model Image** for other uses such as constructing digital phantoms or as input into other parametric mapping programs.
6 PKIN Reference

PKIN implements different types of models with somewhat different properties.

1. Compartment Models require an input curve for the calculation of the expected concentrations in the different compartments. Most of them also support a blood spillover correction term.

2. Reference models do not use explicit blood information. Rather, a reference TAC is specified which must satisfy specific criteria. The reference TAC is used to calculate the response in the tissue of interest.

3. Non-compartment models perform different types of analyses which are not based on a compartment structure of the model. Most of them use the input curve.

Note: It is assumed that all data used has been decay corrected to the same time point.

6.1 Compartment Models

6.1.1 List of PKIN Compartment Models

PKIN features a comprehensive set of compartment models as listed below. The number of tissue compartments ranges from 1 to 3. There are different variants of the same model structure, so that prior information can be entered easily, and that coupling of physiologic parameters across regions is possible.

For instance, the 2-tissue compartment model has the standard parameters $K_1, k_2, k_3, k_4$. An equivalent description is by the parameters $K_1, K_1/k_2, k_3, k_4$. The advantage is, that the $K_1/k_2$, represents the distribution volume of the non-displacement compartment in tissue (free and non-specifically bound tracer), which can often be assumed to be the same across different tissues. Therefore, $K_1/k_2$ can be included as a common parameter in a coupled fit, hereby reducing the number of fitted parameters and thus potentially improving the identifiability of all.

<table>
<thead>
<tr>
<th>Model Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-Tissue Compartment</td>
<td>Most basic compartment model with the plasma compartment and one tissue compartment.</td>
</tr>
<tr>
<td>2-Tissue Compartments</td>
<td>Compartment model with the plasma compartment and two sequential tissue compartments. Often used for receptor studies.</td>
</tr>
<tr>
<td>FDG-2 Tissue Compartments</td>
<td>The 2-Tissue compartment model including the plasma glucose and the lumped constant to calculate the metabolic rate of glucose.</td>
</tr>
<tr>
<td>2-Tissue Compartments, $K_1/k_2$</td>
<td>The same model as the 2-Tissue compartment model, except that $K_1/k_2$ is used as a model parameter instead of $k_2$. This facilitates coupled fitting.</td>
</tr>
<tr>
<td>2-Tissue Compartments, $K_1/k_2$ &amp; $V_s$</td>
<td>The same models as the 2-Tissue compartment model, except that $K_1/k_2$ and the specific or total distribution volumes are used as model parameters instead of $k_2$ and $k_4$. This facilitates coupled fitting and the easy generation of synthetic model curves.</td>
</tr>
<tr>
<td>2-Tissue compartments, $V_s$ Linear Least Squares</td>
<td>2-Tissue compartment model solved by the Linear Least Squares method.</td>
</tr>
<tr>
<td>Model Type</td>
<td>Description</td>
</tr>
<tr>
<td>------------</td>
<td>-------------</td>
</tr>
<tr>
<td>Parallel 2-Tissue Compartments</td>
<td>2-Tissue compartment model with each compartment exchanging independently with blood.</td>
</tr>
<tr>
<td>2-Tissue Compartments, k5</td>
<td>2-Tissue compartment model with an efflux from the last compartment.</td>
</tr>
<tr>
<td>2-Tissue Compartments, Bmax</td>
<td>Non-linear 2-Tissue compartment model for receptor tracer studies accounting for the saturation of receptor sites.</td>
</tr>
<tr>
<td>2-Tissue Compartments, K1/k2, Bmax</td>
<td>Same model as above, but using K1/k2 as fitting parameter instead of k2.</td>
</tr>
<tr>
<td>2-Tissue Compartments, vascular trapping</td>
<td>2-Tissue compartment model with an additional endothelial trapping compartment.</td>
</tr>
<tr>
<td>Irreversible 2-Tissue</td>
<td>In principle the same models as the 2-Tissue compartment model with k4 = 0. The Flux variants allow using the influx as a fitting parameter, or as a fixed parameter set to the result of a Patlak analysis. When using K1/k2 also as a fitting parameter, coupled fitting can be employed.</td>
</tr>
<tr>
<td>3-Tissue Compartments</td>
<td>Compartment model which separates free tracer in tissue from non-specific binding.</td>
</tr>
<tr>
<td>3-Tissue Compartments, K1/k2, Vs</td>
<td>As above, but using but using K1/k2 and the specific distribution volume as fitting parameters.</td>
</tr>
<tr>
<td>3-Tissue Compartments, sequential</td>
<td>Model with three sequential compartments originally developed for FDG uptake in skeletal muscle.</td>
</tr>
<tr>
<td>Flow &amp; Dispersion</td>
<td>Specific model for dynamic H215O-PET Data with implicit deconvolution of the input curve dispersion.</td>
</tr>
<tr>
<td>3-Tissue Compartments, Metabolites</td>
<td>Extends the 2 tissue compartment model by a metabolite compartment with a second input curve of labeled metabolites from the plasma.</td>
</tr>
<tr>
<td>3-Tissue Compartments, Metabolites, K1/k2</td>
<td>As above, but using but using K1/k2 and the specific distribution volume as fitting parameters.</td>
</tr>
<tr>
<td>4-Tissue Compartments, Metabolites, K1/k2</td>
<td>Two parallel 2-compartment models for authentic ligand and metabolites, linked by a transfer constant between the non-specific compartments.</td>
</tr>
<tr>
<td>Triple-injection Protocol for Flumazenil</td>
<td>During a single imaging study three injections are applied: hot ligand first, then cold ligand for displacement, then a mixture of cold &amp; hot ligand. The individual receptor parameters can be estimated.</td>
</tr>
</tbody>
</table>
6.1.2 1-Tissue Compartment Model

The 1-Tissue Compartment Model only separates tissue uptake and washout. All tracer in tissue is included in a single compartment \( C_1 \).

### Differential Equation of the Mass Balance

\[
\frac{dC_1(t)}{dt} = K_1 C_p(t) - k_2 C_1(t)
\]

with input curve \( C_p(t) \), tissue concentration \( C_1(t) \), uptake rate constant \( K_1 [\text{ml/ccm/min}] \) and clearance rate constant \( k_2 [1/\text{min}] \).

### Operational Model Curve

\[
C_{\text{Model}}(t) = (1 - v_B) C_1(t) + v_B C_B(t)
\]

with the concentration \( C_B(t) \) of tracer in whole blood, and the blood volume fraction \( v_B \).

### Parameter Fitting

The model includes the three fitable parameters \( v_B, K_1, k_2 \). Often, \( v_B \) is fixed at a physiologic value of 3-5%.

6.1.3 2-Tissue Compartment Model

The 2-tissue compartment model separates tracer in tissue into two exchanging compartments \( C_1 \) and \( C_2 \) with a serial structure. The arterial plasma exchanges with the first tissue compartment \( C_1 \), which in turn exchanges with the second tissue compartment \( C_2 \). \( K_1 [\text{ml/ccm/min}] \) and \( k_2 [1/\text{min}] \) are the uptake and clearance rate constants, whereas \( k_3 [1/\text{min}] \) and \( k_4 [1/\text{min}] \) describe the exchange between the tissue compartments. The usual interpretation is that \( C_1 \) represents free and non-specifically bound tracer in tissue (non-displaceable compartment), and \( C_2 \) specifically bound tracer.

### Differential Equation of the Mass Balance

\[
\begin{align*}
\frac{dC_1(t)}{dt} &= K_1 C_p(t) - (k_2 + k_3) C_1(t) + k_4 C_2(t) \\
\frac{dC_2(t)}{dt} &= k_3 C_1(t) - k_4 C_2(t)
\end{align*}
\]

with input curve \( C_p(t) \).

### Operational Model Curve

\[
C_{\text{Model}}(t) = (1 - v_B) (C_1(t) + C_2(t)) + v_B C_B(t)
\]

with the concentration \( C_B(t) \) of tracer in whole blood, and the blood volume fraction \( v_B \).
Parameter Fitting
The model includes the 5 fitable parameters $v_B$, $K_1$, $k_2$, $k_3$, $k_4$. Usually, $v_B$ is fixed at a physiologic value of 3-5% to reduce the number of fitted parameters. An alternative is to fit $v_B$ with a TAC from a big representative VOI, and fix it for the TACs from smaller VOIs. For irreversible tracers with trapping in the second compartment, $k_4$ is fixed: $k_4=0$. Please inspect the %SE standard error to get information about the reliability of the parameter estimates.

Alternative Models with VND and VS Fit Parameters
There are two alternative models available which have the same structure but use distribution volumes as fit parameters. The advantage of the reformulation is that in some situations distribution volumes can be assumed to be common among regions and estimated by a coupled fit, or they might already be known and can thus be fixed at a prior value.

6.1.4 2-Tissue Compartment Model, $K_1/k_2$
The 2-tissue compartment model separates tracer in tissue into two exchanging compartments $C_1$ and $C_2$ with a serial structure. The arterial plasma exchanges with the first tissue compartment $C_1$, which in turn exchanges with the second tissue compartment $C_2$. $K_1$ [ml/ccm/min] and $k_2$ [1/min] are the uptake and clearance rate constants, whereas $k_3$ [1/min] and $k_4$ [1/min] describe the exchange between the tissue compartments. The usual interpretation is that $C_1$ represents free and non-specifically bound tracer in tissue (non-displaceable compartment), and $C_2$ specifically bound tracer.

Differential Equation of the Mass Balance
\[
\frac{dC_1(t)}{dt} = K_1 C_p(t) - (k_2 + k_3) C_1(t) + k_4 C_2(t)
\]
\[
\frac{dC_2(t)}{dt} = k_3 C_1(t) - k_4 C_2(t)
\]
with input curve $C_p(t)$.

Operational Model Curve
\[
C_{model}(t) = (1 - v_B) (C_1(t) + C_2(t)) + v_B C_B(t)
\]
with the concentration $C_B(t)$ of tracer in whole blood, and the blood volume fraction $v_B$.

Parameter Fitting using $K_1/k_2$ instead of $k_2$
The 2-Tissue Compartment Model, $K_1/k_2$ model variant uses $K_1/k_2$ as a fitting parameter instead of $k_2$. $K_1/k_2$ represents the distribution volume $V_{ND}$ of the non-displaceable compartment $C_1$. The advantage of this formulation is that in some situations $V_{ND}$ can be assumed to be common among regions and estimated by a coupled fit. Alternatively, $V_{ND}$ might already be known and can thus be fixed at a prior value. The other four fitting parameters $v_B$, $K_1$, $k_3$, $k_4$ are the same as for the 2-Tissue Compartment Model.

6.1.5 2-Tissue Compartment Model, $K_1/k_2$, $V_s$
The 2-tissue compartment model separates tracer in tissue into two exchanging compartments $C_1$ and $C_2$ with a serial structure. The arterial plasma exchanges with the first tissue compartment $C_1$, which in turn exchanges with the second tissue compartment $C_2$. $K_1$ [ml/ccm/min] and $k_2$ [1/min]
are the uptake and clearance rate constants, whereas \( k_3 \) [1/min] and \( k_4 \) [1/min] describe the exchange between the tissue compartments. The usual interpretation is that \( C_1 \) represents free and non-specifically bound tracer in tissue (non-displaceable compartment), and \( C_2 \) specifically bound tracer.

\[
\begin{align*}
\frac{dC_1(t)}{dt} &= K_1 C_p(t) - (k_2 + k_3) C_1(t) + k_4 C_2(t) \\
\frac{dC_2(t)}{dt} &= k_3 C_1(t) - k_4 C_2(t)
\end{align*}
\]

with input curve \( C_p(t) \).

**Operational Model Curve**

\[
C_{\text{model}}(t) = (1 - vB) (C_1(t) + C_2(t)) + vB C_p(t)
\]

with the concentration \( C_{\text{b}}(t) \) of tracer in whole blood, and the blood volume fraction \( v_B \).

**Parameter Fitting using \( K_1/k_2 \) and \( V_s \)**

This **2-Tissue Compartment Model, K1/k2, Vs** model variant uses \( K_1/k_2 \) and \( V_s \) as fitting parameters instead of \( k_2 \) and \( k_4 \). \( V_s (=K_1/k_2*V_2/k_4) \) represents the distribution volume of the specific compartment \( C_2 \). This formulation has particular advantages for the generation of synthetic data with predefined distribution volumes. The other three fitting parameters \( v_B, K_1, \) and \( k_3 \) are the same as for the **2-Tissue Compartment Model**.

### 6.1.6 2-Tissue Compartment Model, K1/k2, Vt

The 2-tissue compartment model separates tracer in tissue into two exchanging compartments \( C_1 \) and \( C_2 \) with a serial structure. The arterial plasma exchanges with the first tissue compartment \( C_1 \), which in turn exchanges with the second tissue compartment \( C_2 \). \( K_1 \) [ml/ccm/min] and \( k_2 \) [1/min] are the uptake and clearance rate constants, whereas \( k_3 \) [1/min] and \( k_4 \) [1/min] describe the exchange between the tissue compartments. The usual interpretation is that \( C_1 \) represents free and non-specifically bound tracer in tissue (non-displaceable compartment), and \( C_2 \) specifically bound tracer.

\[
\begin{align*}
\frac{dC_1(t)}{dt} &= K_1 C_p(t) - (k_2 + k_3) C_1(t) + k_4 C_2(t) \\
\frac{dC_2(t)}{dt} &= k_3 C_1(t) - k_4 C_2(t)
\end{align*}
\]

with input curve \( C_p(t) \).

**Operational Model Curve**
Parameter Fitting using $K_1/k_2$ and $V_t$

This 2-Tissue Compartment Model, $K_1/k_2$, $V_t$ model variant uses $K_1/k_2$ and $V_t$ as fitting parameters instead of $k_2$ and $k_4$. $V_t = (K_1/k_2) \times (1 + k_3/k_4)$ represents the total distribution volume of tracer in tissue. This formulation has particular advantages for the generation of synthetic data with predefined distribution volumes. The other three fitting parameters $v_B$, $K_1$, and $k_4$ are the same as for the 2-Tissue Compartment Model.

6.1.7 FDG 2-Tissue Compartment Model

The 2-tissue compartment model for FDG has the same structure as the standard model.

Differential Equation of the Mass Balance

\[
\frac{dC_1(t)}{dt} = K_1 C_p(t) - (k_2 + k_3) C_1(t) + k_4 C_2(t)
\]

with FDG concentration of in arterial plasma $C_p(t)$, FDG concentration $C_1(t)$ and phosphorylated FDG (FDG-6-P) concentration $C_2(t)$ in tissue. $K_1$ [ml/ccm/min] and $k_2$ [1/min] are the uptake and clearance rate constants, whereas $k_3$ [1/min] represents phosphorylation by hexokinase and $k_4$ [1/min] dephosphorylation.

Operational Model Curve

\[
C_{Mode}(t) = (1 - vB) (C_1(t) + C_2(t)) + vB C_2(t)
\]

with the concentration $C_0(t)$ of tracer in whole blood, and the blood volume fraction $v_B$.

Parameter Fitting

The model includes the 5 fitable parameters $v_B$, $K_1$, $k_2$, $k_3$, $k_4$. Usually, $v_B$ is fixated at a physiologic value of 3-5% to reduce the number of fitted parameters, and dephosphorylation is so slow that $k_4$ is fixed $k_4=0$.

FDG-specific Results

The FDG model has two additional input parameters, the lumped constant (LC) and the plasma glucose concentration (PG [mmol/l]). In combination with the estimated $K_1$, $k_2$, and $k_3$ parameters they allow calculating the metabolic rate of glucose MRGlu

\[
MRGlu = \frac{PG \times K_1 k_3}{LC \times k_2 + k_3}
\]

For convenience, the LC and PG values are maintained when switching back and forth with the Patlak model, if the Model conversion option in the Extras panel is enabled.

Note: The unit conversion is based on an assumed tissue density of 1.04g/ml.
### 6.1.8 2-Tissue Compartment Model, Linear Least Squares Method

In addition to uptake and washout the model handles two exchanging compartments $C_1$ and $C_2$ in tissue.

![Diagram of 2-Tissue Compartment Model](image)

**Differential Equation of the Mass Balance**

\[
\frac{dC_1(t)}{dt} = K_1 C_p(t) - (k_2 + k_3) C_1(t) + k_4 C_2(t)
\]

\[
\frac{dC_2(t)}{dt} = k_3 C_1(t) - k_4 C_2(t)
\]

with input curve $C_p(t)$.

**Operational Model Curve**

With linearized solutions, the differential equations are integrated twice on both sides, substitutions performed and finally rearranged. This can be done in different ways. The current model uses the derivation of Cai et al. [1], equation (6):

\[
C_{Model}(t) = P_1 C_B(t) + P_2 \int_0^t C_p(\tau)d\tau + P_3 \int_0^t C_p(s)ds d\tau + P_4 \int_0^t C_{PET}(\tau)d\tau + P_5 \int_0^t C_{PET}(s)ds d\tau
\]

with input curve $C_p(t)$, the concentration $C_B(t)$ of tracer in whole blood, and the tissue curve to be fitted $C_{PET}(t)$.

**Parameter Fitting**

The multi-linear operational equation can be solved in a least squares sense in one step using a singular value decomposition method. From the resulting 5 coefficients $P_1$, $..$, $P_5$ the target parameters can be calculated as follows:

\[
vB = P_1
\]

\[
K_1 = \frac{P_1 P_3 + P_2}{1 - P_1}
\]

\[
k_3 = -(k_2 + k_4 + P_4)
\]

\[
k_4 = \frac{-P_5}{k_2}
\]

\[
k_2 = -\frac{P_1 P_3 + P_3}{P_1 P_4 + P_2}
\]

**Recommended Use of the Linear Least Squares Method**

The advantages of the linearized approach are fast computation, a well-defined solution, and no problem with local minima. However, it is well known that the method is susceptible to bias, and that small TAC perturbations can cause large changes of the parameter estimates. Therefore it is recommended that the **Linear Least Squares** model is only used for getting a quick solution which is further refined by the iterative methods. In fact, if **Parameters Initialization** option is configured in the **Extras** panel, the initial starting parameters of the compartment models are obtained by the **Linear Least Squares** method.

**Implementation Notes**

The model works as follows for different model configurations:

- **Irreversible model**: if $k_4$ is disabled from fitting, the program automatically sets $k_4 = 0$ and uses a linearized equation which is modified accordingly.
• 1-tissue compartment model: if $k_3$ is disabled from fitting, the program automatically sets $k_3 = k_4 = 0$ and uses a modified equation.

• In all configurations the $v_B$ parameter can optionally be disabled from fitting and fixed at a specific user-defined value.

Reference


6.1.9 Parallel 2-Tissue Compartment Model

This model was developed for the analysis of renal PET data [1]. It consists of two parallel tissue compartments, $C_1$ representing non-specific uptake, and $C_2$ specific binding.

Differential Equation of the Mass Balance

\[
\frac{dC_1(t)}{dt} = K_1 C_p(t) - k_2 C_1(t) \\
\frac{dC_2(t)}{dt} = K_3 C_p(t) - k_4 C_2(t)
\]

with input curve $C_p(t)$, concentration $C_1(t)$ in the first and concentration $C_2(t)$ in the second compartment. $K_1$ [ml/ccm/min] and $k_2$ [1/min] are the uptake and clearance rate constants of $C_1(t)$, $K_3$ [ml/ccm/min] and $k_4$ [1/min] of $C_2(t)$ respectively. In the analysis of Gulaldi et al [1] $C_1$ represents non-specifically bound tracer in tissue and $C_2$ represents specifically bound tracer.

Operational Model Curve

\[
C_{\text{Model}}(t) = (1 - v_B) (C_1(t) + C_2(t)) + v_B C_p(t)
\]

with the concentration $C_{\text{B}}(t)$ of tracer in whole blood, and the blood volume fraction $v_B$.

Parameter Fitting

The model includes the 6 fitable parameters $v_B$, $K_1$, $k_2$, $K_3$, $k_4$. Usually, $v_B$ is fixed at a physiologic value to reduce the number of fitted parameters. In [1] $v_B$ was fixed at 0.15, which is used as the parameter default.

Reference

6.1.10 2-Tissue Compartment Model, k5

In addition to uptake and washout the model handles two exchanging compartments \( C_1 \) and \( C_2 \) in tissue, as well as an efflux from \( C_2 \).

**Differential Equation of the Mass Balance**

\[
\frac{dC_1(t)}{dt} = K_1 C_p(t) - (k_2 + k_3) C_1(t) + k_4 C_2(t)
\]

\[
\frac{dC_2(t)}{dt} = k_3 C_1(t) - (k_4 + k_5) C_2(t)
\]

with input curve \( C_p(t) \), concentration \( C_1(t) \) in the first and concentration \( C_2(t) \) in the second compartment. \( K_1 \ [\text{ml/ccm/min}] \) and \( k_2 \ [\text{1/min}] \) are the uptake and clearance rate constants, whereas \( k_3 \ [\text{1/min}] \) and \( k_5 \ [\text{1/min}] \) describe the exchange between the tissue compartments. The typical interpretation is that \( C_1 \) represents free and non-specifically bound tracer in tissue (non-displaceable compartment), and \( C_2 \) represents specifically bound tracer.

**Operational Model Curve**

\[
C_{\text{Model}}(t) = (1 - v_B) (C_1(t) + C_2(t)) + v_B C_B(t)
\]

with the concentration \( C_B(t) \) of tracer in whole blood, and the blood volume fraction \( v_B \).

**Parameter Fitting**

The model includes the 6 fitable parameters \( v_B, K_1, k_2, k_3, k_4, k_5 \). Usually, \( v_B \) is fixed at a physiologic value of 3-5% to reduce the number of fitted parameters. Per default, \( k_4=0 \) is fixed to model an efflux from the specific compartment rather than an exchange with the non-displaceable compartment.

6.1.11 2-Tissue Compartment Model, Bmax

The saturable receptor-ligand model includes two tissue compartments. \( C_1 \) represents the non-displaceable tissue compartment with free and non-specifically bound tracer, and \( C_2 \) tracer bound to the target receptors. The model accounts for a saturation of receptor sites due to low specific activity \( S_{\text{act}} \) of the injected tracer. In this situation, \( k_3 \) is given by \( k_{\text{on}} (B_{\text{max}} - C_2(t)/S_{\text{act}}) \), and is therefore time-dependent. \( B_{\text{max}} \) is the total receptor concentration, \( k_{\text{on}} \) the bi-molecular association rate, and \( k_4 \) the dissociation rate constant \( k_{\text{on}} \).

**Differential Equation of the Mass Balance**

\[
\frac{dC_1(t)}{dt} = K_1 C_p(t) - k_2 C_1(t) - k_{\text{act}} C_1(t) \left( B_{\text{max}} - \frac{C_2(t)}{S_{\text{act}}} \right) + k_4 C_2(t)
\]

\[
\frac{dC_2(t)}{dt} = k_{\text{act}} C_1(t) \left( B_{\text{max}} - \frac{C_2(t)}{S_{\text{act}}} \right) - k_4 C_2(t)
\]

with input curve \( C_p(t) \).
Operational Model Curve

\[ C_{\text{model}}(t) = (1 - v_B) (C_1(t) + C_2(t)) + v_B C_3(t) \]

with the concentration \( C_0(t) \) of tracer in whole blood, and the blood volume fraction \( v_B \).

Parameter Fitting

The model includes the 6 fitable parameters \( v_B, K_1, k_2, k_3\text{on}, k_4, B_{\text{max}} \). Due to the high number of parameters, it is difficult to use the model for actual fitting. However, it may be valuable for generating synthetic data and assessing the effect of model simplification.

Alternative Model with \( K_1/k_2 \) Fit Parameter

There is an alternative 2-Tissue Compartment Model, \( K_1/k_2, B_{\text{max}} \) model available which has the same structure but uses \( K_1/k_2 \) as a fitting parameter instead of \( k_2 \). \( K_1/k_2 \) represents the distribution volume \( V_{\text{ND}} \) of the non-displaceable compartment \( C_1 \). The advantage of this formulation is that in some situations \( V_{\text{ND}} \) can be assumed to be common among regions and estimated by a coupled fit. Alternatively, \( V_{\text{ND}} \) might already be known and can thus be fixed at a prior value.

6.1.12 2-Tissue Compartment Model, \( K_1/k_2, B_{\text{max}} \)

The saturable receptor-ligand model includes two tissue compartments. \( C_1 \) represents the non-displaceable tissue compartment with free and non-specifically bound tracer, and \( C_2 \) tracer bound to the target receptors. The model accounts for a saturation of receptor sites due to low specific activity \( S_{\text{act}} \) of the injected tracer. In this situation, \( k_3 \) is given by \( k_{\text{on}}(B_{\text{max}} - C_2(t)/S_{\text{act}}) \), and is therefore time-dependent. \( B_{\text{max}} \) is the total receptor concentration, \( k_{\text{on}} \) the bi-molecular association rate, and \( k_4 \) the dissociation rate constant \( k_{\text{off}} \).

Differential Equation of the Mass Balance

\[ \frac{dC_1(t)}{dt} = K_1 C_p(t) - k_2 C_1(t) - k_{\text{act}} C_1(t) \left( B_{\text{max}} - \frac{C_2(t)}{S_{\text{act}}} \right) + k_4 C_2(t) \]
\[ \frac{dC_2(t)}{dt} = k_{\text{on}} C_1(t) \left( B_{\text{max}} - \frac{C_2(t)}{S_{\text{act}}} \right) - k_4 C_2(t) \]

with input curve \( C_p(t) \).

Operational Model Curve

\[ C_{\text{model}}(t) = (1 - v_B) (C_1(t) + C_2(t)) + v_B C_3(t) \]

with the concentration \( C_0(t) \) of tracer in whole blood, and the blood volume fraction \( v_B \).

Parameter Fitting

This model alternative 2-Tissue Compartment Model, \( K_1/k_2, B_{\text{max}} \) uses \( K_1/k_2 \) as a fitting parameter instead of \( k_2 \). \( K_1/k_2 \) represents the distribution volume \( V_{\text{ND}} \) of the non-displaceable compartment \( C_1 \). The advantage of this formulation is that in some situations \( V_{\text{ND}} \) can be assumed to be common among regions and estimated by a coupled fit. Alternatively, \( V_{\text{ND}} \) might already be known and can thus be fixed at a prior value. The other 5 fitable parameters \( v_B, K_1, k_3\text{on}, k_4, B_{\text{max}} \) are the same as for the 2-Tissue Compartment Model, \( B_{\text{max}} \) model.
Due to the high number of parameters, it is difficult to use the model for actual fitting. However, it may be valuable for generating synthetic data and assessing the effect of model simplification.

### 6.1.13 2-Tissue Compartment Model with Vascular Trapping

This model has been developed by Rizzo et al. [1] for the analysis of [11C]-PBR28 uptake in the human brain. In addition to the compartments in the standard 2-tissue compartment model, a compartment $C_{\text{vasc}}$ was introduced which represents slow binding of the tracer to the endothelium of the vasculature.

**Differential Equation of the Mass Balance**

\[
\begin{align*}
\frac{dC_1(t)}{dt} &= K_1 C_p(t) - (k_2 + k_3) C_1(t) + k_4 C_2(t) \\
\frac{dC_2(t)}{dt} &= k_3 C_1(t) - k_4 C_2(t) \\
\frac{dC_{\text{vasc}}(t)}{dt} &= K_b C_p(t)
\end{align*}
\]

with input curve $C_p(t)$.

**Operational Model Curve**

\[C_{\text{Model}}(t) = (1 - vB) (C_1(t) + C_2(t)) + vB (C_B(t) + C_{\text{vasc}}(t))\]

with the concentration $C_B(t)$ of tracer in whole blood, and the blood volume fraction $v_B$.

**Parameter Fitting**

The model includes the 6 fitable parameters $vB$, $K_1$, $k_2$, $k_3$, $k_4$, $K_b$. Usually, $vB$ is fixed at a physiologic value of 3-5%. The default initial parameter values are the ones which were used for PBR28.

**Reference**


### 6.1.14 2-Tissue Compartment Model with Vascular Binding

A model with vascular binding has been developed by Rizzo et al. [1] for the analysis of [11C]-PBR28 uptake in the human brain. In addition to the compartments in the standard 2-tissue compartment model, a compartment $C_{\text{vasc}}$ was introduced which represents slow binding of the tracer to the endothelium of the vasculature. This variant allows for reversible endothelial binding. Additionally, $K_1/k_2$ was introduced as a fit parameter, allowing to reduce the number of fit parameters via regional parameter coupling.
### Operational Model Curve

The model includes the 6 fitable parameters $v_B$, $K_1$, $K_1/k_2$, $k_3$, $k_4$, $K_{b1}$ and $k_{b2}$. Usually, $v_B$ is fixed at a physiologic value of 3-5%. Where possible, the default initial parameter values are the ones which were used for PBR28.

### 6.1.15 Irreversible 2-Tissue Compartment Model

The irreversible 2-tissue compartment model separates tracer in tissue into two exchanging compartments $C_1$ and $C_2$ with a serial structure. The arterial plasma exchanges with the first tissue compartment $C_1$, which in turn exchanges with the second tissue compartment $C_2$ where the tracer is trapped. $K_1$ [ml/ccm/min] and $k_2$ [1/min] are the uptake and clearance rate constants, whereas $k_3$ [1/min] describes the trapping. The usual interpretation is that $C_1$ represents free and non-specifically bound tracer in tissue (non-displaceable compartment), and $C_2$ irreversibly bound or metabolized tracer.

An important parameter of an irreversible configuration is net influx rate $K_i$, the unidirectional uptake rate constant that incorporates both net inward transport and trapping of the tracer in tissue.

\[
Net\ \text{influx}\ = \frac{K_1 k_3}{k_2 + k_3}
\]

Another macroparameter "lambda\*k_4," as been used in the context of enzymatic reactions, where $k_3$ is supposed to be proportional to enzyme concentration.
Logan et al. [1] have shown in simulations that $\lambda k_3$ is preferable to $k_3$ alone in terms of bias and precision.

**Differential Equation of the Mass Balance**

\[
\begin{align*}
\frac{dC_1(t)}{dt} &= K_1 C_p(t) - (k_2 + k_3) C_1(t) \\
\frac{dC_2(t)}{dt} &= k_3 C_1(t)
\end{align*}
\]

with input curve $C_p(t)$.

**Operational Model Curve**

\[
C_{\text{Model}}(t) = (1 - vB) (C_1(t) + C_2(t)) + vB C_B(t)
\]

with the concentration $C_B(t)$ of tracer in whole blood, and the blood volume fraction $v_B$.

**Parameter Fitting**

The model includes the 4 fitable parameters $v_B$, $K_1$, $k_2$, $k_3$, and calculates the macroparameter **Flux**. Usually, $v_B$ is fixed at a physiologic value of 3-5% to reduce the number of fitted parameters. An alternative is to fit $v_B$ with a TAC from a big representative VOI, and fix it for the TACs from smaller VOIs.

**Alternative Models with Flux Fit Parameter**

There are two alternative models available which have the same structure but use the **Flux** as a fit parameter. The advantage of the reformulation is that the **Flux** can be calculated with the robust **Patlak** model, and used as a fixed parameter in the compartment model fit.

**Reference**


### 6.1.16 Irreversible 2-Tissue Compartment Model, Flux

The irreversible 2-tissue compartment model separates tracer in tissue into two exchanging compartments $C_1$ and $C_2$ with a serial structure. The arterial plasma exchanges with the first tissue compartment $C_1$, which in turn exchanges with the second tissue compartment $C_2$ where the tracer is trapped. $K_1$ [ml/ccm/min] and $k_2$ [1/min] are the uptake and clearance rate constants, whereas $k_3$ [1/min] describes the trapping. The usual interpretation is that $C_1$ represents free and nonspecifically bound tracer in tissue (non-displaceable compartment), and $C_2$ irreversibly bound or metabolized tracer.

An important parameter of an irreversible configuration is net influx rate $K_i$, the unidirectional uptake rate constant that incorporates both net inward transport and trapping of the tracer in tissue.
Another macroparameter "lambda*k₃," as been used in the context of enzymatic reactions, where k₃ is supposed to be proportional to enzyme concentration.

\[ \lambda k_3 = (K_1 / k_2) k_3 \]

Logan et al. [1] have shown in simulations that lambda*k₃ is preferable to k₃ alone in terms of bias and precision.

Differential Equation of the Mass Balance

\[ \frac{dC_1(t)}{dt} = K_1 C_p(t) - (k_2 + k_3) C_1(t) \]
\[ \frac{dC_2(t)}{dt} = k_3 C_1(t) \]

with input curve C_p(t).

Operational Model Curve

\[ C_{model}(t) = (1 - vB) (C_1(t) + C_2(t)) + vB C_p(t) \]

with the concentration C_p(t) of tracer in whole blood, and the blood volume fraction v_B.

Parameter Fitting

The model includes the 4 fitable parameters v_B, K_1, Flux, k_3, and calculates the LambdaK3 macroparameter. Usually, v_B is fixed at a physiologic value of 3-5% to reduce the number of fitted parameters.

Parameter Fitting using Flux instead of k_2

The Irreversible 2-Tissue Compartment Model, Flux model variant uses Flux as a fitting parameter instead of k_2. The advantage of the reformulation is that the Flux can be calculated as a regression slope with the robust Patlak-n model, and used as a fixed parameter in the compartment model fit. If the Model Conversion option is enabled, the Flux will be copied when switching from the Patlak model to the Irreversible 2-Tissue Compartment Model, Flux model.

Reference


The irreversible 2-tissue compartment model separates tracer in tissue into two exchanging compartments C₁ and C₂ with a serial structure. The arterial plasma exchanges with the first tissue compartment C₁, which in turn exchanges with the second tissue compartment C₂ where the tracer is trapped. K₁ [ml/ccm/min] and k₂ [1/min] are the uptake and clearance rate constants, whereas k₃ [1/min] describes the trapping. The usual interpretation is that C₁ represents free and non-specifically bound tracer in tissue (non-displaceable compartment), and C₂ irreversibly bound or metabolized tracer.
An important parameter of an irreversible configuration is net influx rate $K_i$, the unidirectional uptake rate constant that incorporates both net inward transport and trapping of the tracer in tissue.

$$\text{Net influx} = K_i = \frac{K_i k_3}{k_2 + k_3}$$

Another macroparameter "lambda*k_3" as been used in the context of enzymatic reactions, where $k_3$ is supposed to be proportional to enzyme concentration.

$$\lambda k_3 = (K_1/k_2)k_3$$

Logan et al. [1] have shown in simulations that lambda*k_3 is preferable to k_3 alone in terms of bias and precision.

**Differential Equation of the Mass Balance**

$$\frac{dC_1(t)}{dt} = K_i C_p(t) - (k_2 + k_3) C_1(t)$$
$$\frac{dC_2(t)}{dt} = k_3 C_1(t)$$

with input curve $C_p(t)$.

**Operational Model Curve**

$$C_{\text{Model}}(t) = (1 - vB)(C_1(t) + C_2(t)) + vB C_p(t)$$

with the concentration $C_B(t)$ of tracer in whole blood, and the blood volume fraction $v_B$.

**Parameter Fitting**

The model includes the 4 fitable parameters $v_B$, $K1/k2$, Flux, and calculates the LambdaK3 macroparameter. Usually, $v_B$ is fixed at a physiologic value of 3-5% to reduce the number of fitted parameters.

**Parameter Fitting using K1/k2 and Flux instead of k_2 and k_3**

The Irreversible 2-Tissue Compartment Model, K1/k2, Flux model variant uses $K1/k2$ and Flux as fitting parameters instead of $k_2$ and $k_3$. The advantage of the reformulation is that the Flux can be calculated as a regression slope with the robust Patlak model, and used as a fixed parameter in the compartment model fit. If the Model Conversion option is enabled, the Flux will be copied when switching from the Patlak model to the Irreversible 2-Tissue Compartment Model, K1/k2, Flux model. Similarly K1/k2 will be copied when switching from a 2-Tissue compartment model. Another potential application is the use of coupled fitting with a common $K1/k2$ across regions.

**Reference**

6.1.18 3-Tissue Compartment Model

This 3-tissue compartment model separates free tracer $C_1(t)$ and non-specifically bound tracer $C_3(t)$ in the non-displaceable compartment. $K_1$ and $k_2$ describe the exchange between arterial plasma and tissue, $k_3$ and $k_4$ between free and specifically bound tracer $C_2(t)$ in tissue, and $k_5$ and $k_6$ between free and non-specifically bound tracer.

\[
\begin{align*}
\frac{dC_1(t)}{dt} &= K_1 C_p(t) - (k_2 + k_3 + k_4) C_1(t) + k_4 C_2(t) + k_6 C_3(t) \\
\frac{dC_2(t)}{dt} &= k_3 C_1(t) - k_4 C_2(t) \\
\frac{dC_3(t)}{dt} &= k_5 C_1(t) - k_6 C_3(t)
\end{align*}
\]

with input curve $C_p(t)$.

Operational Model Curve

\[C_{\text{Model}}(t) = (1 - vB) (C_1(t) + C_2(t) + C_3(t)) + vB C_3(t)\]

with the concentration $C_3(t)$ of tracer in whole blood, and the blood volume fraction $v_B$.

Parameter Fitting

The model in principle includes the 7 fitable parameters $vB, K_1, k_2, k_3, k_4, k_5, k_6$, but reliable fitting cannot be achieved in practice. However, the model can be valuable for educational purposes and for generating simulation data.

6.1.19 3-Tissue Compartment Model, $K_1/k_2, V_s$

This 3-tissue compartment model separates free tracer $C_1(t)$ and non-specifically bound tracer $C_3(t)$ in the non-displaceable compartment. $K_1$ and $k_2$ describe the exchange between arterial plasma and tissue, $k_3$ and $k_4$ between free and specifically bound tracer $C_2(t)$ in tissue, and $k_5$ and $k_6$ between free and non-specifically bound tracer.

\[
\begin{align*}
\frac{dC_1(t)}{dt} &= K_1 C_p(t) - (k_2 + k_3 + k_4) C_1(t) + k_4 C_2(t) + k_6 C_3(t) \\
\frac{dC_2(t)}{dt} &= k_3 C_1(t) - k_4 C_2(t) \\
\frac{dC_3(t)}{dt} &= k_5 C_1(t) - k_6 C_3(t)
\end{align*}
\]
with input curve $C_p(t)$. 

**Operational Model Curve**

$$C_{\text{Model}}(t) = (1 - v_B) (C_1(t) + C_2(t) + C_3(t)) + v_B C_B(t)$$

with the concentration $C_b(t)$ of tracer in whole blood, and the blood volume fraction $v_B$.

**Parameter Fitting using Distribution Volumes**

This variant of the 3-Tissue Compartment Model uses three distribution volumes as fitting parameters: the distribution volume of free tracer ($K1/k2$), of non-specifically bound tracer $V_{ns} = (K_1 * k_5) / (k_2 * k_6)$, and specifically bound tracer $V_s = (K_1 * k_3) / (k_2 * k_4)$ instead of $k_2$, $k_4$ and $k_6$.

### 6.1.20 3-Tissue Compartment Model, Sequential

This model has been developed by Bertoldo et al. [1] for the analysis of FDG uptake in skeletal muscle. In this context $C_1$, $C_2$ and $C_3$ correspond to extracellular, intracellular and phosphorylated intracellular FDG. $K_1$ [ml/ccm/min] and $k_2$ [1/min] are the exchange rate constants between plasma and the extracellular space, $k_3$ [1/min] and $k_4$ [1/min] describe the transport in and out of the cell, $k_5$ [1/min] and $k_6$ [1/min] phosphorylation and dephosphorylation.

**Differential Equation of the Mass Balance**

$$\frac{dC_1(t)}{dt} = K_1 C_p(t) - (k_2 + k_3) C_1(t) + k_4 C_2(t) + k_6 C_3(t)$$

$$\frac{dC_2(t)}{dt} = k_3 C_1(t) - k_4 C_2(t) + k_5 C_3(t)$$

$$\frac{dC_3(t)}{dt} = k_5 C_3(t) - k_6 C_3(t)$$

$$C_{\text{Model}}(t) = (1 - v_B) [C_1(t) + C_2(t) + C_3(t)] + v_B C_B(t)$$

with input curve $C_p(t)$.

**Operational Model Curve**

$$C_{\text{Model}}(t) = (1 - v_B) (C_1(t) + C_2(t) + C_3(t)) + v_B C_B(t)$$

with the concentration $C_b(t)$ of tracer in whole blood, and the blood volume fraction $v_B$.

**Parameter Fitting**

The model includes the 6 fitable parameters $v_B$, $K_1$, $k_2$, $k_3$, $k_4$, $k_5$, $k_6$. Usually, $v_B$ is fixed at a physiologic value of 3-5%, and $k_6$ is fixed: $k_6 = 0$. The influx $K_i$ is given by

6.1.21 3-Tissue Compartments with Metabolites

If a labeled metabolite of the tracer enters tissue, the additional signal has to be accounted for in the model. The current model introduced by Fujita et al. [1] includes a second input curve $C_M(t)$ of a metabolite entering tissue and undergoing non-specific binding.

\[
C_1 = \frac{K_{1p} k_3 k_5}{k_2 k_4 + k_2 k_5 + k_3 k_5}
\]

C\(_1\) represents the non-displaceable compartment of the authentic ligand, C\(_2\) the specific binding of interest, and C\(_3\) metabolized ligand in tissue.

**Differential Equation of the Mass Balance**

\[
\frac{dC_1(t)}{dt} = K_{1p} C_p(t) - (k_{2p} + k_3) C_1(t) + k_4 C_2(t)
\]

\[
\frac{dC_2(t)}{dt} = k_3 C_1(t) - k_4 C_2(t)
\]

\[
\frac{dC_3(t)}{dt} = K_{1m} C_M(t) - k_{2m} C_3(t)
\]

with the input curves of authentic ligand and metabolite, $C_p(t)$ and $C_M(t)$, respectively. The input curve loading menu shows two corresponding entries as soon as the model is selected.

**Operational Model Curve**

\[
C_{\text{Model}}(t) = (1-vB) \left( C_1(t) + C_2(t) + C_3(t) \right) + vB C_3(t)
\]

with the concentration $C_\text{B}(t)$ of tracer in whole blood, and the blood volume fraction $vB$.

**Parameter Fitting**

The model includes the 7 fitable parameters $vB$, $K_{1p}$, $k_{2p}$, $k_3$, $k_4$, $K_{1m}$, $k_{2m}$. Due to the high number of parameters reliable fitting is difficult. There are two other model variants which support coupled fitting by using distribution volumes as fit parameters.

**Models with Alternative Fit Parameters**

There are two alternative models available which have the same structure but use some parameter ratios as fit parameters. The advantage is that these ratios might be candidates as common parameters among regions in a coupled fit, or might be fixed at known values.

**Reference**
6.1.22 3-Tissue Compartments with Metabolites, K1/k2

If a labeled metabolite of the tracer enters tissue, the additional signal has to be accounted for in the model. The current model introduced by Fujita et al. [1] includes a second input curve $C_M(t)$ of a metabolite entering tissue and undergoing non-specific binding. $C_1$ represents the non-displaceable compartment of the authentic ligand, $C_2$ the specific binding of interest, and $C_3$ metabolized ligand in tissue.

Differential Equation of the Mass Balance

$$\frac{dC_1(t)}{dt} = K_{1p} C_p(t) - (k_{2p} + k_3) C_1(t) + k_4 C_2(t)$$
$$\frac{dC_2(t)}{dt} = k_3 C_1(t) - k_4 C_2(t)$$
$$\frac{dC_3(t)}{dt} = K_{1m} C_M(t) - k_{2m} C_3(t)$$

with the input curves of authentic ligand and metabolite, $C_p(t)$ and $C_M(t)$, respectively. The input curve loading menu shows two corresponding entries as soon as the model is selected.

Operational Model Curve

$$C_{Model}(t) = (1 - vB) (C_1(t) + C_2(t) + C_3(t)) + vB C_3(t)$$

with the concentration $C_0(t)$ of tracer in whole blood, and the blood volume fraction $v_B$.

Parameter Fitting using $K_1/k_2$

The 3-Tissue Compartments with Metabolites, K1/k2 model variant uses the parameter ratios $K_{1p}/k_{2p}$, $K_{1m}/k_{2m}$ and $K_{1p}/K_{1m}$, as fitting parameters instead of $k_{2p}$, $K_{1m}$ and $k_{2m}$. The advantage of this formulation is that the ratios may be fixed at known or assumed values, or estimated in a coupled fit. The other four fitting parameters $v_B$, $K_{1p}$, $k_3$, and $k_4$ are the same as for the 3-Tissue Compartments with Metabolites model.

Reference


6.1.23 3-Tissue Compartments with Metabolites, K1/k2, Vš

If a labeled metabolite of the tracer enters tissue, the additional signal has to be accounted for in the model. The current model introduced by Fujita et al. [1] includes a second input curve $C_M(t)$ of a metabolite entering tissue and undergoing non-specific binding. $C_1$ represents the non-
displaceable compartment of the authentic ligand, $C_2$ the specific binding of interest, and $C_3$ metabolized ligand in tissue.

\[
\begin{array}{c}
C_P \\
\uparrow k_{2p} \\
C_1 \\
\rightarrow k_3 \\
C_2 \\
\hline
K_{1p} \\
\downarrow k_2p
\end{array}
\quad
\begin{array}{c}
C_M \\
\uparrow k_{1m} \\
C_3 \\
\hline
K_{1m} \\
\downarrow k_{2m}
\end{array}
\]

**Differential Equation of the Mass Balance**

\[
\begin{align*}
\frac{dC_1(t)}{dt} &= K_{1p} C_P(t) - (k_{2p} + k_3) C_1(t) + k_4 C_2(t) \\
\frac{dC_2(t)}{dt} &= k_3 C_1(t) - k_2 C_2(t) \\
\frac{dC_3(t)}{dt} &= K_{1m} C_M(t) - k_{2m} C_3(t)
\end{align*}
\]

with the input curves of authentic ligand and metabolite, $C_P(t)$ and $C_M(t)$, respectively. The input curve loading menu shows two corresponding entries as soon as the model is selected.

**Operational Model Curve**

\[
C_{\text{Model}}(t) = (1 - vB) (C_1(t) + C_2(t) + C_3(t)) + vB C_3(t)
\]

with the concentration $C_B(t)$ of tracer in whole blood, and the blood volume fraction $v_B$.

**Parameter Fitting using $K_1/k_2$ and $VS$**

The **3-Tissue Compartments with Metabolites, $K_1/k_2$, $VS$** model variant uses the parameter ratios $K_{1p}/k_{2p}$, $K_{1m}/k_{2m}$, $K_{1p}/K_{1m}$, and $VS=K_{1p}/k_{2p} \cdot k_3/k_4$ as fitting parameters instead of $k_{2p}$, $K_{1m}$, $k_{2m}$, and $k_4$. The advantage of this formulation is that the ratios may be fixed at known or assumed values, or estimated in a coupled fit. The other three fitting parameters $v_B$, $K_{1p}$ and $k_3$ are the same as for the **3-Tissue Compartments with Metabolites** model.

**Reference**


**6.1.24 4-Tissue Compartments with Metabolites, $K_1/k_2$**

This is a general compartment model for ligands which have recirculating metabolites entering tissue and contributing to tissue activity. Tissue kinetics of both the authentic ligand ($C_1$ and $C_2$) as well as the metabolites ($C_3$ and $C_4$) are described by a 2-tissue compartment model, with a transfer $k_{\text{trans}}$ constant linking the two exchangeable compartments which represents metabolites generated in tissue.
Two input curves are required: the authentic ligand in plasma $C_p(t)$, and the metabolites exchanging with tissue $C_M(t)$. The input curve loading menu shows two corresponding entries as soon as the model is selected.

### Operational Model Curve

$$C_{\text{Model}}(t) = (1 - vB) \left( C_1(t) + C_2(t) + C_3(t) + C_4(t) \right) + vB\ C_B(t)$$

with the concentration $C_B(t)$ of tracer in whole blood, and the blood volume fraction $v_B$.

### Parameter Fitting

The model includes more parameters than can be reliably fitted at once. Therefore it has been implemented with $K_1/k_2$ and $K_{1m}/k_{2m}$ as fitting parameters to allow fixing the distribution volume of the exchangeable compartment, or to use them as common parameters in a coupled fit.

### 6.1.25 Flumazenil Triple-injection Model

This model corresponds to a triple-injection protocol developed by Delforge et al. [1] for [11C] flumazenil PET. The setup includes three injections of labeled and/or unlabeled ligand during a single acquisition. The model is based on a 2-tissue compartment structure and accounts for the saturation of the benzodiazepine receptor. The uptake and binding of both labeled and unlabeled ligand is modeled by two parallel compartment models linked by common model parameters.

Differential Equation of the Mass Balance

They are linked by common model parameters and the system of differential equations.
Note that only the labeled ligand can be measured, both in PET and in the blood samples. Therefore the input curve of unlabeled ligand \( C_{u}(t) \) must be derived from the input curves of labeled ligand \( C_{p}(t) \). This task is complicated by the fact of metabolite build-up in plasma which must be corrected for. With \( T_{1}, T_{2} \) and \( T_{3} \) denoting the time of the three injections, the input curve \( C_{p}(t) \) of hot (labeled) tracer is calculated from the total plasma activity \( C_{\text{plasma}}(t) \) as follows:

\[
\frac{dC_{1}(t)}{dt} = k_{1}C_{p}(t) - k_{2}C_{1}(t) - \frac{k_{\text{met}}}{V_{R}} C_{1}(t)[B_{\max} - C_{2}(t) - C_{u}(t)] + k_{\text{off}}C_{2}(t)
\]

\[
\frac{dC_{2}(t)}{dt} = \frac{k_{\text{met}}}{V_{R}} C_{1}(t)[B_{\max} - C_{2}(t) - C_{u}(t)] - k_{\text{off}}C_{2}(t)
\]

\[
\frac{dC_{u1}(t)}{dt} = k_{1}C_{p}(t) - k_{2}C_{1}(t) - \frac{k_{\text{met}}}{V_{R}} C_{1}(t)[B_{\max} - C_{2}(t) - C_{u}(t)] + k_{\text{off}}C_{2}(t)
\]

\[
\frac{dC_{u2}(t)}{dt} = \frac{k_{\text{met}}}{V_{R}} C_{1}(t)[B_{\max} - C_{2}(t) - C_{u}(t)] - k_{\text{off}}C_{2}(t)
\]

This metabolite correction is implemented as the input curve model **Multiinjection, HOT**.

The input curve \( C_{u}(t) \) of cold (unlabeled) tracer is derived from \( C_{p}(t) \) by the expression

\[
C_{u}(t) = \begin{cases} 
C_{\text{plasma}} \left( A_{1}e^{B_{1}(t-T_{1})} + A_{2}e^{B_{2}(t-T_{2})} \right) & T_{1} < t < T_{3} \\
C_{\text{plasma}} \left( A_{1}e^{B_{1}(t-T_{1})} + A_{2}e^{B_{2}(t-T_{2})} \right) & t > T_{3}
\end{cases}
\]

which is implemented as the input curve model **Multiinjection, COLD**.

**Operational Model Curve**

Since PET only measures the radioactively labeled compounds the operational equation is given by

\[ C_{\text{mode}}(t) = (1 - v_{B}) (C_{1}(t) + C_{2}(t)) + v_{B} C_{g}(t) \]

with the concentration \( C_{g}(t) \) of labeled compounds in whole blood, and the blood volume fraction \( v_{B} \).

**Parameter Fitting**

The model includes six parameters \( v_{B}, K_{1}, k_{2}, K_{\text{on/VR}}, k_{\text{off}}, B_{\max} \) which are normally not possible to fit at once. However, the complex injection protocol results in more complex tissue TACs than with the usual single-injection protocols. As a consequence, the model has proven to result in reliable estimates of \( B_{\max} \).

**Reference**


**Acknowledgement**

The model implementation was supported in helpful discussions by Dr. Philippe Millet, University Hospital Geneva, Switzerland.
6.1.26 Water PET Model with Flow and Dispersion

This model implements the 1-tissue compartment model for $\text{H}_2\text{H}_1\text{O}$ water PET studies developed by E. Meyer [1].

![Model Diagram]

Operational Model Curve

It is assumed that at the arrival in tissue the true input curve $C_p(t)$ has been dispersed. This dispersion is modeled by convolving $C_p(t)$ with an exponential kernel

$$d(t) = \frac{1}{\tau} e^{-t/\tau}$$

whereby the parameter $\tau$ is called Dispersion. Incorporating the convolution into the 1-tissue compartment model and solving it using the Laplace transform yields the following analytic formula for the tissue concentration

$$C_1(t) = \tau K_1 C_p(t) + (1 - \tau k_2)K_1 e^{-k_2 t} \int_0^t C_p(\tau) e^{k_2 \tau} d\tau$$

$K_1$ represents flow, and $k_2$ flow divided by the partition coefficient $p$.

The operational equation is given by

$$C_{\text{Model}}(t) = (1 - vB) C_1(t) + vB C_B(t)$$

Note that in this particular case there is no difference between the plasma and blood concentrations $C_p(t)$ and $C_B(t)$ because water is a freely diffusible tracer.

Parameter Fitting

The tissue model includes four fitable parameters $vB$, $K_1$, $k_2$, (Flow/Part.Coeff), $\tau$ (Dispersion). Per default, however, $v_B$ is fixed at $v_B=0$ as in the original model. For water studies with the blood activity sampled in a peripheral artery, the resulting input curve is delayed respect to the arrival in tissue. In this case, the model should be fitted using the Fit region and blood delay button.

Implementation Note

Unlike the standard compartment models in PKIN the model curve of the operational equation is not calculated by integrating a set of differential equations, but rather using the analytical solution given above. The convolution integral is approximated by summing rectangles of 0.1sec in length, assuming that the acquisition time of dynamic water studies is short.

Reference


6.2 Compartment Models for Cardiac PET

<table>
<thead>
<tr>
<th>Model Name</th>
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<tr>
<td>Card. NH3 (de Grado)</td>
<td>1-Tissue compartment model with two spillover terms. Has been developed for</td>
</tr>
<tr>
<td></td>
<td>cardiac NH$_3$-PET studies with bolus administration. The left</td>
</tr>
</tbody>
</table>
Assumptions of the Cardiac Models

- The cardiac models usually apply a geometrical spillover with two correction terms: correction for signal from the left and right ventricular cavity.
- For calculating the myocardial perfusion per g tissue a density of 1.04 g/ml for myocardial tissue is applied.
- Usually the PET signal from the left ventricle or the atrium is used as a measure of the tracer activity in blood. Depending on the tracer, corrections are applied to this blood curve for calculation the concentration of parent tracer with is required as the input curve of the models.

Exceptions to these rules are specified in the description of the individual models.

6.2.1 Cardiac Perfusion from Ammonia PET

There are three model variants available for the quantification of myocardial blood flow from $^{13}$NH$_3$ ammonia bolus PET data, the 1-tissue compartment model and the 2-tissue compartment model with metabolic trapping. The 1-tissue model is preferably used with the first 2-4 minutes of a dynamic measurement, while the 2-tissue model is also adequate for longer durations. This may compensate to some extent the increased vulnerability to identifiability problems due to the higher number of fit parameters in the 2-tissue model.

6.2.1.1 Card. NH3 (de Grado)

The Card. NH3(De Grado) model has been developed by DeGrado et al. [1] for cardiac PET studies using $^{13}$NH$_3$ ammonia bolus injection. It is based on a 1-tissue compartment model with plasma activity $C_p(t)$ and total uptake in myocardium $C_{myo}(t)$.

![Operational Model Curve](image-url)
The plasma activity is calculated from the total activity in the left ventricular cavity $C_L(t)$ by a linear metabolite correction with factor $m_{\text{Corr}}$

$$C_p(t) = (1 - m_{\text{Corr}} \cdot t)C_L(t)$$

A value of $m_{\text{Corr}}=0.077 \ [1/\text{min}]$ has been found in humans [1]. The resulting differential equation for myocardial uptake is

$$\frac{dC_{\text{mys}}(t)}{dt} = K_1 C_p(t) - k_2 C_{\text{mys}}(t)$$

where $K_1$ equals myocardial blood flow $F$. The model curve incorporates a cardiac dual spillover correction, resulting in the operational equation

$$C_{\text{model}}(t) = (1 - V_{LV} - V_{RV})C_{\text{mys}}(t) + V_{RV}C_{LV}(t) + V_{RP}C_{RV}(t)$$

where

- $V_{LV} =$ spillover fraction of the blood activity in the left ventricle $C_{LV}(t)$,
- $V_{RV} =$ spillover fraction of the blood activity in the right ventricle $C_{RV}(t)$.

DeGrado et al. recommend to only use the first 4 minutes of data after injection of the tracer to reduce the effects of metabolite buildup and washout.

**Implementation**

When using the model from the PCARD module, the data are transferred appropriately. When using it in PKIN the blood data have to be loaded as follows:

- The left ventricle curve must be loaded as the **Plasma activity** curve. No metabolite correction needs to be enabled on the **Blood** panel of PKIN because it is included in the tissue model.
- The right ventricle curve must be loaded as the **Whole blood** curve.

The following automatic adjustments are performed within the model:

- The spillover fraction from the right ventricle $V_{RV}$ is automatically fixed to zero if the string "Sep" is not contained in the name a region. The assumption is that such a TAC is not from septal tissue and should thus be modeled with spillover from the left ventricle only.
- The valid flag for all data samples after 4 minutes is set to false, and they are consequently not considered in the fit.

**Parameter Fitting**

The model includes the 4 fitable parameters $F$, $v_{RV}$, $v_{LV}$ and $k_2$. Please inspect the %SE standard error to get information about the reliability of the parameter estimates.

**Reference**


6.2.1.2 Card. NH3 (UCLA, 2 Parameters)

This model implements a 2-tissue compartment model with only two fitted tissue rate constants. It was developed by Choi et al. for cardiac PET studies using $^{13}$NH$_3$ ammonia bolus injection and was called the **Modified Two-Parameter Model**, [1] eq (5).
Operational Model Curve

The system of compartmental differential equations for this compartment structure is given by

$$\frac{dC_1(t)}{dt} = K_1 C_1(t) - (k_2 + k_3) C_1(t)$$
$$\frac{dC_2(t)}{dt} = k_3 C_1(t)$$

with $K_1$ corresponding to myocardial blood flow $F$. The $k_3$ rate constant corresponding to glutamine formation was converted into a $K_1$-dependent value by the relation

$$k_3 = K_1 \left(1.65 e^{1.25/K_1} - 1\right)$$

Similarly, the $k_2$ rate constant of ammonia washout was converted into a $K_1$-dependent value by

$$k_2 = K_1 / V_{ND}$$

with the $V_{ND}$ representing the distribution volume of ammonia in the $C_1$ compartment which was assumed to be constant ($V_{ND} = 0.8$).

In contrast to Choi’s method, the model incorporates not only a spillover from the left ventricle, but for septal regions also from the right ventricle. The operational equation is then given by

$$C_{\text{Model}}(t) = (1 - V_{LV} - V_{RV}) (C_1(t) + C_2(t)) + V_{LV} C_{LV}(t) + V_{RV} C_{RV}(t)$$

where

$V_{LV} =$ spillover fraction of the blood activity in the left ventricle $C_{LV}(t)$,

$V_{RV} =$ spillover fraction of the blood activity in the right ventricle $C_{RV}(t)$.

Because the model does not include a metabolite correction, only the samples within 2 minutes after injection should be used for the fit [1].

Implementation Notes:

When using the model from the PCARD module, the data are transferred appropriately. When using it in PKIN the blood data have to be loaded as follows:

- The left ventricle curve must be loaded as the Plasma activity curve.
- The right ventricle curve must be loaded as the Whole blood curve.

The following automatic adjustments are performed within the model:

- The spillover fraction from the right ventricle $V_{RV}$ is automatically fixed to zero if the string "Sep" is not contained in the name a region. The assumption is that such a TAC is not from septal tissue and should thus be modeled with spillover from the left ventricle only.
- The valid flag for all data samples after 2 minutes is set to false, and they are consequently not considered in the fit.

Parameter Fitting

The model includes the three fitable parameters $F$, $v_{LV}$, $v_{RV}$. Please inspect the %SE standard error to get information about the reliability of their estimates. The input parameters $V_{nd}$ (0.8), EF scale (1.65) and EF Exp (1.25) are related to the conversion of $k_2$ and $k_3$ as described above and can be adjusted if needed.

Reference

6.2.1.3 Card. NH3 (2 Compartments)

The Card NH3 (2-Tissue) model developed by Hutchins et al. [1] is an implementation of the irreversible 2-tissue-compartment model for cardiac PET studies using $^{13}$NH₃ ammonia bolus injection. The compartment model has the following structure:

\[
\begin{array}{c}
\text{C}_p \\
\text{K}_1 \\
\text{K}_2 \\
\text{C}_1 \\
\text{C}_2
\end{array}
\]

where $C_1$ is free tracer in tissue, and $C_2$ is metabolically trapped tracer in the form of $^{13}$N glutamine. Because ammonia is considered in this model as freely diffusible across the capillary wall, the unidirectional uptake parameter $K_1$ equals the myocardial perfusion.

Operational Model Curve

The system of differential equations is:

\[
\begin{align*}
\frac{dC_1(t)}{dt} &= K_1C_p(t) - (k_2 + k_3)C_1(t) \\
\frac{dC_2(t)}{dt} &= k_3C_1(t)
\end{align*}
\]

To allow the fitting of data over an extended period, the model includes the exponential metabolite correction described by van den Hoff et al. [2] with a delay $t_0=0.48 \text{ min}$ and half-time $T_{1/2}=6.69 \text{ min}$. $C_{LV}(t)$ is the total tracer concentration measured in the left ventricular cavity, including metabolites.

Additionally, the model incorporates a cardiac dual spillover correction by the operational equation

\[
C_{\text{Model}}(t) = (1-V_{LV}V_{RV})(C_1(t) + C_2(t)) + V_{LV}(t)C_{LV}(t) + V_{RV}(t)C_{RV}(t)
\]

where

\[
V_{LV} = \text{spill-over fraction of the blood activity in the left ventricle } C_{LV}(t),
\]

\[
V_{RV} = \text{spill-over fraction of the blood activity in the right ventricle } C_{RV}(t).
\]

Implementation Notes:

When using the model from the PCARD module, the data are transferred appropriately. When using it in PKIN the blood data have to be loaded as follows:

- The left ventricle curve must be loaded as the Plasma activity curve. No metabolite correction needs to be enabled on the Blood panel of PKIN because it is included in the tissue model.
- The right ventricle curve must be loaded as the Whole blood curve.

The following automatic adjustments are performed within the model:

- The spill-over fraction from the right ventricle $V_{RV}$ is automatically fixed to zero if the string "Sep" is not contained in the name a region. The assumption is that such a TAC is not from septal tissue and should thus be modeled with spill-over from the left ventricle only.
- The valid flag for all data samples after 2 minutes is set to false, and they are consequently not considered in the fit.

Parameter Fitting

The model includes the five fitable parameters $F$, $v_{LV}$, $v_{RV}$, $k_2$, $k_3$. Please inspect the %SE standard error to get information about the reliability of their estimates. The input parameters MC...
T0 (0.48) and MC T12 (6.69) are related to the metabolite correction as described above and can be adjusted if needed.

Alternate Model Card NH3 (2-Tissue, K1/k2)
Due to the increased number of fit parameters it has been found, that the Card NH3 (2-Tissue) may suffer from identifiability problems. Therefore, the variant Card NH3 (2-Tissue, K1/k2) has been developed using the parameter K1/k2 (distribution volume of the first compartment) as a fit parameter instead of k2. In this configuration physiological restrictions can be imposed on K1/k2, or K1/k2 can be used as a common parameter in a coupled fit.

References

6.2.1.4 Card. NH3 (2 Compartments, K1/k2)
The Card NH3 (2-Tissue) model developed by Hutchins et al. [1] is an implementation of the irreversible 2-tissue-compartment model for cardiac PET studies using 13NH3 ammonia bolus injection. The compartment model has the following structure

\[
\begin{array}{ccc}
C_P & K_1 & C_1 \\
& k_2 & \\
& k_3 & C_2 \\
\end{array}
\]

where C1 is free tracer in tissue, and C2 is metabolically trapped tracer in the form of 13N glutamine. Because ammonia is considered in this model as freely diffusible across the capillary wall, the unidirectional uptake parameter K1 equals the myocardial perfusion.

Operational Model Curve
The system of differential equations is

\[
\begin{align*}
\frac{dC(t)}{dt} &= K_1C_P(t) - (k_2 + k_3)C_1(t) \\
\frac{dC_1(t)}{dt} &= k_3C_1(t)
\end{align*}
\]

To allow the fitting of data over an extended period, the model includes the exponential metabolite correction described by van den Hoff et al. [2]

\[
C_P(t) = \begin{cases} 
C_{LV}(t) & t \leq t_0 \\
\frac{1}{1 - e^{-k_2(t-t_0)}}C_{LV}(t) & t > t_0
\end{cases}
\]

with a delay \(t_0=0.48\) min and half-time \(T_{1/2} = 6.69\) min. \(C_{LV}(t)\) is the total tracer concentration measured in the left ventricular cavity, including metabolites.

Additionally, the model incorporates a cardiac dual spillover correction by the operational equation

\[
C_{Model}(t) = (1 - V_{LV} - V_{RV})C_1(t) + V_{LV}C_{LV}(t) + V_{RV}C_{RV}(t)
\]

where \(V_{LV}\) = spill-over fraction of the blood activity in the left ventricle \(C_{LV}(t)\), \(V_{RV}\) = spill-over fraction of the blood activity in the right ventricle \(C_{RV}(t)\).
Implementation Notes:
When using the model from the PCARD module, the data are transferred appropriately. When using it in PKIN the blood data have to be loaded as follows:

- The left ventricle curve must be loaded as the **Plasma activity** curve. No metabolite correction needs to be enabled on the **Blood** panel of PKIN because it is included in the tissue model.
- The right ventricle curve must be loaded as the **Whole blood** curve.

The following automatic adjustments are performed within the model:

- The spill-over fraction from the right ventricle \( V_{RV} \) is automatically fixed to zero if the string "Sep" is *not* contained in the name a region. The assumption is that such a TAC is not from septal tissue and should thus be modeled with spill-over from the left ventricle only.
- The valid flag for all data samples after 2 minutes is set to false, and they are consequently not considered in the fit.

Parameter Fitting
The model includes the five fitable parameters \( F, v_{LV}, v_{RV}, K_1/k_2, k_3 \). The parameter \( K_1/k_2 \) (distribution volume of the first compartment) is used as a fit parameter instead of \( k_2 \). In this configuration physiological restrictions can be imposed on \( K_1/k_2 \), or \( K_1/k_2 \) can be used as a common parameter in a **coupled fit** [3]. \( k_2 \) is derived from the estimated \( F (=K_1) \) and \( K_1/k_2 \). Please inspect the %SE standard error to get information about the reliability of their estimates.

The input parameters \( MC\ T_0 \) (0.48) and \( MC\ T_{12} \) (6.69) are related to the metabolite correction as described above and can be adjusted if needed.

References

### 6.2.2 Cardiac Perfusion from Rubidium-82 PET

Two kinetic models are available for the quantification of myocardial perfusion from \( ^{82}\)Rb bolus PET data.

1. A 1-tissue compartment model with geometrical spillover correction and correction for flow-dependent extraction.
2. A 2-tissue compartment model with recovery and spillover correction.

#### 6.2.2.1 Card. Rb-82 (1 Compartment)

The **Card. Rb82(1 Compartment)** model has been developed by Lortie et al. [1] for cardiac PET studies using \( ^{82}\)Rb bolus injection. It is based on a 1-tissue compartment model with plasma activity \( C_p(t) \) and total uptake in myocardium \( C_{myo}(t) \).

![Operational Model Curve](image)

The differential equation is given by
The model uses the activity in the left ventricular cavity $C_{LV}(t)$ as the input curve ($C_p(t) = C_{LV}(t)$). Rb is known to have a flow-dependent extraction fraction, so that $K_1$, which is the product of flow $F$ times extraction fraction $E$, is described by the expression

$$K_1 = F E = F \left(1 - A e^{-B/F}\right)$$

This expression is inserted for $K_1$ into the differential equation, so that $F$ becomes a fit parameter, and $K_1$ is derived from it. The values of the extraction parameters reported by Lortie et al. [1] are

$$A = 0.77$$
$$B = 0.63 \text{ [ml/min/g]}$$

They can be entered as input parameters into the model for changing the form of the extraction function. The model curve incorporates a cardiac dual spillover correction, resulting in the operational equation

$$C_{mode}(t) = (1 - V_{LV} - V_{RV}) C_{myo}(t) + V_{RV} C_{LV}(t) + V_{RV} C_{RV}(t)$$

where

$V_{LV} = \text{spill-over fraction of the blood activity in the left ventricular cavity } C_{LV}(t)$,
$V_{RV} = \text{spill-over fraction of the blood activity in the right ventricular cavity } C_{RV}(t)$

Implementation

When using the model from the PCARD module, the data are transferred appropriately. When using it in PKIN the blood data have to be loaded as follows:

- The left ventricle curve must be loaded as the Plasma activity curve.
- The right ventricle curve must be loaded as the Whole blood curve.

The following automatic adjustment is performed within the model:

- The spill-over fraction from the right ventricle $V_{RV}$ is automatically fixed to zero if the string "Sep" is not contained in the name a region. The assumption is that such a TAC is not from septal tissue and should thus be modeled with spill-over from the left ventricle only.

Parameter Fitting

The model includes the 4 fitable parameters $F$, $v_{RV}$, $v_{LV}$ and $k2$. Please inspect the %SE standard error to get information about the reliability of the parameter estimates. The scale factor $A$ (EF scale) and the exponent $B$ (EF exp) of the extraction function can be entered as input parameters.

Reference


6.2.2.2 Card. Rb-82 (2 Compartments)

The Card. Rb-82 (2 Compartments) model has been implemented according to the method described and evaluated by Herrero et al [1]. The model is based on the following compartment structure to describe the kinetics of rubidium in the myocardium:
where \( C_1(t) \) represents the fast exchangeable compartment (vascular and interstitial spaces), and \( C_2(t) \) the slow exchangeable compartment (intracellular space), myocardium flow \( F \), and rate constants \( k_1 \) and \( k_2 \).

**Operational Model Curve**

The differential equations for the activity concentrations in the different compartments are given by

\[
\frac{dC_1(t)}{dt} = F(C_p(t) - C_1(t)) - k_1 C_1(t) + k_2 C_2(t)
\]

\[
\frac{dC_2(t)}{dt} = k_1 C_1(t) - k_2 C_2(t)
\]

The model uses the activity in the left ventricular cavity \( C_{LV}(t) \) is used as the input curve \( (C_p(t) = C_{LV}(t)) \). The operational equation which is fitted to the measured data is

\[
C_{Model} = F_{MM} (C_1(t) + C_2(t)) + F_{BM} C_{LV}(t)
\]

where \( F_{MM} \) denotes the tissue recovery coefficient and \( F_{BM} \) the blood to myocardium spillover fraction.

**Implementation**

When using the model from the PCARD module, the data are transferred appropriately. When using it in PKIN the left ventricle curve must be loaded as the *Plasma activity* curve. The right ventricle curve is not used.

**Parameter Fitting**

The model encompasses 5 fitable parameters. However, in practice it is impossible to estimate so many parameters from a time-activity curve with reasonable identifiability. Therefore, at least the recovery coefficient \( F_{MM} \) is usually fixed. The default from Herrero et al. [1] is 0.65, but it depends on the image resolution and should be determined experimentally.

**Reference**


### 6.2.3 Cardiac Perfusion from Water PET

The **Card. H2O (Tissue fraction)** model has was developed by Hermannsen et al. [1] for cardiac PET studies using \( \text{H}_2^{15}\text{O} \) water bolus injection. The distribution of the freely diffusible inert tracer \( \text{H}_2^{15}\text{O} \) in myocardium can be described by a 1-tissue compartment model with myocardial blood flow \( F \) and the partition coefficient of water \( p \), which is defined as (ml water/ml tissue)/(ml water/ml blood).

![Cardinal H2O model diagram](image)

**Operational Model Curve**

The differential equation for the tracer concentration in myocardium is

\[
\frac{dC_{myo}(t)}{dt} = FC_p(t) - \frac{F}{p} C_{myo}(t)
\]

In practice, the left ventricular time-activity curve is used as the input curve \( C_p(t) \).
The model curve incorporates two spill-over terms from blood in the left and the right ventricular cavities which are relatively displaced in time. So the operational equation which is fitted to the measured data is

\[ C_{\text{Model}}(t) = TF C_{\text{myo}}(t) + V_{\text{LV}} C_{\text{LP}}(t) + V_{\text{RV}} C_{\text{RP}}(t) \]

where TF = tissue fraction, \( V_{\text{LV}} \) = spill-over fraction from the left ventricle, \( V_{\text{RV}} \) = spill-over fraction from the right ventricle.

**Implementation**

As opposed to most models in PKIN a tissue fraction is used for calculating the model curve rather than a strict geometrical correction for compliance with the original model. When using the model from the PCARD module, the data are transferred appropriately. When using it in PKIN the blood data have to be loaded as follows:

- The left ventricle curve must be loaded as the **Plasma activity** curve.
- The right ventricle curve must be loaded as the **Whole blood** curve.

The following automatic adjustment is performed within the model:

- The spill-over fraction from the right ventricle \( V_{\text{RV}} \) is automatically fixed to zero if the string "Sep" is not contained in the name a region. The assumption is that such a TAC is not from septal tissue and should thus be modeled with spill-over from the left ventricle only.

**Parameter Fitting**

The model encompasses the 5 fitable parameters \( F, TF, v_{\text{LV}}, v_{\text{RV}} \) and \( p \). However, in practice it is impossible to estimate so many parameters from a time-activity curve with reasonable identifiability. The partition coefficient is therefore fixed per default at a value of 0.96 [1], and the spill-over fraction from right ventricular blood \( v_{\text{RV}} \) is only fitted for septal tissue TACs.

**Reference**


### 6.2.4 Cardiac Perfusion from Acetate PET

Van den Hoff et al. [1] have investigated and validated \(^{11}\)C-acetate as a flow tracer. This methodology is implemented as the **Card Acetate (1 Compartment)** model. It employs a single tissue compartment model

![Diagram](https://via.placeholder.com/150)

with tracer exchange between arterial plasma \( C_p(t) \) and myocardial tissue \( C_{\text{myo}}(t) \).

**Operational Model Curve**

The differential equation \( C_{\text{myo}}(t) \) is given by

\[
\frac{dC_{\text{myo}}(t)}{dt} = K_1 C_p(t) - k_2 C_{\text{myo}}(t) = EF C_p(t) - k_2 C_{\text{myo}}(t)
\]

\( K_1 \) is the product of flow \( F \) and extraction \( E \)

\( K_1 = E \cdot F \)

For acetate, the extraction is flow dependent [1]
Furthermore a metabolite correction is necessary to derive the plasma activity from whole blood measured in the left cavity \([1]\)

\[
C_p = 0.91 e^{-\frac{b}{T_{1/2}}} C_{LV}(t)
\]

with \(T_{1/2}=5.3\) min. Including these relations into the differential equation yields

\[
\frac{dC_{\text{mod}}(t)}{dt} = F(1 - 0.64 e^{-1.2/F}) 0.91 e^{-\frac{b}{T_{1/2}}} C_{LV}(t) - k_2 C_{\text{myo}}(t)
\]

which is non-linear in \(F\). The model curve incorporates a cardiac dual spillover correction, resulting in the operational equation

\[
C_{\text{model}}(t) = (1-V_{LV}-V_{RV}) C_{\text{myo}}(t) + V_{RV} C_{LV}(t) + V_{RV} C_{RV}(t)
\]

where
- \(V_{LV}\) = spillover fraction of the blood activity in the left ventricular cavity \(C_{LV}(t)\),
- \(V_{RV}\) = spillover fraction of the blood activity in the right ventricular cavity \(C_{RV}(t)\)

### Implementation

When using the model from the PCARD module, the data are transferred appropriately. When using it in PKIN the blood data have to be loaded as follows:

- The left ventricle curve must be loaded as the **Plasma activity** curve. No metabolite correction needs to be enabled on the **Blood** panel of PKIN because it is included in the tissue model.
- The right ventricle curve must be loaded as the **Whole blood** curve.

The following automatic adjustment is performed within the model:

- The spillover fraction from the right ventricle \(V_{RV}\) is automatically fixed to zero if the string "Sep" is not contained in the name a region. The assumption is that such a TAC is not from septal tissue and should thus be modeled with spillover from the left ventricle only.

### Parameter Fitting

The model includes the 4 fitable parameters \(F\), \(k_2\), \(v_{LV}\), \(v_{RV}\). Please inspect the %SE standard error to get information about the reliability of the parameter estimates. The parameters for the correction of the flow-dependent extraction (EF scale, EF exp) and the metabolites (MC scale, MC T12) are initialized with the published values \([1]\), but can be edited if needed. The \(K1\) parameter of the 1-tissue compartment model equivalent to the EF product is provided as a macro parameter.

### Reference


### 6.3 Compartment Models for Cardiac SPECT

Due to recent improvements of SPECT cameras regarding sensitivity and time resolution, similar protocols as for cardiac perfusion PET have been developed and tested. To date, however, there is no consensus regarding the analysis of such data. In order for researchers to experiment with such data, PMOD offers one of the recently published methods, which fits well into the existing framework.
6.3.1 Cardiac Perfusion from $^{99m}$Tc Tetrofosmin SPECT

The Card. $^{99m}$Tc Tetrofosmin (Shrestha) model has been developed by Shrestha et al. [1] for cardiac SPECT studies using $^{99m}$Tc Tetrofosmin bolus injection. It is based on a 1-tissue compartment model with plasma activity $C_p(t)$ and total uptake in myocardium $C_{myo}(t)$.

![Diagram of plasma activity and myocardium uptake](image)

Operational Model Curve

The differential equation is given by

$$\frac{dC_{myo}(t)}{dt} = K_1 C_p(t) - k_2 C_{myo}(t)$$

The model uses the activity in the left ventricular cavity $C_v(t)$ as the input curve ($C_p(t) = C_v(t)$). Rb is known to have a flow-dependent extraction fraction, so that $K_1$, which is the product of flow $F$ times extraction fraction $E$, is described by the expression

$$K_1 = F E = F (1 - A e^{-B/F})$$

This expression is inserted for $K_1$ into the differential equation, so that $F$ becomes a fit parameter, and $K_1$ is derived from it. The values of the extraction parameters reported by Shrestha et al. [1] are

$$A = 0.91$$
$$B = 0.34 \text{ [ml/min/g]}$$

They can be entered as input parameters into the model for changing the form of the extraction function. The model curve incorporates a cardiac dual spillover correction, resulting in the operational equation

$$C_{Model}(t) = (1-V_{LV} - V_{RV}) C_{myo}(t) + V_{RV} C_{LV}(t) + V_{LV} C_{RF}(t)$$

where

- $V_{LV}$ = spillover fraction of the blood activity in the left ventricular cavity $C_{LV}(t)$,
- $V_{RV}$ = spillover fraction of the blood activity in the right ventricular cavity $C_{RV}(t)$

Implementation

When using the model from the PCARD module, the data are transferred appropriately. When using it in PKIN the blood data have to be loaded as follows:

- The left ventricle curve must be loaded as the Plasma activity curve.
- The right ventricle curve must be loaded as the Whole blood curve.

The following automatic adjustment is performed within the model:

- The spill-over fraction from the right ventricle $V_{RV}$ is automatically fixed to zero if the string "Sep" is not contained in the name a region. The assumption is that such a TAC is not from septal tissue and should thus be modeled with spill-over from the left ventricle only.

Parameter Fitting

The model includes the 4 fitable parameters $F$, $v_{RV}$, $v_{LV}$ and $k_2$. Please inspect the %SE standard error to get information about the reliability of the parameter estimates. If necessary, the scale factor $A$ (EF scale) and the exponent $B$ (EF exp) of the extraction function can be interactively edited.

Reference
PKIN Reference


6.4 Reference Tissue Models

PKIN features the following list of reference models which are mostly used for the analysis of receptor ligand studies:

<table>
<thead>
<tr>
<th>Model Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>4 Parameter Reference Tissue Model</td>
<td>4 Parameter Reference Tissue Model by Lammertsma et al.</td>
</tr>
<tr>
<td>SRTM</td>
<td>Simplified Reference Tissue Model by Lammertsma and Hume</td>
</tr>
<tr>
<td>SRTM2</td>
<td>Modification of Simplified Reference Tissue Model by Wu and Carson to reduce BP bias induced by noise</td>
</tr>
<tr>
<td>MRTM0</td>
<td>Original Multilinear Reference Tissue Model by Ichise et al.</td>
</tr>
<tr>
<td>MRTM</td>
<td>Multilinear Reference Tissue Model by Ichise et al modified to reduce bias induced by noise</td>
</tr>
<tr>
<td>MRTM2</td>
<td>Multilinear Reference Tissue Model by Ichise et al modified to reduce bias and variability induced by noise</td>
</tr>
<tr>
<td>Logan Non-Invasive</td>
<td>Reference tissue method which is based on an average $k_2$</td>
</tr>
<tr>
<td>2 Tissue Reference Model</td>
<td>Watabe’s reference tissue model with a 2-tissue compartment model in the reference region</td>
</tr>
<tr>
<td>MP4A RLS (Nagatsuka)</td>
<td>Multi-linear Reference Tissue Model for [11C]-MP4A (RLS)</td>
</tr>
</tbody>
</table>

6.4.1 Coupled fitting for Determination of $k_2'$ in Reference Tissue Models

Some of the reference methods require an a priori average value of $k_2'$, while other methods such as the MRTM or the SRTM reference methods estimate $k_2'$ together with the other parameters in every pixel.

Averaging $k_2'$ from Separate Regional Fits

In response to a request from a PMOD user Dr. Ichise recommended the following approach for calculating $k_2'$:

1. $k_2'$ is the tissue clearance rate from the reference region, e.g., cerebellum. This reference tissue is always a region (ROI), not a voxel for both ROI based PKIN or PXMOD parametric imaging. If you define a reference region, there should be only one correct $k_2'$ value for that particular subject (scan).
2. Logan in her original formulation of her reference tissue model suggested to determine $k_2'$ by using arterial data for a group of subjects and use this mean $k_2'$.
3. However, we showed that this $k_2'$ can be estimated for each individual without arterial data using MRTM or SRTM (three parameter estimation, one of the three parameters is $k_2'$).
4. I prefer to use \( k' \) estimated this way for each subject for the subsequent MRTM2 or SRTM2. This would be more accurate than the mean \( k' \) estimated for a group of subjects as above.

5. Now the accuracy of \( k' \) estimation depends on the following (see [52]): A) noise in the PET data, B) the magnitude of \( k' \) and C) the ratio of \( k'/k \) or \( k/k' \) determined by MRTM or SRTM.

6. The bias and variability of \( k' \) estimation by MRTM is less as \( k' \) is larger and \( k'/k \) or \( k/k' \) ratio is further away from unity. This ratio for fallypride using cerebellum and striatum should be greater than 3, I think. In that case, \( k' \) estimation from cerebellum and striatum (use ROIs) should be minimum (see figs in the paper).

7. Even using the ROIs, \( k' \) estimation is affected by noise and hence it is good to run MRTM a few times choosing ROIs with high BPND areas (say right stratum and left stratum) \( k'/k \) ratio is further away from 1) and average the \( k' \) values. This averaging is within the subject and totally different from population average.

8. Please use the \( k' \) determined as above for estimation of BPND for the cortical regions. The \( k' \) estimated with cerebellum and cortical regions is not accurate because \( k'/k \) is closer to unity.

9. One advantage of MRTM over SRTM: To use SRTM, both cerebellum and target must be 1 tissue (1T) kinetics (use of the SRTM for 2T will bias BPND). However, MRTM is good for tracers with 2T kinetics such as Fallypride. The only thing here is that you have to give a t* value.

**Estimating k2' using a Fit with Regional Coupling**

Alternatively, the a-priori knowledge that \( k' \) should always have the same value can be exploited using coupled fitting in PKIN with SRTM2 as described here. The \( k' \) value obtained in PKIN can then be used in PXMOD.

**Maintaining of k2' when Switching Reference Tissue Models**

Note that an estimated \( k' \) can be maintained when switching between reference tissue models by enabling Parameter Propagation. This allows for instance to keep \( k' \) estimated with SRTM2 when switching to the Logan reference tissue model.

### 6.4.2 Full Reference Tissue Model

**Model Configuration and Assumptions**

The Full Reference Tissue method (or 4 Parameter Reference Tissue Method) of Lammertsma et al. is used for the analysis of studies with reversibly binding neuroreceptor tracers. A reference tissue devoid of receptors is required which can be modeled by a single-tissue compartment model. Tissues of interest which include the target receptor are modeled by a 2-tissue compartment model.

In the model equations \( C_i'(t) \) represents the tissue TAC from the reference region \( (k=0 \text{ in the 2-tissue compartment model}) \), and \( C_j(t) \) the tissue TAC from a receptor-rich region \( (k>0) \). It is assumed that the distribution volume of the non-displaceable compartment is the same for tissues of interest and the reference tissue: \( K_2/K_4 = K'_2/K'_4 \). Furthermore the reference tissue should not affected by the pathology investigated under study.
Operational Model Curve
Defining the ratio of tracer delivery $R_1$ as $K_1/K_1'$ and the binding potential $BP_{ND}$ as $k_3/k_4$, the following operational equation can be derived for the measured tissue TAC in a receptor-rich region:

$$C_{mode}(t) = R_1 \left( C_T'(t) + aC_T'(t) \otimes e^{-at} + bC_T'(t) \otimes e^{-bt} \right)$$

The equation parameters and their relation to the model are given by:

$$a = (k_3 + k_4 - c)(c - r)/p$$
$$b = (d - k_3 - k_4)(d - r)/p$$
$$c = (s + p)/2$$
$$d = (s - p)/2$$
$$p = \sqrt{s^2 - q}$$
$$q = 4k_2k_4$$
$$r = k_2/R_1$$
$$s = k_2 + k_4$$

For convolution with the exponentials, the reference tissue TAC $C_T'(t)$ is resampled on a regular grid, which can be specified by the Resampling parameter.

Parameter Fitting
The operational equation includes four unknowns: $R_1$, $k_2$, $k_3$, and $BP_{ND}$ (after substitution of $BP_{ND}=k_3/k_4$), which can be fitted using nonlinear fitting techniques.

Reference

6.4.3 Simplified Reference Tissue Model (SRTM)

Model Configuration and Assumptions
The Simplified Reference Tissue Model (SRTM) of Lammertsma and Hume [1] is used for the analysis of studies with reversibly binding neuroreceptor tracers. A reference tissue devoid of receptors is required which can be modeled by a single-tissue compartment model.

The assumptions of the model are:

1. The distribution volume is the same for the tissue of interest and the reference tissue: $K_1/K_1' = K_1'/K_1$.
2. The kinetics in the receptor-rich tissue of interest is such that it is difficult to distinguish between the specific and the non-displaceable compartment; i.e. the tissue TAC can be fitted by a 1-tissue compartment model with an uptake rate constant $k_2a = k_2/(1+BP_{ND})$. Note that this assumption may not be valid for all tracers, and in this case SRTM calculates biased $BP_{ND}$ estimates.
The model is a further development of the 4 Parameter Reference Tissue method. It has been shown to be less dependent on the initial parameter values, more stable in the presence of noise, and provide more consistent estimates of relative tracer delivery $R_1$.

**Operational Model Curve**

Defining the ratio of tracer delivery $R_1$ as $K_1/K_1'$ and the binding potential $BP_{ND}$ as $k_3/k_4$, the following operational equation can be derived for the measured tissue TAC in a receptor-rich region:

$$C_{Model}(t) = R_1 C_1'(t) + \left[ k_2 - R_1 k_2' \left( 1 + BP_{ND} \right) \right] C_1'(t) \otimes e^{-k_2' \left( 1 + BP_{ND} \right)}$$

For convolution with the exponentials, the reference tissue TAC $C'(t)$ is resampled on a regular grid, which can be specified by the Resampling parameter.

**Parameter Fitting**

The operational equation includes four unknowns: $R_1$, $k_2'$, and $BP_{ND}$, which can be fitted using nonlinear fitting techniques. $k_2'$, the transfer of tracer from the reference tissue back to the plasma, is provided as a derived parameter, as well as $k_{2a}$.

**Note:** The reference methods MRTM2 and SRTM2 require $k_2'$ as an input parameter. The $k_2'$ resulting from the SRTM method is a suitable estimate. Therefore, when switching in PKIN from the SRTM model to the SRTM2 or MRTM2, $k_2'$ is automatically copied from SRTM, in case the Model conversion option in the Extras panel is enabled.

**Reference**


6.4.4 **Simplified Reference Tissue Model 2 (SRTM2)**

**Model Configuration and Assumptions**

The Simplified Reference Tissue Model 2 (SRTM2) was derived by Wu and Carson [1] from the SRTM with the aim of improved parametric mapping. The model structure is the same.
as well as the assumptions:

1. The distribution volume is the same for the tissue of interest and the reference tissue:
   \[ K_1/k_2 = K_1'/k_2'. \]

2. The kinetics in the receptor-rich tissue of interest is such, that it is difficult to distinguish between the specific and the free/non-specific compartment; i.e. the TAC can be fitted by a 1-tissue compartment model with an uptake rate constant \( k_{2a} = k_2/(1+BP_{ND}) \).

**Operational Model Curve**

The operational equation of the SRTM was re-written to allow for fixing of \( k_2' \). This is relevant for parametric mapping because the model in each pixel TAC uses the same reference TAC and therefore should employ the same \( k_2' \). Defining the ratio of tracer delivery \( R_1 \) as \( K_1/K_1' \) and the binding potential \( BP_{ND} \) as \( k_3/k_4 \), the following operational equation can be derived for the measured TAC in a receptor-rich region:

\[
C_{Model}(t) = R_1 C_T'(t) + R_1 [k_2' - k_{2a}] C_T'(t) \otimes e^{-k_2't}.
\]

The three unknowns \( R_1 \), \( k_2' \) and \( k_{2a} \) in this equation can be fitted using nonlinear regression techniques. The binding potential can then be calculated as

\[
BP_{ND} = R_1 \frac{k_2'}{k_{2a}} - 1.0
\]

For convolution with the exponentials, the reference tissue TAC \( C_T'(t) \) is resampled on a regular grid, which can be specified by the **Resampling** parameter.

**Parameter Fitting**

The operational equation includes three unknowns: \( R_1 \), \( k_2' \), and \( k_2a \) which can be fitted using nonlinear fitting techniques. Note that \( k_2' \), the transfer of tracer from the reference tissue back to the plasma, can also be fitted. This can be employed by first estimating \( k_2' \) with a coupled fit and adequate TACs, and then fixing it for all regional fits as described above. Furthermore, fixing of \( k_2' \) allows studying the bias introduced by an inadequate \( k_2' \).

**Note:** As with the SRTM method, BP_{ND} estimates from SRTM2 tend to be biased if the 1-tissue compartment model assumption does not apply. The magnitude of the bias is even larger, most likely because the fixed \( k_2' \) can not compensate any more a part of the model inadequacy.

**Reference**

6.4.5 Linear Simplified Reference Tissue Model (LSRTM)

This model was developed by Alpert et al. [1] to extend the concept of activation studies for including measurements targeting neurotransmitters and specific receptor populations. It uses the fact that cognitive activation increases neuronal firing rate, hereby increasing the endogenous neurotransmitter level and altering the kinetics of specifically bound radioligands. The methodology uses a single PET injection. The scan starts with an acquisition of the subject at rest (baseline), followed by an acquisition with the subject performing a task (activation). The data (after motion correction of the lengthy scan) is analyzed modeling the change of the endogenous neurotransmitter level as an effect which occurs immediately after the task onset, and diminishes exponentially over time. The method has been applied to [11C]raclopride [1] and more recently to [18F]fallypride data [2,3].

Note: The model has been called LSSRM [1,2] and LSRRM [3], but was renamed to LSRTM to correspond to the commonly used notation of the Simplified Reference Tissue Model SRTM.

Model Configuration and Assumptions

The model uses the same assumptions as the Simplified Reference Tissue Model (SRTM):

1. A reference tissue devoid of receptors exists which can be modeled by a single-tissue compartment model.
2. The distribution volume is the same for the tissue of interest and the reference tissue: $K_{1}'/k_{2}' = K_{1}/k_{2}$.
3. The kinetics in the receptor-rich tissue of interest is such that it is difficult to distinguish between the specific and the non-displaceable compartment; ie. the tissue TAC can be fitted by a 1-tissue compartment model with an apparent uptake rate constant $k_{2a} = k_{2}/(1+BP_{ND})$.

Additionally, it is assumed that at the onset of the activation, $k_{2a}$ is increased by the amplitude $\gamma$ of ligand displacement, and that the effect dies away with a decay constant $\tau$. This behavior is described by a time-dependent $k_{2a}$ as follows:

$$k_{2a}(t) = \begin{cases} k_{2a} + \gamma h(t) & \text{if } t < \text{Begin} \\ 0 & \text{if } t \geq \text{Begin, } u = t - \text{Begin} \end{cases}$$

$$h(t) = e^{-\tau u}$$

Operational Model Curve

Defining the ratio of tracer delivery $R_t$ as $K_{1}/K_{1}'$, the following operational equation can be derived for the measured tissue TAC in a receptor-rich region:

$$C_{model}(t) = R_t C_t(t) + k_{2} \int_{0}^{t} C_t(s)ds - k_{2a} \int_{0}^{t} C_t(s)ds - \gamma \int_{0}^{t} h(s)C_t(s)ds$$

From the parameters in the equation the binding potential $BP_{ND}$ can be calculated as
The main outcome of the model, however, is the amplitude of ligand displacement, for which a t-score value and its p-value for a one-tailed t-test are also provided.

### Parameter Fitting

The user has to define the **Activation** start time with the corresponding input parameter. The **Resampling** parameter defines the interpolation grid for the exponential convolution, which corresponds to the last integral term in the operational equation.

The parameters of the non-linear fitting procedure include R1, k2, k2a, gamma and tau. To reduce the number of fitted parameters tau is usually fixed at a value of 0.03min^{-1} (~23min half-life). From the fit parameters, BPnd is calculated as well as the t-score and the p-value of gamma.

### References


### 6.4.6 Logan's Reference Tissue Model based on Average k2'

#### Model Configuration and Assumptions

Logan et al. [1] developed a reference tissue method for reversible receptor ligands which does not depend on a specific model structure of the reference tissue. Assuming the presence of reference region TAC C_{r}'(t) with an average tissue-to-plasma clearance k_{2}'_{r}, the target tissue TAC C_{t}(t) is transformed and plotted as a function of the transformed reference TAC, as illustrated below.

The operational equation resembles a linear equation with the distribution volume ratio (DVR = BP_{nd}+1) as the slope plus an error term which decreases over time. Therefore the late part starting...
from a time \( t^* \) of the plotted samples can be fitted by a regression line and the slope used for calculating \( \text{BP}_{ND} \). The time \( t^* \) can be determined as the time after which no further significant increases in slope are observed.

### Operational Model Curve

The graphical plot of the Logan Reference Tissue method is described by the following equation with the form resembling a linear equation:

\[
\int_0^t \frac{C_T(t')dt'}{C_T(t)} = \text{DVR} \left[ \frac{\int_0^t \left( C_T(t') + \frac{C_T(t')}{k_2'} \right) dt'}{C_T(t)} \right] + b = (\text{BP}_{ND} + 1) \left[ \frac{\int_0^t \left( C_T(t') + \frac{C_T(t')}{k_2'} \right) dt'}{C_T(t)} \right] + b
\]

\( k_2' \) in the original publication was the population average \( k_2 \) determined for the reference tissue using blood sampling, but using the subject's own \( k_2' \) may be preferable.

### Parameter Fitting

After switching to the Logan Reference model in PKIN a suitable reference region must be selected. \( k_2' \) is an input parameter which must be manually entered, or estimated by using other models such as the MRTM or SRTM model. Logan Reference fits a linear regression to the tail of the plot starting at time \( t^* \). Note that \( t^* \) is specified in regular acquisition time, which is translated into the “funny time” of the x-axis and shown as the Start parameter. The result parameter of interest is the \( \text{BP}_{nd} \) which is calculated from the slope.

A reasonable value of \( t^* \) can be estimated by the based on the error criterion Max Err. For instance, if Max Err. is set to 10% and the fit box of \( t^* \) is checked, the model searches the earliest sample such that the deviation between the regression and all measurements is less than 10%. Samples earlier than the \( t^* \) time are disregarded for regression and thus painted in gray. In order to apply the analysis to the same data segment in all regions, please switch off the fit box of \( t^* \), propagate the model with the Copy to all Regions button, and then activate Fit all regions.

**Note:** The \( k_2' \) resulting from the SRTM or MRTM method might be a reasonable estimate for the average clearance rate \( k_2' \). Therefore, when switching in PKIN from the SRTM or MRTM model to Logan Reference model, \( k_2' \) of those methods is automatically copied to \( k_2' \), in case Model conversion in the Configuration panel is enabled. See above for recommendations how to estimate \( k_2' \).

### Reference


### 6.4.7 Ichise's Original Multilinear Reference Tissue Model (MRTM0)

#### Model Configuration and Assumptions

Starting from the operational equation of the blood-based Logan plot Ichise et al. derived three multi-linear reference tissue model variants MRTM0, MRTM and MRTM2 [1,2]. They all assume an initial equilibration time \( t^* \) from which on the derived multi-linear relation holds. However, if kinetics in the target tissue can be described by a 1-tissue compartment model (an assumption required for the SRTM), all data can be used for the fitting \((t^*=0)\). Otherwise an adequate \( t^* \) value has to be determined.

Assuming the presence of receptor-devoid reference region TAC \( C_r'(t) \), the target tissue TAC \( C_T(t) \) is transformed and plotted as a function of the transformed reference TAC, as illustrated below.
For the calculation of $BP_{ND}$ it is assumed that the non-displaceable distribution volumes in the tissue and reference regions are identical.

Operational Model Curve

The MRTM0 model curve is described by

$$\int_0^t \frac{C_T(r) dr}{C_T(t)} = \frac{V_T}{V_T'} \int_0^t \frac{C_T'(r) dr}{C_T'(t)} + \frac{V_T}{V_T'k_2'} \frac{C_T'(t)}{C_T(t)} + b$$

where $V_T$ and $V_T'$ are the total distribution volumes of $C_T(t)$ and $C_T'(t)$, $k_2'$ is the clearance rate constant from the reference region to plasma, and $b$ is the intercept term, which becomes constant for $T > t^*$. The multi-linear relationship above can be fitted using multi-linear regression, yielding three regression coefficients. From the first coefficient the binding potential can be calculated by

$$BP_{ND} = \frac{V_T}{V_T'} - 1.0 = DVR - 1$$

Parameter Fitting

After switching to the Ichise NonInvasive MRTM0 model a suitable reference region must be selected. It allows fitting a multi-linear regression to the operational equation within a range defined by the parameters $t^*$ and End Lin. The results are the three regression coefficients, and the derived binding potential $BP_{nd}$.

A reasonable value of $t^*$ can be estimated by the based on the error criterion Max Err. For instance, if Max Err. is set to 10% and the fit box of $t^*$ is checked, the model searches the earliest sample such that the deviation between the regression and all measurements is less than 10%. Samples earlier than the $t^*$ time are disregarded for regression and thus painted in gray. In order to apply the analysis to the same data segment in all regions, please switch off the fit box of $t^*$, propagate the model with the Copy to all Regions button, and then activate Fit all regions.

For receptor ligands with 1-tissue kinetics such as [11C]DASB the multi-linear equation is correct from $t^*=0$, and $b$ is equal to $(-1/k_2)$, where $k_2$ is the clearance rate constant from the tissue to plasma. Furthermore, $R_1 = K_r/K_n$, the relative radioligand delivery, can be calculated from the ratio of the second and third regression coefficients.

References


6.4.8 Ichise's Multilinear Reference Tissue Model (MRTM)

Model Configuration and Assumptions

Starting from the operational equation of the blood-based Logan plot, Ichise et al. derived three multi-linear reference tissue model variants MRTM0, MRTM and MRTM2 [1]. They all assume an initial equilibration time \( t^* \) from which on the derived multi-linear relation holds. However, if kinetics in the target tissue can be described by a 1-tissue compartment model (an assumption required for the SRTM), all data can be used for the fitting \((t^*=0)\). Otherwise an adequate \( t^* \) value has to be determined.

Assuming the presence of receptor-devoid reference region TAC \( C_T'(t) \), the target tissue TAC \( C_T(t) \) is plotted as a function of the transformed tissue TACs as illustrated below. For the calculation of \( BP_{ND} \) it is assumed that the non-displaceable distribution volumes in the tissue and reference regions are identical.

![Operational Model Curve](image)

Operational Model Curve

To reduce noise-related bias effects arising in the MRTM0 method Ichise et al. applied a strategy known to be effective in reducing the noise-induced bias for the models requiring blood data. To this end the equation of the MRTM0 method was rearranged to remove the noisy tissue radioactivity term \( C_T(t) \) from the independent variables. This approach resulted in a new method called MRTM with following operational equation for \( C_T(t) \):

\[
C_{\text{MRTM}}(t) = - \frac{V_T}{V_T + b} \int_0^t C_T'(\tau)d\tau + \frac{1}{b} \int_0^t C_T(\tau) d\tau - \frac{V_T}{V_T + k_2 b} C_T(t)
\]

The multi-linear relationship above can be fitted using multi-linear regression, yielding three regression coefficients. The binding potential can then be calculated by dividing the first two regression coefficients.
Furthermore, division of the first by the third regression coefficient yields an estimate of \( k_2' \).

### Parameter Fitting

After switching to the Ichise NonInvasive MRTM model a suitable reference region must be selected. It allows fitting the multi-linear operational equation to the tissue TAC using the data segment starting from \( t^* \). The results are the three regression coefficients, and the binding potential \( \text{BP}_{\text{ND}} \) and \( k_2' \).

A reasonable value of \( t^* \) can be estimated by the based on the error criterion \text{Max Err}. For instance, if \text{MaxErr} is set to 10% and the fit box of \( t^* \) is checked, the model searches the earliest sample such that the deviation between the regression and all measurements is less than 10%. Samples earlier than the \( t^* \) time are disregarded for regression and thus painted in gray. In order to apply the analysis to the same data segment in all regions, please switch off the fit box of \( t^* \), propagate the model with the Copy to all Regions button, and then activate Fit all regions.

For receptor ligands with 1-tissue kinetics such as [11C]DASB the multi-linear equation is correct from \( t^* = 0 \), and the clearance rate constant from the tissue to plasma \( k_2 \) is equal to the negative value of the second regression coefficient, \(-(1/b)\). Furthermore, \( R_1 = k_1/k_1' \), the relative radioligand delivery, equals the third regression coefficient.

**Note:** The reference methods MRTM2 and SRTM2 require \( k_2' \) as an input parameter. The \( k_2' \) resulting from the MRTM method above might be a suitable estimate. Therefore, when switching in PKIN from the MRTM model to MRTM2 or SRTM2, \( k_2' \) is automatically copied from MRTM, if the Model conversion option in the Extras panel is enabled.

### Reference


### 6.4.9 Ichise’s Multilinear Reference Tissue Model 2 (MRTM2)

#### Model Configuration and Assumptions

Starting from the operational equation of the blood-based Logan plot, Ichise et al. derived three multi-linear reference tissue model variants MRTM0, MRTM and MRTM2 [1]. They all assume an initial equilibration time \( t^* \) from which on the derived multi-linear relation holds. However, if kinetics in the target tissue can be described by a 1-compartment model (an assumption required for the SRTM), all data can be used for the fitting (\( t^* = 0 \)). Otherwise an adequate \( t^* \) value has to be determined.

Assuming the presence of receptor-devoid reference region TAC \( C_{r}'(t) \), the target tissue TAC \( C_r(t) \) is plotted as a function of the transformed tissue TACs as illustrated below. For the calculation of \( \text{BP}_{\text{ND}} \) it is assumed that the non-displaceable distribution volumes in the tissue and reference regions are identical.
Operational Model Curve

When applied to noisy data such as single-pixel TACs in parametric mapping, the MRTM method still suffers from a high variability. Assuming a known value of the reference tissue clearance rate $k_2'$ the MRTM operational equation can be reformulated as the MRTM2 operational equation:

$$C_{\text{Model}}(t) = \frac{V_T}{V_T'} \frac{1}{b} \left( \int_0^t C_T'(\tau) d\tau + \frac{1}{k_2'} C_T'(t) \right) + \frac{1}{b} \int_0^t C_T(\tau) d\tau$$

with only two regression coefficients $V_T/(V_T' b)$ and $1/b$ for $T > t^*$. The multi-linear relationship above can be fitted using multi-linear regression, yielding three regression coefficients. The binding potential is then calculated from the ratio of the two regression coefficients as

$$BP_{ND} = \frac{V_T}{V_T'} - 1.0 = DVR - 1$$

For receptor ligands with 1-tissue kinetics such as $[\text{11C}]$DASB the multi-linear equation is correct from $t^*=0$, and the clearance rate constant from the tissue to plasma $k_2'$ is equal to the negative value of the second regression coefficient, $-(1/b)$. Furthermore, $R_1 = K_1/K_1'$, the relative radioligand delivery, equals the first regression coefficient divided by $k_2'$.

Parameter Fitting

After switching to the Ichise NonInvasive MRTM2 model a suitable reference region must be selected. It allows fitting the multi-linear operational equation to the tissue TAC using the data segment starting from $t^*$ and a fixed value of $k_2'$ which needs to be specified. The results are the two regression coefficients, and the binding potential $BP_{ND}$.

A reasonable value of $t^*$ can be estimated based on the error criterion Max Err. For instance, if Max Err. is set to 10% and the fit box of $t^*$ is checked, the model searches the earliest sample such that the deviation between the regression and all measurements is less than 10%. Samples earlier than the $t^*$ time are disregarded for regression and thus painted in gray. In order to apply the analysis to the same data segment in all regions, please switch off the fit box of $t^*$, propagate the model with the Copy to all Regions button, and then activate Fit all regions.

Note: The $k_2'$ resulting from the SRTM or MRTM method may be a suitable estimate for use in MRTM2. Therefore, when switching in PKIN from the SRTM or MRTM model to MRTM2, $k_2'$ is automatically copied, if the Model conversion option in the Extras panel is enabled. Further suggestions how to obtain an adequate $k2'$ value are given above.
Reference


6.4.10 Watabe’s Reference Tissue Model with 2 Compartments

Model Configuration and Assumptions

The 2 Tissue Reference Tissue method proposed by of Watabe [1] has been developed for ligands which do not follow the usual assumptions of the reference tissue models. Instead of a 1-tissue compartment model, the reference tissue is described by a 2-tissue model as illustrated below. The second compartment $C_2'$ may represent specific or non-specific binding.

Operational Model Curve

Assuming that the non-specific distribution volume of both tissues is the same, the following operational equation for fitting the tissue TAC can be derived (see Millet et al [2], appendix C):

$$ C_{model}(t) = R_1 \left[ C_{T'}(t) + aC_{T'}(t) \otimes e^{-k_{1}-k_{2}} + bC_{T'}(t) \otimes e^{-k_{3}'} \right] $$

When the parameters $k_{2}'$, $k_{3}'$, and $k_{4}'$ are fixed to constants determined with receptor-rich tissue, only $R_1$ and $k_2$ need to be fitted for estimating the binding potential $BP_{ND}$. The equation parameters and their relation to the model are given by:

$$ a = \frac{k_{2}'k_{3}'}{k_{3}'+k_{4}'-k_{2}} $$

$$ a = \frac{k_{2}^2 - k_2(k_{3}' + k_{4}' + k_{3}' + k_{4}')} {k_{3}'+k_{4}'-k_{2}} $$

$$ BP_{ND} = \frac{V_T}{V_T'} - 1 = \frac{K_i k_{3}'}{K_i k_{2}'(1+k_{3}' / k_{4}') - 1} $$

For convolution with the exponentials, the reference tissue TAC $C_{T'}(t)$ is resampled on a regular grid, which can be specified by the Resampling parameter.

Parameter Fitting

After switching to the 2 Tissue Reference Model a suitable reference region must be selected. The operational equation includes six fitting parameters: $R_1$, $k_2$, $k_2'$, $k_3'$, $k_4'$. Per default, the fit boxes of $k_2'$, $k_3'$ and $k_4'$ are off with the idea, that the values are determined externally. However, any of them may also be fitted. Fitting is performed using nonlinear techniques, and the main outcome is $BP_{ND}$.

References
6.4.11 Patlak Reference Tissue Model

The Patlak plot has been developed by Patlak and Blasberg [1] for tracers undergoing irreversible trapping. Most often it is applied for the analysis of FDG, which can be modeled as a 2-tissue compartment model with $k_4=0$.

However, this model structure is not necessary for the application of the method. It is sufficient to have any compartment in the system which binds irreversibly.

When the plasma activity is not available, the Patlak plot can be employed as a reference method provided that there exists some tissue wherein tracer is not irreversibly trapped. The procedure simply replaces the input curve by the reference tissue TAC.

**Operational Model Curve**

The Patlak plot belongs to a group of *Graphical Analysis* techniques, whereby the measured tissue TAC $C'(t)$ undergoes a mathematical transformation and is plotted against some sort of "normalized time". The Patlak plot using reference tissue is given by the expression

$$\frac{C'_T(t)}{C'_T'(t)} = K \int_0^t \frac{C'_T(\tau)}{C'_T'(\tau)} d\tau + V$$

with the reference tissue TAC $C'_T(t)$. This means that the measured PET activity is divided by the reference tissue activity, and plotted at a "normalized time" (integral of the reference TAC from the injection time divided by the instantaneous reference activity). For systems with irreversible compartments this plot will result in a straight line after an equilibration time $t^*$.

Under several assumptions, including a common $K_1/k_2$, the slope of the linear regression represents the following relation

$$slope = K = \frac{k_3}{(k_2 + k_3)(1 - K_{eq})}$$

with the equilibrium constant $K_{eq}$.
The reference Patlak plot has been applied for the FDOPA PET tracer for calculating an index of the influx $K_i$. Both the cerebellum and the occipital lobe have been used as the reference [2].

Parameter Fitting

The Patlak Reference model calculates and displays the transformed measurements as described by the formula above. It allows fitting a regression line to the data segment starting at time $t^*$. The results are the slope $K$ and the intercept of the regression line.

$t^*$ can be specified manually, or a value estimated using an error criterion Max Err. For instance, if Max Err. is set to 10% and the fit box of $t^*$ is checked, the model searches the earliest sample so that the deviation between the regression line and all measurements is less than 10%. Samples earlier than the $t^*$ time are disregarded for regression and thus painted in gray. Note that $t^*$ must be specified in real acquisition time, although the x-axis units are in "normalized time". The corresponding normalized time which can be looked up in the plot is shown as a non-fitable result parameter Start. In order to apply the analysis to the same data segment in all regions, please switch off the fit box of $t^*$, propagate the model with the Copy to all Regions button, and then activate Fit all regions.

References:

6.4.12 Multi-linear Reference Tissue Model for [11C]-MP4A (RLS)

Model Configuration and Assumptions

The MP4A RLS (Nagatsuka) model has been developed for the non-invasive quantification method (RLS) of the acetylcholinesterase (AChE) activity in the human brain from measurements with the $^{11}$C-MP4A acetylcholine analog [1]. In contrast to reference methods for receptor tracers which use a reference devoid of specific binding, the present method uses a reference with very high AChE activity which immediately traps the tracer so there is no washout. $C_T(t)$ is the TAC from a cortical target region, and $C_{T'}(t)$ the TAC from the reference region (striatum or cerebellum). $k_3$ represents the rate of tracer hydrolysis by AChE.

Operational Model Curve

By applying the method of Blomqvist, the following multi-linear equation is derived
It can be fitted to $C_t(t)$ using multi-linear regression, yielding three regression coefficients from which three parameters of interest can be calculated:

$$R_1 = K_1 / K_1' = p_3$$
$$k_2 = -(p_3 + p_2 / p_1)$$
$$k_3 = p_2 / p_1$$

**Parameter Fitting**

After switching to the **MP4A RLS (Nagatsuka)** model a suitable reference region must be selected. The findings in different publications indicate that cerebellum yields more stable results than striatum, most likely due to the higher impact of motion on the signal from the small striatum than the large cerebellum.

While the regression coefficients represent the fitting parameters, the resulting rate constants are shown in the derived parameter section.

**Reference**


### 6.5 Non-Compartmental Models

PKIN includes some additional models, which are neither explicit compartment nor reference models. They include the important linear regression methods **Logan Plot** and **Patlak Plot**, recent alternative methods for estimating the total distribution volume, as well as simple methods to quickly calculate ratios between tissue TACs and/or the plasma concentration.

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<th>Description</th>
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<td><strong>Ichise MA1</strong></td>
<td>Reformulation of the Logan plot as a multilinear analysis to reduce noise-induced bias. Result is the total distribution volume.</td>
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6.5.1 Logan Plot

The Logan plot has been developed by Logan et al. [1] for ligands that bind reversibly to receptors and enzymes and is used for estimating the total distribution volume \( V_T \). Its results can be interpreted with respect to the 1- and 2-tissue compartment models.

![Operational Model Curve](image)

**Operational Model Curve**

The Logan plot belongs to a group of Graphical Analysis techniques, whereby the measured tissue TAC \( C_T(T) \) undergoes a mathematical transformation and is plotted against some sort of "normalized time". The Logan plot is given by the expression

\[
\frac{\int C_T(t) \, dt}{C_T(t)} = K \frac{\int C_p(t) \, dt}{C_T(t)} + b
\]

with the input curve \( C_p(t) \). This means that the tissue activity integrated from the time of injection is divided by the instantaneous tissue activity, and plotted at a "normalized time" (integral of the input curve from the injection time divided by the instantaneous tissue activity). For systems with reversible compartments this plot will result in a straight line after an equilibration time \( t^* \).

In the derivation of the Logan plot the PET signal is described as a sum of tissue activity plus a fractional plasma signal

\[
C_{model}(t) = C_T(t) + v_p C_p(t)
\]

unlike the operational equation of the compartment model. Under these premises the slope represents the total distribution volume \( V_T \) plus the plasma space \( v_p \) in the VOI, which is usually neglected. Alternatively an explicit blood volume correction is available via the \( v_B \) input parameter. In the case of \( v_B=0 \) (default setting), no correction is performed. For \( v_B>0 \), the tissue TAC is corrected by the scaled whole blood activity before the actual analysis.

\[
C_T(t) = \frac{C_{PET}(t) - v_B C_p(t)}{1 - v_B}
\]
The relation of the slope to the parameters of the compartment models is given by

\[ K \equiv V_T = \frac{k_1}{k_2} \quad \text{1 - tissue compartment model} \]
\[ K \equiv V_T = \frac{k_1}{k_2 (1 + k_3/k_4)} \quad \text{2 - tissue compartment model} \]

It has been found that the Logan plot is susceptible to noise in the data. Noise causes the true \( V_T \) to be underestimated, to a degree which not only depends on the noise level, but also on the local kinetics. The underestimation problem is particularly relevant in parametric mapping, where the pixelwise TACs suffer from a high noise level.

The reason for the underestimation effect is the fact that noise is not only present in the y-values (dependent variable) as the linear regression assumes, but also in the x-values (independent variable). To arrive at more accurate results it was therefore proposed to measure the residuals perpendicular to the regression line, rather than vertical to the x-axis [2].

### Parameter Fitting

The Logan Plot model calculates and displays the measurements transformed as described by the formula above. It allows to fit a regression line within a range defined by the start time of the linear section \( t^* \). The results are the distribution volume (slope) and the intercept. There is also an error criterion **Max Err** to fit \( t^* \). For instance, if **Max Err** is set to 10% and the fit box of \( t^* \) is checked, the model searches the earliest sample so that the deviation between the regression and all measurements is less than 10%. Samples earlier than the \( t^* \) time are disregarded for regression and thus painted in gray. Note that \( t^* \) must be specified in real acquisition time, although the x-axis units are in "normalized time". The corresponding normalized time which can be looked up in the plot is shown as a non-fitable result parameter **Start**.

The regression line is calculated using the traditional and the perpendicular distances, resulting in the total distribution volumes \( V_t \) and \( V_t_{\text{perpend}} \), respectively. It has been found that \( V_t \) has a tendency to underestimate the distribution volume at increasing noise levels. \( V_t_{\text{perpend}} \) shows less bias due to noise, but suffers from a somewhat increased variability.

### References


### 6.5.2 Patlak Plot

The Patlak plot has been developed by Patlak et al. [1] for tracers undergoing irreversible trapping. Most often it is applied for the analysis of FDG, which can be modeled as a 2-tissue compartment model with \( k_4 = 0 \).

The Patlak plot belongs to a group of Graphical Analysis techniques, whereby the measured tissue TAC \( C_2(T) \) undergoes a mathematical transformation and is plotted against some sort of "normalized time". The Patlak plot is given by the expression
with the input curve $C_p(t)$. This means that the measured PET activity is divided by plasma activity, and plotted at a "normalized time" (integral of the input curve from the injection time divided by the instantaneous plasma activity). For systems with irreversible compartments this plot will result in a straight line after an equilibration time $t^*$.

The slope $K$ and the intercept $V$ must be interpreted according to the underlying compartment model. For the FDG model mentioned, the slope $K$ equals $K_1k_3/(k_2+k_3)$ and represents the metabolic flux, while the intercept $V$ equals $V_0+v_B$ with the distribution volume $V_0$ of the reversible compartment $C_1$ and the fractional blood volume $v_B$.

Note that an explicit blood volume correction is available via the $v_B$ input parameter. In the case of $v_B=0$ (default setting), no correction is performed. For $v_B>0$, the tissue TAC is corrected by the scaled whole blood activity before the actual analysis.

Parameter Fitting

The Patlak Plot model calculates and displays the transformed measurements as described by the formula above. It allows fitting a regression line to the data segment starting at time $t^*$. The results are the Slope and the Intercept of the regression line.

$t^*$ can be specified manually, or a value estimated using an error criterion Max Err. For instance, if Max Err. is set to 10% and the fit box of $t^*$ is checked, the model searches the earliest sample so that the deviation between the regression line and all measurements is less than 10%. Samples earlier than the $t^*$ time are disregarded for regression and thus painted in gray. Note that $t^*$ must be specified in real acquisition time, although the x-axis units are in "normalized time". The corresponding normalized time which can be looked up in the plot is shown as a non-fitable result parameter Start. In order to apply the analysis to the same data segment in all regions, please switch off the fit box of $t^*$, propagate the model with the Copy to all Regions button, and then activate Fit all regions.

For the analysis of FDG data, the Lumped Constant (LC) and the Plasma glucose level (PG) of the patient should be entered. The metabolic rate of glucose $MRGlu$ is then obtained from the regression slope by

$$MRGlu = K_1 \frac{PG}{LC}$$

Note: The unit conversion is based on an assumed tissue density of 1.04g/ml.
Reference:

6.5.3 Spectral Analysis, vB

The Spectral Analysis, vB model allows performing a regular Spectral Analysis (SA) given a TAC and an input curve according to Cunningham and Jones [1].

Operational Model Curve of Spectral Analysis

The operational equation of SA is given by

\[ C_{Tissue}(t) = \sum_{i=1}^{N} a_i C_p(t) \otimes e^{-\beta_i t} = \sum_{i=1}^{N} a_i B_i(t) \]

that is, tissue uptake is modeled as a sum of N possible tissue responses, and like for the compartment models a fixed blood volume fraction is supported.

Due to the constraint of first order tracer kinetics, the coefficients \( a_i \) and the decay constants \( \beta_i \) must be non-negative. In practice, a discrete set of the decay constants \( \beta_i \) is selected which covers the physiologically reasonable range, typically logarithmically spaced in the range \([10^{-5},1]\text{sec}^{-1}\). The corresponding tissue responses

\[ B_i(t) = C_p(t) \otimes e^{-\beta_i t} \]

are the Basis Functions of spectral analysis. When fitting the operational equation above to a tissue TAC, the only unknowns are the coefficients \( a_i \), because only a pre-defined set of discrete \( \beta_i \) values is considered. Therefore, the problem is that of a non-negative linear least squares estimation (NNLS) with the constraint of non-negative coefficients.

An advantage of SA is the fact that no particular compartment structure is imposed. Rather, its result can be used to estimate how many kinetic tissue compartments can be resolved by PET. To this end, the results are plotted as a spectrum with the selected decay constants \( \beta_i \) along the x-axis (as the "frequencies") and the estimated coefficients \( a_i \) along the y-axis (as the "amplitudes"). Because of the large range, \( \log(\beta_i) \) is used in spectrum plotting rather than \( \beta_i \). The number of peaks in this spectrum corresponds to the number of distinct compartments. A peak appearing to the far left (low frequency, slow component) indicates irreversible trapping. A peak to the far right (high frequency, fast component) corresponds to kinetics indistinguishable from the input curve, thus to vascular contributions. Intermediate peaks represent compartments which exchange reversibly with plasma or with other tissue compartments.

Parameter Fitting

In the Spectral Analysis, vB model #Exponentials determines the number of discrete \( \beta_i \) decay constants which are logarithmically spaced in the range between Beta-min and the Beta-max. There is no well-defined minimum value as in the case of decay-corrected TACs where it is equal to the decay constant of the isotope. vB is the fractional volume of whole-blood in the signal, which is subtracted before the SA. Flux is the influx into the irreversible compartment and the main result of the model. vBest is the remaining blood component after subtracting the whole-blood contribution.

Estimates of the tissue uptake constant \( K1 \) and the total distribution volume \( Vt \) can be calculated based on the peaks found in the spectrum. nPeak is the number of peaks in the spectrum, whereby immediately neighboring spikes are grouped into a single peak.
The SA results can be visualized in 3 different ways with a corresponding radio button setting:

1. **Display TAC** shows the TAC together with the model curve calculated from the detected peaks.
2. **Display Spectrum** shows the peaks in the spectrum.
3. **Display unit IRF** shows the Impulse Response Function which depends on the **Exclude HF** setting.

### Reference


### 6.5.4 Spectral Analysis SAIF

The **Spectral Analysis SAIF** model allows performing a [Spectral Analysis (SA)] with an explicit trapping compartment [1].

#### Operational Model Curve of Spectral Analysis

The operational equation for this type of SA is given by

$$
C_{\text{Tissue}}(t) = a_0 \int_0^t C_p(t) dt + \sum_{i=1}^{N} a_i C_p(t) e^{-\beta_i t} = a_0 \int_0^t C_p(t) dt + \sum_{i=1}^{N} a_i B_i(t)
$$

$$
C_{\text{Model}}(t) = (1-vB) C_{\text{Tissue}}(t) + vB C_{\text{Blood}}(t) \geq C_{\text{PET}}(t)
$$

that is, tissue uptake is modeled as a sum of N possible tissue responses plus a flux $a_0$ into the irreversible compartment. Like for the compartment models a fixed blood volume fraction $vB$ is supported.

Due to the constraint of first order tracer kinetics, the coefficients $a_i$ and the decay constants $\beta_i$ must be non-negative. In practice, a discrete set of the decay constants $\beta_i$ is selected which covers the physiologically reasonable range, typically logarithmically spaced in the range $[10^{-5}, 1] \text{sec}^{-1}$. The corresponding tissue responses

$$
B_i(t) = C_p(t) \otimes e^{-\beta_i t}
$$

are the Basis Functions of spectral analysis. When fitting the operational equation above to a tissue TAC, the only unknowns are the coefficients $a_i$, because only a pre-defined set of discrete $\beta_i$ values is considered. Therefore, the problem is that of a non-negative linear least squares estimation (NNLS) with the constraint of non-negative coefficients.

An advantage of SA is the fact that no particular compartment structure is imposed. Rather, its result can be used to estimate how many kinetic tissue compartments can be resolved by PET. To this end, the results are plotted as a spectrum with the selected decay constants $\beta_i$ along the x-axis (as the “frequencies”) and the estimated coefficients $a_i$ along the y-axis (as the “amplitudes”). Because of the large range, $\log(\beta_i)$ is used in spectrum plotting rather than $\beta_i$. The number of peaks in this spectrum corresponds to the number of distinct compartments. A peak appearing to the far left (low frequency, slow component) indicates irreversible trapping. A peak to the far right (high frequency, fast component) corresponds to kinetics indistinguishable from the input curve, thus to vascular contributions. Intermediate peaks represent compartments which exchange reversibly with plasma or with other tissue compartments.

The Spectral Analysis with Iterative Filter (SAIF) approach uses a bandpass filter for selecting the real equilibrating compartments. The passband is defined by a range $[\beta_L, \beta_H]$. It is assumed that all $\beta_i < \beta_L$ are shifted noise components, and all $\beta_i > \beta_H$ shifted blood components. In an iterative process the following two steps are repeated...
1. Subtract the noise and blood components ($\beta_i$, outside $[\beta_L, \beta_H]$) from the signal, and estimate $a_0$ and $v_B$.
2. Subtract the trapping and blood components ($a_0$ and $v_B$) from the signal and estimate the equilibrating components.

The iteration is repeated until the residuals from step 1 and 2 are very similar.

**Parameter Fitting**

In the **Spectral SAIF** model **#Exponentials** determines the number of discrete $\beta_i$ decay constants which are logarithmically spaced in the range between **Beta-min** and the **Beta-max**. There is no well-defined minimum value as in the case of decay-corrected TACs where it is equal to the decay constant of the isotope. The passband of equilibrating compartments is defined by **Beta-cutoff low** ($\beta_L$) and **Beta-cutoff high** ($\beta_H$). $v_B$ is the fractional volume of whole-blood in the signal, which is subtracted before the SA. **Flux** is the influx into the irreversible compartment and the main result of the model. **vBest** is the remaining blood component after subtracting the whole-blood contribution.

Estimates of the tissue uptake constant $K_1$ and the total distribution volume $V_t$ of all reversible compartments can be calculated based on the peaks found in the spectrum. $nPeak$ is the number of peaks in the spectrum, whereby immediately neighboring spikes are grouped into a single peak.

The SA results can be visualized in 3 different ways with a corresponding radio button setting:

1. **Display TAC** shows the TAC together with the model curve calculated from the detected peaks.
2. **Display Spectrum** shows the peaks in the spectrum.
3. **Display unit IRF** shows the Impulse Response Function which depends on the **Exclude HF** setting.

**Reference**


### 6.5.5 Ichise Multilinear Analysis MA1

Ichise's MA1 analysis method [1] is a technique for calculating the total distribution volume for receptor tracers with reversible binding. It is a further development of the **Logan Plot** [17] aimed at minimizing the bias induced by noise in the measurements.

**Operational Model Curve**

The following bilinear relationship was derived

$$C_{Model}(t) = -\frac{V_T}{b} \int_0^t C_p(t) dt + \frac{1}{b} \int_0^t C_T(t) dt$$

where $C_T(t)$ represents the tissue time-activity curve, $C_p(t)$ the plasma activity, $V_T$ the total distribution volume, and $b$ the intercept of the Logan plot which becomes constant after an equilibration time $t^*$. 
Note that an explicit blood volume correction is available via the \( vB \) input parameter. In the case of \( vB=0 \) (default setting), no correction is performed. For \( vB>0 \), the tissue TAC is corrected by the scaled whole blood activity before the actual analysis as follows.

\[
C_T(t) = \frac{C_{PET}(t) - vBC_B(t)}{1 - vB}
\]

Based on simulation and experimental data the authors show that MA1 demonstrates the largest bias reduction among several methods. Therefore they conclude, that MA1 is the method of choice for calculating the total distribution volume, if \( t^* \) can accurately be defined.

### Parameter Fitting

The bilinear regression is performed using singular value decomposition. The result are the two coefficients \( VT/b \) and \( b \), from which \( Vt \) can easily be derived.

The MA1 method requires specification of the equilibration time as the \( t^* \) parameter. As it equals the equilibration time of the Logan plot, it is possible to perform a Logan plot first and use the resulting Logan \( t^* \). \( t^* \) can also be estimated using an error criterion Max Err. For instance, if Max Err. is set to 10% and the fit box of \( t^* \) is checked, the model searches the earliest sample so that the deviation between the regression and all measurements is less than 10%. Samples earlier than the \( t^* \) time are disregarded for regression and thus painted in gray. In order to apply the analysis to the same data segment in all regions, please switch off the fit box of \( t^* \), propagate the model with the Copy to all Regions button, and then activate Fit all regions.

### Reference


### 6.5.6 Ichise Multilinear Analysis MA2

Ichise's MA2 analysis method [1] is another alternative technique developed to calculate the total distribution volume of reversible receptor systems with minimal bias. It is particularly suitable for tracers with slow kinetics and low to moderate noise.

The MA2 method has two advantages:

1. It is independent of an equilibration time, so that the data from the first acquisition can be included into the regression.
2. An estimate of the specific distribution volume is also obtained. The authors state, that although the method has been derived with the 2-tissue model, it still shows a good performance with the data representing only 1-tissue characteristics.
Operational Model Curve

Based on the 2-tissue compartment model equations the following multi-linear relationship was derived [34]:

\[
C_{\text{Model}}(t) = p_1 \int_0^t C_p(t) \, dt + p_2 \int_0^t C_T(t) \, dt + p_3 \int_0^t C_T(t) \, dt + p_4 \int_0^t C_p(t) \, dt
\]

where \(C_T(t)\) represents the tissue time-activity curve, and \(C_p(t)\) the plasma activity.

A multi-linear regression can be performed to calculate the four regression coefficients from the transformed data. The total distribution volume \(V_T\) is then calculated as the ratio of the first two regression coefficients, and the distribution volume of specific binding \(V_S\) by the expressions

\[
V_T = \frac{p_1}{p_2}, \quad V_S = \frac{-p_2(p_1 + p_3 p_4) + p_3 p_4^2}{p_2(p_1 + p_3 p_4)}
\]

Note that an explicit blood volume correction is available via the \(v_B\) input parameter. In the case of \(v_B=0\) (default setting), no correction is performed. For \(v_B>0\), the tissue TAC is corrected by the scaled whole blood activity before the actual analysis as follows.

\[
C_T(t) = \frac{C_{PET}(t) - v_B C_B(t)}{1 - v_B}
\]

Parameter Fitting

The multi-linear regression is performed using a singular value decomposition, resulting in the 4 regression parameters \(p_1, p_2, p_3, p_4\), and the derived distribution volumes \(V_T\) and \(V_S\). Although no equilibration time is required for MA2, there is a \(t^*\) parameter to disregard early samples from the regression as for the Logan plot and MA1.

Reference


6.5.7 MLAIR (Multiple Linear Analysis for Irreversible Radiotracers)

The MLAIR method has been developed by Kim et al. [1] for tracers undergoing irreversible trapping. It is an alternative to the Patlak plot analysis which is dependent on the specification of an equilibration time \(t^*\) which may vary among tissues, and may suffer from bias when applied to noisy data. It has applied for the analysis of FDG, which can be modeled as a 2-tissue compartment model with \(k_4=0\), and 11C-labeled MeNTI [1].
Operational Model Curve

The authors develop two approaches, both of which are included in the Irreversible Ki (MLAIR) model. The 2-tissue compartment model structure with $k_4=0$ is assumed. A blood volume fraction $v_B$ is taken into account, but applied to the metabolite-corrected input curve $C_p(t)$ rather than the whole-blood activity.

The operational equation of method M Lair1 for the measured tissue TAC $C_T(t)$ is given by

$$C_{Mode}(t) = P_1 C_p(t) + P_2 \int C_p(\tau) d\tau + P_3 \int C_T(\tau) d\tau + P_4 \int\int C_p(s) ds d\tau$$

The linear coefficients are related to the model parameters as follows, allowing to calculate the influx $K_i$ from $P_3$ and $P_4$:

$$P_1 = v_B, \quad P_3 = -(k_2 + k_3), \quad P_2 = K_1k_3, \quad P_4 = -P_4/P_3$$

In order to avoid the $P_4/P_3$ division of M Lair1 which might introduce artefacts for noisy data, a second multilinear operational equation M Lair2 was developed which estimates $K_i$ as a direct regression parameter. M Lair2 has the following operational equation for the integrated tissue activity curve:

$$C_{Mode}(t) = \int C_T(\tau) d\tau = P_1 C_p(t) + P_2 \int C_p(\tau) d\tau + P_3 C_T(t) + P_4 \int\int C_p(s) ds d\tau$$

$$P_1 = v_B/(k_2 + k_3), \quad P_3 = -1/(k_2 + k_3), \quad P_2 = K_1/(k_2 + k_3) + v_B, \quad P_4 = K_i$$
Parameter Fitting

As soon as the **Irreversible Ki (MLAIR)** model is selected it fits the MLAIR1 and MLAIR2 model curves to the data using singular value decomposition and shows the regression coefficients. The parameter of interest is $K_i$ which is calculated with both methods: $K_i(\text{MLAIR1})$ and $K_i(\text{MLAIR2})$.

The curve area initially shows the tissue TAC together with the MLAIR1 fit, whereas the MLAIR2-related curves are hidden because they have a completely different dynamic range. However, they can easily be shown by enabling the respective check boxes in the curve control area, as illustrated above.

Reference:

### 6.5.8 RE-GP Analysis

In 2009 Zhou et al. introduced a new graphical method, the Relative-Equilibrium (RE) plot [1]. It can be applied with a plasma input curve for the calculation of the distribution volume, and with a reference tissue curve for the calculation of the binding potential. It was shown with Raclopride data and with simulations, that unlike the Logan plot the RE plot is not suffering from bias due to high noise levels. As a consequence, the results obtained with VOI-averaged TACs is consistent to the results obtained in pixel-wise applications.

However, it was found that violation of the relative equilibrium condition did introduce bias. To compensate this bias Zhou et al [2] combined the RE plot with the Patlak plot in a bi-graphical manner called the **RE-GP Analysis**.

**Operational Model Curve**

The operational equation of the RE Plot is given by:

$$\frac{\int_0 T \cdot C_T(t) \, dt}{C_p(t)} = V_{T \cdot RE} \frac{\int_0 T \cdot C_p(t) \, dt}{C_p(t)} + \alpha$$
For the RE plot to be applicable there must exist a time t* after which two conditions are fulfilled:

1. The plasma input curve must be mono-exponential. This condition can be verified by fitting a single exponential to the late part of the plasma curve on the Blood tab of PKIN.
2. The ratio of \( C_T/C_p \) is constant. This condition can be verified by switching the KM model to the Tissue/Plasma Ratio model.

Under these conditions the tracer in all tissue compartments reaches equilibrium relative to plasma. Note that the conditions must be verified explicitly, because the linear appearance of the RE plot is not a sufficient criterion.

Violation of the relative equilibrium condition above introduces bias. To compensate this bias the RE plot was combined with the Patlak plot

\[
\frac{C_T(t)}{C_p(t)} = \frac{K \int_0^t C_p(\tau)d\tau}{C_p(t)} + \beta
\]

using the same t* for fitting two respective regression lines. A consistent and unbiased distribution volume is then obtained by combining the slopes and intercepts of the two plots:

\[
V_T = V_{TRE} - \frac{\alpha K}{\beta}
\]

For the pixel-wise application of the RE-GP Analysis the results of the Patlak plot are smoothed, so that the calculation turns into

\[
V_T = V_{TRE} - \frac{\alpha K_s}{\beta_s}
\]

where \( K_s \) and \( \beta_s \) are obtained from spatially smoothed maps of K and \( \beta \).
Parameter Fitting

The **RE-GP Analysis** model calculates and displays the measurements transformed as described by the RE plot formula above. It allows fitting a regression line using the data points after a time \( t^* \). Note that \( t^* \) must be specified in real acquisition time, although the x-axis units are in "normalized time". The corresponding normalized time which can be looked up in the plot is shown as a non-fitable result parameter **Start**.

If \( t^* \) is changed to define a new data segment, the program finds the closest acquisition start time, fits the two regression lines to the RE plot and the Patlak plot, and updates the calculated parameters. The main outcome is the distribution volumes calculated with the RE-GP analysis (\( V_{t-REGP} \)). For a comparison, \( V_t \) calculated with the RE plot alone (\( V_{t-RE} \)) is also shown.

Per default, only the curves of the RE plot and its regression line are shown in the curve area. However, the Patlak plot and its regression line can also be visualized by enabling their boxes in the curve control area. Note that the values of the Patlak plot may have a very different dynamic range than the RE plot. Therefore it is recommended switching off the RE curves when switching on the Patlak ones.

References


6.5.9 Ito Plot

In 2010 Ito et al. introduced a new graphical method for the analysis of reversible receptor ligand binding [1]. It is applied with a plasma input curve for the calculation of the total distribution volume \( V_T \), the distribution volume of the nondisplaceable compartment \( V_{ND} \), and therefore also allows calculation of the binding potential \( BP_{ND} \). Furthermore, the shape of the plot gives an indication whether there the tissue includes is specific binding or not not.

**Operational Model Curve**

The equation of the graphical plot called is given by the equation

\[
\frac{C_T}{\int_0^t C_P(\tau) d\tau} = \frac{a}{b} - \frac{\int_0^t C_T(\tau) d\tau}{\int_0^t C_P(\tau) d\tau}
\]

The following cases can be distinguished:

1. The plot is a straight line, indicating the absence of specific binding. In this case the line equation parameters have the following interpretation in terms of a 2-tissue compartment model: \( a = K_1 \) (y-intercept), \( b = k_2 \) (slope), and the x-intercept equals \( V_{ND} \).

2. The plot has a curved shape as illustrated below, indicating the presence of specific binding. In this case two regression lines are fitted to the plot: a line to the early part within a short time segment \( T_1 \) to \( T_2 \), and a line to the end part after an equilibration time \( t^* \). The parameters of the resulting lines can be interpreted as follows:
   - Early line: \( a_1 = K_1 \), \( b_1 = k_2 \), x-intercept = \( V_{ND} \) of a 2-tissue compartment model.
   - Late line: \( a_2 = K_1 \), \( b_2 = k_2 \), x-intercept = \( V_T \) of a 1-tissue compartment model.
The binding potential can finally be calculated by $BP_{ND} = (V_T - V_{ND})V_{ND}$.

**Parameter Fitting**

The Ito Plot model calculates and displays the measurements transformed as described by the Ito plot formula above. It allows fitting two regression lines as follows:

1. The first regression line is fitted within the data segment specified by the times $T1$ and $T2$ (in acquisition time).
2. The second regression line is fitted to the late data segment starting at equilibration time $t^*$.

If any of the times is changed to define a new data segment, the program finds the closest acquisition start time, fits the two regression lines, and updates the calculated parameters.

$t^*$ can be specified manually, or a value estimated using an error criterion $\text{Max Err}$. For instance, if $\text{Max Err.}$ is set to 10% and the fit box of $t^*$ is checked, the model searches the earliest sample so that the deviation between the regression line and all measurements is less than 10%. Samples earlier than the $t^*$ time are disregarded for regression and thus painted in gray. Note that $t^*$ must be specified in real acquisition time, although the x-axis units are in "normalized time". The corresponding normalized time which can be looked up in the plot is shown as a non-fitable result parameter $\text{Start}$. In order to apply the analysis to the same data segment in all regions, please switch off the fit box of $t^*$, propagate the model with the Copy to all Regions button, and then activate Fit all regions.

Per default, only the Ito Plot and the late regression line is shown in the curve area. However, the early regression line can also be visualized as illustrated above by enabling its box in the curve control area.

**References**


**6.5.10 Partial Saturation Method, Data-driven**

Wimberley et al. [1] have developed a data driven strategy for the partial saturation approach (ddPSA) [2] for determining reliable regional estimates of receptor density $B_{avail}$ and in vivo affinity $1/appK_D$. It is a reference tissue approach applicable to 11C raclopride which makes it ideal for use in longitudinal studies of neurodegenerative diseases in the rodent.

**Operational Equation**

The non-linear 2-tissue compartment model with saturable specific binding is illustrated below.
The differential equations for a target tissue can be rearranged to

\[
\frac{C_S(t)}{C_{ND}(t)} = \frac{B_{avail} - C_S(t)}{appK_D} = \frac{dC_S(t)}{dt} \cdot \frac{1}{k_{off} C_{ND}(t)} = \frac{B_{avail} - C_S(t)}{appK_D} - DET(t)
\]

where \(C_S\) represents the concentration of specifically bound ligand, \(C_{ND}\) the concentration of non-displaceable ligand, \(k_{off}\) the dissociation constant (taken from in vitro data) and \(appK_D\) is the reciprocal of the in vivo receptor-ligand affinity. DET is a term related to the dynamic equilibrium of the system.

The ddPSA approach uses the scaled concentration of a reference tissue as an approximation of the \(C_{ND}\) in the target tissue

\[C_{ND}(t) = r C_{Ref}(t)\]

where \(r\) is determined in a presaturation study.

The time course \(C_S(t)\) is then calculated by

\[C_S(t) = C_I(t) - C_{ND}(t)\]

where \(C_I(t)\) represents the activity concentration in the target tissue.

Using \(C_S(t)\), \(C_{ND}(t)\) and the in vitro \(k_{off}\), DET(t) is plotted. When the DET is negligible compared to the \(C_S/C_{ND}\) value, the PET scan is in a dynamic equilibrium state and the \(C_S\) and \(C_{ND}\) values can be used for estimation of \(B_{avail}\) and \(appKD\).

**Parameter Fitting**

The *Partial Saturation, data-driven* model is a reference tissue model. The scale factor \(r\) for the reference tissue is an input parameter to be determined in a separate experiment. The specific activity \(Spec.\ Act.\) is required for converting kBq/cc data to pmol/ml. \(t_{start}\) and \(t_{end}\) define the range within which the data is considered for the fit. \(k_{off}\) is the in vitro dissociation constant, and \(Threshold\) defines the cutoff value for DET relative to \(C_S/C_{ND}\).

The fitting procedure switches off all time points with \(|DET(t)|\) above the threshold, and estimates \(B_{avail}\) and \(appKD\). As illustrated below, the various curves involved in the model calculations can be enabled for display, for instance DET(t).
6.5.11 Bolus/Infusion Optimization

When a compartment system has equilibrated, the total distribution volume can be calculated easily as the ratio of tracer concentration in tissue to the metabolite-corrected plasma concentration with a single static scan. It has been found that the time required to reach equilibrium can be shortened by an optimized tracer delivery. A setup which does not require a sophisticated tracer delivery system is to apply an initial bolus and continue with a constant infusion.

Carson et al. [1] have developed a method to optimize the ratio between the activity given as the initial bolus and the activity level of the subsequent infusion for quickly reaching an equilibrium (Appendix B in [1]). It is assumed that tissue TACs as well as the input curve have been measured beforehand in a bolus experiment. Ideally then, the measured activities represent the impulse response of the system. Under this assumption it is possible to calculate the tissue response for any given input curve by just convolving the TACs with the input curve.

Operational Model Curve

For a bolus and infusion (B/I) experiment the input function can be described as an initial impulse followed by a step function, and the convolution can be analytically solved. The resulting model curve

\[
C_{\text{Mode}}(t) = \frac{K_{\text{bol}} C_I(t) + \int_0^t C_I(\tau) \, d\tau}{K_{\text{bol}} + T}
\]
is a function of the bolus fraction and the infusion duration. $K_{bol}$ [min] defines the dose of the bolus expressed as minutes of infusion, and $T$ is the total duration of the infusion (end of last frame).

Note that an explicit blood volume correction is available via the $vB$ input parameter. In the case of $vB=0$ (default setting), no correction is performed. For $vB>0$, the tissue TAC is corrected by the scaled whole blood activity before the actual analysis.

$$C_T(t) = \frac{C_{PET}(t) - vB C_D(t)}{1 - vB}$$

**Parameter Fitting**

It is assumed that the data of a bolus study has been loaded. When the **Bolus/Infusion Opt** model is selected, the expected tissue activity according to the formula above is calculated using the default parameters, and shown as the model curve.

$K_{bol}$ can be estimated. The user has to enter a reasonable equilibration time as the input parameter **Start Equilibr.** When **Fit current region** is activated, the optimal $K_{bol}$ is calculated. The criterion for an equilibrium is that the calculated activities from **Start Equilibr** to the end of the scan are as constant as possible. Using the duration of the current data, the % **Fraction** of the dose in the syringe is displayed which is to be injected as the initial bolus.

Also, please note the following:

1. The same formulation holds for both TACs and input curves. However, $K_{bol}$ can only be fitted with the TAC curve, not the input curve. For the blood curve, the **Bolus/Infusion Opt** model is only usable for visualizing the plasma response with an entered $K_{bol}$ value which was obtained in TAC fitting.

2. If the **Bolus/Infusion Opt** model is applied to a TAC and the **Fit blood** is enabled in the **Extras** panel, the **Bolus/Infusion Opt** model is also applied to the plasma curve during the fit using the plasma model curve. Therefore the **plasma model should be an interpolation model, NOT THE B/I model/Kbol** is a common parameter in the fit, and the criterion is that both the expected TAC and the expected input curve are constant.

3. The **Bolus/Infusion Opt** model can be used for coupled fitting to find the optimal dose for several tissues. In a coupled fit, only the TACs are used, and the input curve is currently neglected (as opposed to 2).

4. The residual weighting of all curves involved in the fit should be set to constant weighting.

**Reference**


6.5.12 Cumulated Activity (OLINDA, IDAC)

The **Cumulated Activity (OLINDA, IDAC)** model performs pre-processing steps for the calculation of the absorbed dose from diagnostic or therapeutic radiopharmaceuticals. Given the time-course of the activity in a volume of tissue, it calculates the **Normalized Cumulated Activity** [1] (OLINDA *Residence Time*). The normalized cumulated activity represents the total number of disintegrations which have occurred during an integration time per unit administered activity. Ideally, integration is performed from the time of administration to infinity.

The normalized cumulated activity values resulting from the **Cumulated Activity (OLINDA, IDAC)** model can serve as input to a program such as OLINDA [2] or IDAC2.1 for the actual calculation of the absorbed organ doses.

Note that all steps required for the analysis of dosimetry data are explained in detail in the PMOD Workflow for Dosimetry Preprocessing application guide.

**Operational Equations**

Given an activity $A_0$ applied to a subject, and a measured (NOT decay corrected) activity $A(t)$ in an organ, the Normalized Cumulated Activity $\tau$ is calculated by

$$\tau = \frac{\int_0^\infty A(t) dt}{A_0}$$

Note that the unit of $\tau$ is usually given as [Bq·hr/Bq]. The measured part of the organ activity can easily be numerically integrated by the trapezoidal rule. For the unknown remainder of $A(t)$ till infinity it is a conservative assumption to apply the radioactive decay of the isotope. This exponential area can easily be calculated and added to the trapezoidal area. Alternatively, if the measured activity curve has a dominant washout shape, a sum of exponentials can be fitted to the measured data and the entire integral algebraically calculated.

The **Cumulated Activity (OLINDA, IDAC)** model in PKIN supports both the trapezoidal integration approach as well as the use of fitted exponentials. It furthermore supports some practicalities such as

- conversion of an average activity concentration in [kBq/cc] to VOI activity in [Bq] by multiplication with the VOI volume;
- reversion of decay correction;
Model Input Parameters

The Cumulated Activity (OLINDA, IDAC) model has 4 input parameters which need to be specified interactively by the user. These settings can easily be propagated from one region to all others with the Model & Par button.

<table>
<thead>
<tr>
<th>Decay corrected</th>
<th>Check this box if the loaded curves have been decay corrected as is usually the case for PET imaging studies. The Halflife of the radioisotope will be used to undo the decay correction.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Activity concentration</td>
<td>Check this box if the loaded curves represent activity concentration rather than the total activity in a region. The volume of the region will be multiplied with the signal to calculate and show the activity in [Bq].</td>
</tr>
</tbody>
</table>
| Isotope tail | Check this box to use the radioactive decay shape for the integration from the end of the last frame to infinity. This option is only relevant if exponentials are fitted to the measurement.  
For the Rectangle and Trapezoid methods it will always be enabled. For the exponentials it will also be enabled if the exponential halftime is longer than the isotope half-life. |
| Integration Selection | The integration method can be selected in the option list:  
| The Rectangle method is the natural selection for PET values which represent the time average during the frame duration. If there are gaps between the PET frames, the uncovered area is approximated by the trapezoidal rule.  
The Trapezoid method uses trapezoidal areas between the frame mid-times. |
With an **Exponential** selection, the specified number of decaying exponentials is fitted to the measurements, and used for analytical integration. Note that samples marked as invalid are neglected in the fit.

<table>
<thead>
<tr>
<th><strong>Exp integ from 0</strong></th>
<th>Check this box to use the exponential from time zero for the integration. If the box is unchecked, rectangular integration is used for samples before <strong>Exp begin</strong>.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Exp begin</strong></td>
<td>Defines the time (frame mid), from when on the exponentials are fitted to the measurement.</td>
</tr>
</tbody>
</table>

**Editing of Constituents**

Depending on the import of the activity curves, isotope and activity information may or may not be available. They can be edited in the Subject and Study Information window as illustrated below. Activate **Edit Subject** in the Tools list, select **SUV PARAMETERS**, and enter **Radionuclide half-life** and **Calibrated dose in syringe**. Note that the activity needs to be calibrated to time 0 of the activity curve.

The regional VOI volume is available if the TAC was generated in PMOD's View tool and transferred to PKIN. Often, not the entire volume of the organ can be outlined, because it is not clearly visible, or because (like muscle), it is not compact. In this case, standard organ volumes can be used for extrapolating VOI uptake to organ uptake. An organ can be assigned to a defined VOI as illustrated below:

1. Select the VOI in the list.
2. Open the VOI properties editor, and select the **List** button.
3. Select an appropriate name list. For instance, the **OLINDA 2 Human (Adult Female)** contains the organ definitions (name, volume, mass) for the adult male phantom in OLINDA 2.
4. Choose the organ from the list. The predefined **ORGAN volume** and **mass** are shown and may be edited.
5. Close with the **Set VOI name and color** button.
An alternative approach is to edit the VOI and organ volumes as follows:

1. Activate **Edit Data** in the **Tools** list
2. Select a region in the upper list.
3. Choose **Edit volume** from the curve tools
4. Enter the **Volume** of the VOI in the dialog window.
5. Change The **Organ Volume** in the lower part. (Note: currently, Organ mass is not used for the calculation).
6. Close the **PKIN data explorer** with **Ok**.

**Model Output Parameters**

As soon as the **Fit Region** button is activated, the fits and calculations are performed, resulting in a list of parameters:
| **Organ cumulated activity [Bq hr/Bq]** | Total number of disintegrations in the organ volume from time 0 until infinity per unit injected activity. This is the main result and equal to \( \text{Cumulated activity/cc} \times \text{Organ Volume} \). Note that it is zero as long as the Organ volume is not defined. |
| **VOI cumulated activity [Bq hr/Bq]** | Total number of disintegrations in the regional VOI volume from time 0 until infinity per unit injected activity, calculated as \( \frac{\text{VOI AUC}}{\text{Activity}} \). |
| **Cumulated activity/cc [Bq/cc hr/Bq]** | Total number of disintegrations per unit volume and unit injected activity. Equal to \( \frac{\text{VOI cumulated activity}}{\text{VOI Volume}} \). |
| **Dose [MBq]** | Administered activity at the time of scan start. Shown for verification only. Can be changed via Edit Patient. |
| **Halflife [sec]** | Isotope half-life. Shown for verification only. Can be changed via Edit Patient. |
| **Organ Volume [ccm]** | Volume of the organ which corresponds to the regional VOI. Shown for verification only. Can be changed via Edit Data. |
| **Organ Mass [g]** | Volume of the organ which corresponds to the regional VOI. Shown for verification only. Can be changed via Edit Data. |
| **VOI Volume [ccm]** | Volume of the VOI in which the uptake is measured. Shown for verification only. Can be changed via Edit Data. |
| **VOI AUC [Bq-hr]** | Integral of the VOI activity curve from time 0 till infinity. |
| **Amplitude 1, 2, 3 [Bq]** **Halftime 1, 2, 3 [min]** | Amplitudes and halftimes of the fitted exponentials. In the case of 1 exponential a linear regression is applied, whereas iterative fitting is used with 2 or 3 exponentials. |

### Generate Output File for Dosimetry Programs

After the **Cumulated Activity (OLINDA, IDAC)** model has been configured and fitted for all organs, a `.cas` file for direct import into OLINDA or a `.idac` file for import into IDAC2.1 can be created as follows:

1. Select **Edit Data** from the **Tools** list in the lower right.
2. Switch the operation below the curve list to \textbf{Dosimetry OLINDA/EXM Export} or \textbf{Dosimetry IDAC Export}.

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure.png}
\caption{OLINDA/EXM Output}
\end{figure}

The appearing dialog window lists the phantoms supported by OLINDA/EXM. Please select the appropriate one, but note that OLINDA/EXM doesn't interpret this information during loading of the \texttt{.cas} file. The option \textbf{Use default Olinda dose} should normally be on. Otherwise, the dose information in the patient SUV panel will be used which will most probably cause problems upon import, because OLINDA/EXM crashes when the dose values are not integer.

Finally specify a location and a name for saving the \texttt{.cas} file which can be loaded in OLINDA/EXM using the \textbf{Retrieve Case} button.
Note that the region names must be spelled exactly as below for the export/import to work properly.

Adrenals
Brain
Breasts_including_skin
Gallbladder_contents
LLI_contents
SI_contents
Stomach_contents
ULI_contents
Heart_contents
Heart_wall
Kidneys
Liver
Lungs
Muscle
Ovaries
Pancreas
Red_marrow
Cortical_bone
Trabecular_bone
Spleen
Testes
Thymus
Thyroid
Urinary_bladder_contents
Uterus
Fetus
Placenta

The difference between the injected activity and the activity accumulated in all organs is treated as the Remainder.

**OLINDA 2 Output**

The appearing dialog window lists the phantoms supported by the export. Note that within OLINDA 2 the phantoms can be easily switched, whereby the activities are propagated.

![OLINDA 2 Output](image)

Please select the appropriate one and save the file which is stored with .cas suffix. When opening the file in Olinda 2, the isotope and the model are selected, and the cumulated activities can be inspected on the Set Kinetics panel.
The **Remainder** value is automatically calculated from the total normalized number of disintegrations of the isotope (\(=\text{Half-time}[\text{h}] / \ln(2)\)) minus the activity found in the organs (excluding the bladder). It shows up in the **Total Body/Remainder** field.

**IDAC Output**

The appearing dialog window

Please select the appropriate one and save the file which is stored with `.idac` suffix. When opening the file in IDAC2.1 the cumulated activity will be shown and available for calculating the doses.
Note that the VOI names for the organs must be defined exactly as follows:

<table>
<thead>
<tr>
<th>Female</th>
<th>Male</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adipose tissue</td>
<td>Adipose tissue</td>
</tr>
<tr>
<td>Adrenals</td>
<td>Adrenals</td>
</tr>
<tr>
<td>Alveolar-interstitial</td>
<td>Alveolar-interstitial</td>
</tr>
<tr>
<td>Brain</td>
<td>Brain</td>
</tr>
<tr>
<td>Breast</td>
<td>Breast</td>
</tr>
<tr>
<td>Bronchi basal cells</td>
<td>Bronchi basal cells</td>
</tr>
<tr>
<td>Bronchi secretory cells</td>
<td>Bronchi secretory cells</td>
</tr>
<tr>
<td>Bronchiolar secretory cells</td>
<td>Bronchiolar secretory cells</td>
</tr>
<tr>
<td>Endosteal cells</td>
<td>Endosteal cells</td>
</tr>
<tr>
<td>ET1 basal cells</td>
<td>ET1 basal cells</td>
</tr>
<tr>
<td>ET2 basal cells</td>
<td>ET2 basal cells</td>
</tr>
<tr>
<td>Extrathoracic lymph nodes</td>
<td>Extrathoracic lymph nodes</td>
</tr>
<tr>
<td>Gall bladder</td>
<td>Gall bladder</td>
</tr>
<tr>
<td>Heart wall</td>
<td>Heart wall</td>
</tr>
<tr>
<td>Kidneys</td>
<td>Kidneys</td>
</tr>
<tr>
<td>Left colon</td>
<td>Left colon</td>
</tr>
<tr>
<td>Lens of eye</td>
<td>Lens of eye</td>
</tr>
<tr>
<td>Liver</td>
<td>Liver</td>
</tr>
<tr>
<td>Muscle</td>
<td>Muscle</td>
</tr>
<tr>
<td>Oesophagus</td>
<td>Oesophagus</td>
</tr>
<tr>
<td>Oral mucosa</td>
<td>Oral mucosa</td>
</tr>
<tr>
<td>Ovaries</td>
<td>Pancreas</td>
</tr>
<tr>
<td>Pancreas</td>
<td>Pituitary gland</td>
</tr>
<tr>
<td>Pituitary gland</td>
<td>Prostate</td>
</tr>
<tr>
<td>Rectosigmoid colon</td>
<td>Rectosigmoid colon</td>
</tr>
<tr>
<td>Red (active) marrow</td>
<td>Red (active) marrow</td>
</tr>
<tr>
<td>Right colon</td>
<td>Right colon</td>
</tr>
<tr>
<td>Salivary glands</td>
<td>Salivary glands</td>
</tr>
<tr>
<td>Skin</td>
<td>Skin</td>
</tr>
<tr>
<td>Small intestine</td>
<td>Small intestine</td>
</tr>
</tbody>
</table>
Use of Organ Lists for VOI Definition

The OLINDA and IDAC organ lists have been implemented in the VOI facility to support correct VOI naming. Illustrated below is use of these lists to assign proper name and organ mass to a defined VOI.

References


6.5.13 Area under Curve (AUC)

This is an auxiliary model which cannot be used for fitting. It calculates the following curves from the input curve $C_F(t)$ and the selected regional tissue TAC $C_T(t)$ at the frame mid-times:
1. The AUC of the tissue TAC as a function of time:
   \[ AUC_T(t) = \int_0^t C_T(\tau) d\tau \]

2. The AUC of the input curve as a function of time:
   \[ AUC_p(t) = \int_0^t C_p(\tau) d\tau \]
   Note that the integration uses the interpolation model selected for the blood curves involved in the calculation of the input curve.

3. The ratio of the two AUCs as a function of time
   \[ AUC_{ratio}(t) = \frac{\int_0^t C_T(\tau) d\tau}{\int_0^t C_p(\tau) d\tau} \]
   It provides an indication of the tissue accumulation of the tracer [1].

As the ratio curve has a completely different scaling than the curve integrals, it appears along the zero line. It can be shown with appropriate scaling by hiding all other curves. The easiest way to export the values is using the Save all curves or the View values option in the context menu (right click into curve area).

**Reference**

Under equilibrium conditions this ratio is equal to the total distribution volume of tracer in tissue. Therefore, the plot of the Tissue/Input curve Ratio against time is a quick means to check whether the system has equilibrated (plot becomes constant) or not.

Note that an explicit blood volume correction is available via the \( vB \) input parameter. In the case of \( vB=0 \) (default setting), no correction is performed. For \( vB>0 \), the tissue TAC is corrected by the scaled whole blood activity before the actual analysis as follows.

\[
C_T(t) = \frac{C_{PET}(t) - vBC_b(t)}{1-vB}
\]

### 6.5.15 Tissue Ratio Methods

This auxiliary model is aimed at receptor tracers for which a reference tissue without specific binding is available. It allows getting a quick estimate of the binding potential \( BP_{ND} \) using two different methods as described by Ito et al. [1].

At the time, when specific binding \( C_2 \) peaks, its derivative equals zero (transient equilibrium) and \( BP_{ND} = \frac{k_3}{k_4} = C_2/C_1 \) (non-specific/specific binding).

In practice, however, neither the \( C_1 \) part nor the \( C_2 \) part of a measured tissue TAC are known. As a pragmatic solution it is assumed that the non-displaceable compartment is equal among tissues, so the \( C_T'(t) \) of reference tissue without specific binding is regarded as \( C_1' \), and \( C_2 \) is calculated by \( C_T(t) - C_T'(t) \).

**Operational Model Curve**

Two different ratios are calculated by the Tissue Ratio Methods model:

\[
BP_{ND} = \frac{k_3}{k_4} = \frac{C_2(t_{\text{trans.equ.}})}{C_1(t_{\text{trans.equ.}})} = \frac{C_T(t_{\text{trans.equ.}}) - C_T'(t_{\text{trans.equ.}})}{C_T'(t_{\text{trans.equ.}})}
\]
This ratio is calculated for all times, but it is only valid at the time of transient equilibrium. There is no automatic routine for detecting this peak. Rather, the user has to find this time by looking at the difference curve (enable check box).

The second ratio calculated is called the interval method in [1].

\[ BP_{ND} = \frac{k_4}{k_4} = \frac{\int_{t_b}^{t_e} C_2(t) dt}{\int_{t_b}^{t_e} C_1(t) dt} \]

Here, the two curves are integrated in a time interval \([t_b, t_e]\) which must include the time of transient equilibrium \((t_b=9\text{ min}, t_e=45 \text{ min})\) for raclopride, where the time of transient equilibrium is in the range 20 to 24 min [1]).

Implementation
1. After switching to Tissue Ratio Methods a suitable reference tissue without specific binding must be selected.
2. The integration start time of the interval method must be specified by the user as the input parameter Integral start time.
3. The ratios are calculated for all times (for the integral method only after the Integral start time) and presented as curves. These curves can easily be exported as values using the save button [2].

Reference

6.5.16 Tracer Retention Fraction

The Retention Fraction is the fraction of the total tracer delivered to an organ that is extracted into and retained by the tissue. It is the residual after clearances of the vascular component and the portion of the tracer that rapidly back diffuses from tissue to blood. Usually this fraction of tracer is sequestered in more slowly turning over metabolic or membrane binding processes. [1]

Operational Equation
Given a tissue TAC \(C_t(t)\) and the input curve \(C_p(t)\) this auxiliary model allows calculating the retention fraction \(R\) by [2]:

\[ R(t) = \frac{\int_0^t C_t(t) dt}{\int_0^t C_p(t) dt} \]
The denominator with the input curve integrated from the time of injection represents the available tracer, the numerator with the integral of the late tissue uptake the actually extracted tracer. Other authors [3] have used a simplified version of tracer retention which is calculated by dividing the tracer concentration by the input curve integral.

Note that an explicit blood volume correction is available via the \( vB \) input parameter. In the case of \( vB=0 \) (default setting), no correction is performed. For \( vB>0 \), the tissue TAC is corrected by the scaled whole blood activity before the actual analysis as follows.

\[
C_T(t) = \frac{C_{PET}(t) - vB \cdot C_B(t)}{1 - vB}
\]

### Implementation

The Tracer Retention auxiliary model has two input parameters **Start** and **End** for the specification of \( t_b \) and \( t_e \) in the integration formula above, respectively. The curve display shows the input curve interpolated at the frame mid-times. The result parameters are calculated in % and %/min for the two calculation methods:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>( R_{\text{window}} ) [%]</td>
<td>Retention of tracer in the time window (Tissue-Integral in window)/(Plasma-Integral from start)</td>
</tr>
<tr>
<td>( R_{\text{wind_min}} ) [%/min]</td>
<td>Retention of tracer in the time window (Tissue-Average in window)/(Plasma-Integral from start)</td>
</tr>
<tr>
<td>( R_{\text{last}} ) [%]</td>
<td>Retention of tracer in the last frame of the acquisition (Tissue-Integral in window)/(Plasma-Integral from start)</td>
</tr>
<tr>
<td>( R_{\text{last_min}} ) [%/min]</td>
<td>Retention of tracer in the last frame of the acquisition (Tissue-Average in window)/(Plasma-Integral from start)</td>
</tr>
</tbody>
</table>

### References


### 6.5.17 Fractal Dimension

The Fractal Dimension model measures the complexity of a 2-dimensional structure by calculating its box-counting dimension [1]. This concept has been applied in oncologic studies for assessing the heterogeneity of tissue kinetics [2].

#### Operational Equation

The idea is to subdivide the area under the tissue TAC into a number of square boxes and simply count the number of boxes containing some part of the structure. The mesh size is defined as \( s \), so \( 1/s \) gives the number of segments in each of the 2 dimensions. For instance, specifying \( 1/s=5 \) therefore means a subdivision into \( 5*5=25 \) boxes.
The counting process is performed with increasing number of intervals up to the specified \( 1/s \). Next, the data are plotted in a double-logarithmic way, namely \( \log(N(s)) \) on the y axis and \( \log(1/s) \) on the x-axis.

The box-counting dimension is finally obtained as the slope of a linear regression through the plotted points.

**Implementation**

After switching to the Fractal dimension model, two input parameters are available for specifying the box-counting process: \( 1/s \), and Maximal value the highest TAC value which might occur in the data.

**References**


### 6.6 Interpolation Functions for Whole-Blood and Plasma Activity

PKIN supports multiple functions which can be fitted to measured blood activity data and allow interpolating intermediate values. Exceptions are the functions for the dispersion deconvolution, as described above.

#### 6.6.1 Linear Activity Interpolation

The blood and plasma activity concentrations require interpolation procedures for kinetic modeling, as described above.

**Operational Model Curve**

The Lin. Interpolation model interpolates linearly between the measured activity values. Outside the measured values the following interpolation rules apply:

- If the first measurements occurs at \( t<0 \), the activity is assumed to be zero for all earlier times.
If the first measurements occurs at \( t>0 \), the activity is assumed to be zero for \( t<=0 \). Activity values from \( t=0 \) up to the first measurement are calculated by linear interpolation from activity 0 at \( t=0 \).

The activity is assumed to remain constant after the last measurement. For short blood curves it is therefore recommended to replace the tail of the measurements by a fitted \textit{tri-exponential} model.

Parameter Fitting

The \textit{Lin. Interpolation} model has no fitting parameter.

The \textit{Delay} parameter serves for correcting a timing offset between tissue and blood data. Positive delays correspond to delayed blood information and hence shift the blood curve to earlier times (to the left). The \textit{Delay} is only relevant during the fitting of the tissue model where it can be fitted as an additional parameter to the parameters of the kinetic model.

6.6.2 Decaying Exponentials

The blood and plasma activity concentrations require interpolation procedures for kinetic modeling, as described above.

Operational Model Curve

The \textit{3 Exponentials} model allows replacing the tail of the blood activity curve by a fitted sum of up to three exponentials. The \textit{Begin} parameter defines at what time the models switches from linear to tri-exponential interpolation. The exponentials are each defined by an \textit{Amplitude} and a \textit{Halftime} [min] of the decay. The number of exponentials can easily be reduced by fixing one or two of the amplitudes at a value of 0.

\[
f_{\text{Plasma/Pl}}(t) = \begin{cases} 
\text{Lin.Interpol} & \text{if } t < \text{Begin} \\
\sum_{i=1}^{3} A_i e^{-\ln 2 T_i} & \text{if } t \geq \text{Begin}, u = t - \text{Begin}
\end{cases}
\]
Parameter Fitting

Per default only two exponentials are enabled, and the Begin is initialized to the time of the blood peak. Use the Fit plasma activity or Fit whole blood button to fit the two exponentials to the measurements after the specified Begin. If the shape cannot be described well enough, enable the third exponential by enabling the fit boxes and set Amplitude 3 to a positive number, then Fit again. Possibly, Begin has to be set to a later time for obtaining the signal tail.

The Delay parameter serves for correcting a timing offset between tissue and blood data. Positive delays correspond to delayed blood information and hence shift the blood curve to earlier times (to the left). The Delay is only relevant during the fitting of the tissue model where it can be fitted as an additional parameter to the parameters of the kinetic model.

6.6.3 Compartment Model with 3 Eigenvalues

The blood and plasma activity concentrations require interpolation procedures for kinetic modeling, as described above.

Operational Model Curve

The Compart, 3 EV model has been developed for the FDG tracer [1]. It is the result from modeling the distribution and delivery of FDG in the circulatory system by a compartment model and results in an analytical solution with 3 eigenvalues (EV):

\[
C_{\text{continuous}}(t) = \begin{cases} 0 & t < \text{Begin} \\ \left[A_1 u - A_2 - A_3 \right] e^{\ln 2 / T_1} + A_2 e^{\ln 2 / T_2} + A_3 e^{\ln 2 / T_3} & t \geq \text{Begin}, u = t - \text{Begin} \end{cases}
\]
Parameter Fitting
Similar to the 3 Exponentials model it includes a Begin parameter and three Amplitudes ($A_1$, $A_2$, $A_3$) and a Halftimes ($T_1$, $T_2$, $T_3$). However, the functional form supports a peak rising from time Begin followed by an exponentially decreasing tail. Use the Fit plasma activity or Fit whole blood button to fit the operational model curve to the measurements. Note that all samples are considered, not only the ones after Begin.

The form of the curve reacts very sensitively to changes of the parameters. The example above illustrates a peak which could not fully reach the maximum value. This model may be more suited for simulations rather than actual data processing.

The Delay parameter serves for correcting a timing offset between tissue and blood data. Positive delays correspond to delayed blood information and hence shift the blood curve to earlier times (to the left). The Delay is only relevant during the fitting of the tissue model where it can be fitted as an additional parameter to the parameters of the kinetic model.

Reference

6.6.4 Compartment Model with 2 Eigenvalues
The blood and plasma activity concentrations require interpolation procedures for kinetic modeling, as described above.

Operational Model Curve
The Compart, 2 EV model is an alternative model to the Compart, 3 EV model for the plasma activity curve of FDG [1]. It assumes repeated eigenvalues (EV) and has the following functional form:

$$C_{\text{continuous}}(t) = \begin{cases} 0 & t < \text{Begin} \\ [A_1 u - A_2] e^{-\ln 2 / T_1} + A_2 e^{-\ln 2 / T_2} & t \geq \text{Begin}, u = t - \text{Begin} \end{cases}$$

Parameter Fitting
Similar to the 3 Exponentials model it includes a Begin parameter and two Amplitudes ($A_1$, $A_2$) and a Halftimes ($T_1$, $T_2$). However, the functional form supports a peak rising from time Begin followed by an exponentially decreasing tail. Use the Fit plasma activity or Fit whole blood button to fit the operational model curve to the measurements. Note that all samples are considered, not only the ones after Begin.
The form of the curve reacts very sensitively to changes of the parameters. The example above illustrates a result is model may be more suited for simulations rather than actual data processing.

The **Delay** parameter serves for correcting a timing offset between tissue and blood data. Positive delays correspond to delayed blood information and hence shift the blood curve to earlier times (to the left). The **Delay** is only relevant during the fitting of the tissue model where it can be fitted as an additional parameter to the parameters of the kinetic model.

**Reference**


### 6.6.5 Gamma Function

The blood and plasma activity concentrations require interpolation procedures for kinetic modeling, as described above.

**Operational Model Curve**

The **Gamma Function** model is defined by the following functional form:

\[
C_{\text{continuous}}(t) = \begin{cases} 
0 & t < \text{Begin} \\
A u^B e^{-uC} & t \geq \text{Begin}, u = t - \text{Begin}
\end{cases}
\]

describing a sharp peak governed followed by a quick decay towards zero.

**Parameter Fitting**

The **Gamma Function** model includes a **Begin** time parameter for positioning the peak. The factor **A** scales the curve, the exponent **B** governs the rise and constant **C** the decay towards zero. Use the **Fit plasma activity** or **Fit whole blood** button to fit the operational model curve to the measurements. Note that all samples are considered, not only the ones after **Begin**.

The form of the curve reacts very sensitively to changes of the parameters. The example above illustrates the inability to fit model a peak followed by a moderate to slow decay appropriately. This model may therefore be more suited for simulations rather than actual data processing.

The **Delay** parameter serves for correcting a timing offset between tissue and blood data. Positive delays correspond to delayed blood information and hence shift the blood curve to earlier times (to the left). The **Delay** is only relevant during the fitting of the tissue model where it can be fitted as an additional parameter to the parameters of the kinetic model.
6.6.6 Gamma Function plus two Decaying Exponentials

The blood and plasma activity concentrations require interpolation procedures for kinetic modeling, as described above. 

Operational Model Curve

This Gamma + 2 Exponentials model is very similar to the Compartment Model with 3 Eigenvalues. However, it includes a power term \( u^B \) which allows better modeling of sharp peaks.

\[
C_{\text{continuous}}(t) = \begin{cases} 
A_1 u^B - A_2 - A_3 e^{-\frac{ln 2}{T_1}} + A_2 e^{-\frac{ln 2}{T_2}} + A_3 e^{-\frac{ln 2}{T_3}} & t < \text{Begin} \\
0 & t \geq \text{Begin}, u = t - \text{Begin} 
\end{cases}
\]

Parameter Fitting

The Gamma + 2 Exponentials model includes a Begin parameter, three Amplitudes (\( A_1, A_2, A_3 \)) and a Halftime (\( T_1, T_2, T_3 \)), and a Power Exponent (\( B \)). This functional form supports a peak rising from time Begin followed by an exponentially decreasing tail. The initial default of the Power Exponent is a fixed value of 1 which corresponds to the Compartment Model with 3 Eigenvalues. Use the Fit plasma activity or Fit whole blood button to fit the operational model curve to the measurements. Then, for better fitting of the peak, enable the Power Exponent and fit again. The form of the curve reacts very sensitively to changes of the parameters. Note that all samples are considered, not only the ones after Begin.

The Delay parameter serves for correcting a timing offset between tissue and blood data. Positive delays correspond to delayed blood information and hence shift the blood curve to earlier times (to the left). The Delay is only relevant during the fitting of the tissue model where it can be fitted as an additional parameter to the parameters of the kinetic model.

6.6.7 Simulation of Bolus/Infusion Tracer Application

When a compartment system has equilibrated, the total distribution volume can be calculated easily as the ratio of tracer concentration in tissue to the metabolite-corrected plasma concentration with a single static scan. It has been found that the time required to reach equilibrium can be shortened by an optimized tracer delivery. A setup which does not require a sophisticated tracer delivery system is to apply an initial bolus and continue with a constant infusion.

Carson et al. [1] have developed a method to optimize the ratio between the activity given as the initial bolus and the activity level of the subsequent infusion for quickly reaching an equilibrium (Appendix B in [1]). This method is implemented as a tissue model in PKIN. The current blood model Bolus/Infusion is only a visualization model and should not be used for fitting measured blood data. It allows entering the \( K_{bol} \) value found with the tissue model optimization and inspecting the plasma activity curve under the bolus/infusion regime.
Operational Model Curve

For a bolus and infusion (B/I) experiment the expected input function can be derived from the input curve \(C_p(t)\) of a bolus-only experiment by

\[
C_{\text{Model}}(t) = \frac{K_{\text{bol}} C_p(t) + \int_0^t C_p(\tau) d\tau}{K_{\text{bol}} + T}
\]

\(K_{\text{bol}}\) [min] defines the dose of the bolus expressed as minutes of infusion, and \(T\) is the total duration of the infusion (end of last frame).

Parameter Fitting

This model cannot be used for data fitting. It assumes the plasma activity curve of a bolus experiment has been loaded and visualizes the expected input curve if \(K_{\text{bol}}\) [min] equivalents of the infusion are applied as an initial bolus. The criterion of fast equilibration is that the model curve gets soon constant and remains so.

Reference


6.6.8 Flumazenil, HOT Input Curve

The Multiinjection, HOT model is used in combination with the tissue model for multi-injection studies with \(^{11}\text{C}\)-Flumazenil as described by Delforge et al. [1]. Please refer to the documentation of the implemented tissue model for a more detailed description.

It is assumed that the total plasma activity of the tracer \(C_{\text{Plasma}}(t)\) is loaded from the menu as illustrated below.

Operational Model Curve

The input curve is then calculated by applying an exponential metabolite correction which considers the two injections of hot ligand at time \(T_1\) and \(T_3\):

\[
C_p(t) = \begin{cases} 
C_{\text{Plasma}}(A_1 e^{B_1(t-T_1)} + A_2 e^{B_2(t-T_2)}) & T_1 < t < T_3 \quad \text{decay since 1st injection} \\
C_{\text{Plasma}}(A_1 e^{B_1(t-T_1)} + A_2 e^{B_2(t-T_2)}) & t > T_3 \quad \text{decay since 3rd injection}
\end{cases}
\]
Parameter Fitting

Note that although the corresponding parameters have a fit box, they should normally be used for entering pre-determined values of injection times and metabolite correction parameters. The default parameters of the for the bi-exponential metabolite correction ($A_1$, $T_1$, $A_2$, $B_2$) used in this model are from [1].

Reference


6.6.9 Flumazenil, COLD Input Curve

The Multiinjection, COLD model is used in combination with the tissue model for multi-injection studies with $^{11}$C-Flumazenil as described by Delforge et al. [1]. Please refer to the documentation of the implemented tissue model for a more detailed description.

It is assumed that the total plasma activity of the labeled tracer $C_{\text{Plasma}}(t)$ is loaded from the menu as illustrated below.

Operational Model Curve

The plasma activity of unlabeled (cold) ligand is calculated by using the information of the labeled (hot) ligand, scaling it proportionally to the applied dose, and applying the same metabolite correction. The metabolite corrected plasma activity of the labeled compound $C_p(t)$ is calculated by

$$C_p(t) = \begin{cases} C_{\text{Plasma}} \left( A_1 e^{B_1(t-T_1)} + A_2 e^{B_2(t-T_2)} \right) & T_1 < t < T_3 \quad \text{decay since 1\textsuperscript{st} injection} \\ C_{\text{Plasma}} \left( A_1 e^{B_1(t-T_1)} + A_2 e^{B_2(t-T_2)} \right) & t > T_3 \quad \text{decay since 3\textsuperscript{rd} injection} \end{cases}$$

The calculation of the plasma activity of the unlabeled ligand $C_u(t)$ is also performed differently in two time segments:
\[ C_{mP}(t) = \begin{cases} \frac{Dose_p^{\text{Cold}}}{Dose_p^{\text{Hot}}} C_p(t - T_2^*) & T_2^* < t < T_3^* \\ \frac{Dose_p^{\text{Cold}}}{Dose_p^{\text{Hot}}} C_p(t - T_3^*) + C_{mP}(T_3^*) & t > T_3^* \end{cases} \]

1. Following the cold-only injection at \( T_2 \) the input curve of the first hot injection at \( T_1 \) is scaled according to the dose ratio and shifted to start at \( T_2^* \).

2. Following the injection at \( T_3 \) with hot and cold ligand the input curve of the hot ligand is scaled, and a contribution of the first cold injection added which is decreasing by the metabolite build-up.

Parameter Fitting

Note that although the corresponding parameters have a fit box, they should normally be used for entering pre-determined values of injection times and metabolite correction parameters. The default parameters of the for the bi-exponential metabolite correction (\( A_1, T_1, A_2, B_2 \)) used in this model are from [1].

Reference

6.7 Interpolation Functions for Plasma/Whole-Blood Fraction

The following functions for converting whole-blood activity into plasma activity are available in PKIN. Note that it is feasible to combine the conversion into plasma activity and then into authentic tracer activity into a single conversion function. In this case the parent fraction can remain the identity function, which is its default.

6.7.1 Fix Plasma Fraction

The sequential steps for converting blood measurements to an input curve are explained in a dedicated section. The current plasma fraction model serves for converting a whole-blood activity curve into a plasma-activity curve.

It is assumed that a whole-blood activity curve has been loaded with Load Whole Blood Activity entry in the menu.

With the Fix model the activity concentration in plasma is assumed to be proportional to the activity concentration in whole blood. The proportionality constant Fraction has to be specified by the user and has a default of 1.

Note that as long as no other plasma-related data is loaded or the plasma fraction changed, the Fix model is applied so that the whole-blood activity curve will be used as the input curve.

Parameter Fitting
There is nothing to fit with the Fix model.

6.7.2 Linear Interpolation of Plasma/Whole-blood Fraction

The sequential steps for converting blood measurements to an input curve are explained in a dedicated section. The current plasma fraction model serves for converting a whole-blood activity curve into a plasma-activity curve.

It is assumed that a curve representing the measured fractions of plasma/whole-blood activity has been loaded with Load Plasma/WB Fraction.

The Lin. Interpolation model interpolates linearly between the measured fraction values. Outside the measured values the following interpolation rules apply:

- The fraction at time t=0 is assumed to equal 1.
- The fraction is assumed to remain constant after the last measured fraction.

For incomplete fraction data it is therefore recommended to replace the tail of the measurements by a fitted tri-exponential model.
Parameter Fitting
There is nothing to fit with the Lin. Interpolation model.

6.7.3 Exponential Plasma/Whole-blood Fraction

The sequential steps for converting blood measurements to an input curve are explained in a dedicated section. The current plasma fraction model serves for converting a whole-blood activity curve into a plasma-activity curve.

It is assumed that a curve representing the measured fractions of plasma/whole-blood activity has been loaded with Load Plasma/WB Fraction.

Operational Model Curve
The 3 Exponentials model allows replacing the tail of the fraction curve by a fitted sum of up to three exponentials. The Begin parameter defines at what time the model switches from linear to tri-exponential interpolation. The exponentials are each defined by an Amplitude and a Halftime [min] of the decay. The number of exponentials can easily be reduced by fixing one or two of the amplitudes at a value of 0.

\[
f_{\text{Plasma/WB}}(t) = \begin{cases} 
\text{Lin.Interpol} & t < \text{Begin} \\
\sum_{i=1}^{3} A_i e^{-\ln(2)/T_i} & t \geq \text{Begin}, u = t - \text{Begin} 
\end{cases}
\]

The illustration below shows an example with the exponentials fitted following 2 minutes (Begin = 120sec). Note that the linear interpolation assumes a fraction of 1 at t=0.
Parameter Fitting

The model supports the fitting of the parameters **Begin**, **Amplitude 1, 2, 3** ($=A_1, A_2, A_3$) and **Halftime 1, 2, 3** ($=T_1, T_2, T_3$). Fitting is started with the **Fit plasma fraction** button, which only takes the samples after the **Begin** time into consideration.

Appropriate default parameters for FDG scans:

- Rat experiments [1]: Plasma/whole_blood(t) = 0.51 e^{-(\ln 2/4.79)t} + 0.3 e^{-(\ln 2/337)t} + 0.8
- Mouse Experiments [2]: Plasma/whole_blood(t) = 0.386e^{-(0.191t)} + 1.165

Use without Measured Fraction

If no measured plasma/whole-blood fraction data is available, the **3 Exponentials** model can still be applied. In this case the **Begin** time is disregarded, and the functional defined by the entered parameters applied.

References


6.7.4 Hill Function

The sequential steps for converting blood measurements to an input curve are explained in a dedicated section. The current plasma fraction model serves for converting a whole-blood activity curve into a plasma-activity curve.
It is assumed that a curve representing the measured fractions of plasma/whole-blood activity has been loaded with **Load Plasma/WB Fraction**.

**Operational Model Curve**

The **Hill** model allows replacing the tail of the fraction curve by an analytical function. The **Begin** parameter defines at what time the models switches from linear interpolation to the Hill function.

\[
f_{\text{Plasma/WB}}(t) = \begin{cases} 
\text{Lin.Interpol} & t \leq \text{Begin} \\
\text{scale} \cdot \left(1 - \frac{Au^B}{u^B + C}\right) & t > \text{Begin}, u = t - \text{Begin} 
\end{cases}
\]

The illustration below shows an example with the Hill function fitted following 2 minutes (**Begin** = 120sec). The the weighting has been changed for putting more emphasis on the tail. Note that the linear interpolation assumes a fraction of 1 at **t=0**.

**Parameter Fitting**

The model supports the fitting of the parameters **Begin**, **scale**, **A**, **B**, **C**. Fitting is started with the **Fit plasma fraction** button, which only takes the samples after the **Begin** time into consideration.

**Use without Measured Fraction**

If no measured plasma/whole-blood fraction data is available, the **Hill** model can still be applied. In this case the **Begin** time is disregarded, and the functional defined by the entered parameters applied.

**References**


**6.7.5 Watabe Function**

The sequential steps for converting blood measurements to an input curve are explained in a dedicated section. The current plasma fraction model serves for converting a whole-blood activity curve into a plasma-activity curve.

It is assumed that a curve representing the measured fractions of plasma/whole-blood activity has been loaded with **Load Plasma/WB Fraction**.
Operational Model Curve

The Watabe model allows replacing the tail of the fraction curve by an analytical function. The Begin parameter defines at what time the models switches from linear interpolation to the Hill function.

\[
 f_{\text{Plasma/WB}}(t) = \begin{cases} 
 \text{Lin.Interpol} & t \leq \text{Begin} \\
 \frac{\text{scale}}{1 + (A u)^B} & t > \text{Begin}, u = t - \text{Begin} 
\end{cases}
\]

The illustration below shows an example with the Hill function fitted following 2 minutes (Begin = 120 sec). The weighting has been changed for putting more emphasis on the tail. Note that the linear interpolation assumes a fraction of 1 at t=0.

Parameter Fitting

The model supports the fitting of the parameters Begin, scale, A, B, C. Fitting is started with the Fit plasma fraction button, which only takes the samples after the Begin time into consideration.

Use without Measured Fraction

If no measured plasma/whole-blood fraction data is available, the Watabe model can still be applied. In this case the Begin time is disregarded, and the functional defined by the entered parameters applied.

References


6.7.6 Sigmoid Function

The sequential steps for converting blood measurements to an input curve are explained in a dedicated section. The current plasma fraction model serves for converting a whole-blood activity curve into a plasma-activity curve.

It is assumed that a curve representing the measured fractions of plasma/whole-blood activity has been loaded with Load Plasma/WB Fraction.
Operational Model Curve

The Sigmoid model allows replacing the tail of the fraction curve by an analytical function. The Begin parameter defines at what time the models switches from linear interpolation to the Hill function.

\[ f_{\text{Plasma/Whole Blood}}(t) = \begin{cases} \text{Lin.Interpol} & t \leq \text{Begin} \\ \text{scale} \left[ 1 - \frac{u^3}{u^3 + 10^4} \right]^B + C & t > \text{Begin}, u = t - \text{Begin} \end{cases} \]

The illustration below shows an example with the Hill function fitted following 2 minutes (\(\text{Begin} = 120\) sec). The the weighting has been changed for putting more emphasis on the tail. Note that the linear interpolation assumes a fraction of 1 at \(t=0\).

Parameter Fitting

The model supports the fitting of the parameters Begin, scale, A, B, C. Fitting is started with the Fit plasma fraction button, which only takes the samples after the Begin time into consideration.

Use without Measured Fraction

If no measured plasma/whole-blood fraction data is available, the Sigmoid model can still be applied. In this case the Begin time is disregarded, and the functional defined by the entered parameters applied.

References


6.7.7 Linear Function

The sequential steps for converting blood measurements to an input curve are explained in a dedicated section. The current plasma fraction model serves for converting a whole-blood activity curve into a plasma-activity curve.
It is assumed that a curve representing the measured fractions of plasma/whole-blood activity has been loaded with **Load Plasma/WB Fraction**.

The **Linear** model provides a linear function with **Slope** and **Intercept** parameters which can be fitted to the plasma fraction values.

### 6.8 Interpolation Functions for Parent/Plasma Fraction

The following parent fraction models are available in PKIN.

#### 6.8.1 Fix Parent/Plasma Fraction

The sequential steps for converting blood measurements to an input curve are explained in a dedicated section. The current **parent fraction** model serves for converting a plasma activity curve into an input curve (authentic, unchanged tracer in plasma).

**Operational Model Curve**

The **Fix** model assumes that the concentration of authentic tracer in plasma is in constant proportion to the concentration of total tracer in plasma. The fraction $f_{\text{parent}}(t)$ of authentic tracer to total tracer in plasma is therefore given by

$$f_{\text{parent}}(t) = f_p$$

with $f_p$ representing the free fraction of authentic tracer in plasma, i.e. the fraction of tracer not bound to plasma proteins.

**Note:** The **Fix** model is the default as soon as blood data is loaded. This is adequate if a metabolite correction is not necessary or has been performed in a prior step, for instance externally before loading the plasma activity.

**Parameter Fitting**

There is nothing to fit with the **Lin. Interpolation** model. The only input parameter is the free fraction $f_p$ which defaults to 1, as it is experimentally difficult to measure.
6.8.2 Linear Interpolation Fraction

The sequential steps for converting blood measurements to an input curve are explained in a dedicated section\(^{12}\). The current parent fraction\(^{12}\) model serves for converting a plasma activity curve into an input curve (authentic, unchanged tracer in plasma).

The Lin. Interpolation model assumes that the concentration ratio of authentic tracer to total tracer in plasma has been determined at certain times during the acquisition and loaded with the Load Fraction entry in the menu.

Operational Model Curve

The Lin. Interpolation model interpolates linearly between the measured fraction values \(F_k\).

\[
f_{\text{parex}}(t) = f_p \cdot \text{Interpol}(F_k)
\]

and multiplies with \(f_p\) representing the free fraction of authentic tracer in plasma, i.e. the fraction of tracer not bound to plasma proteins.

Outside the measured values the following interpolation rules apply:

- The fraction at time \(t=0\) is assumed to equal 1.
- The fraction is assumed to remain constant after the last measured fraction. For incomplete fraction data it is therefore recommended to replace the tail of the measurements by a fitted tri-exponential\(^{12}\) model.

Note that fraction values \(>1\) are not allowed.

Parameter Fitting

There is nothing to fit with the Lin. Interpolation model. The only input parameter is the free fraction \(f_p\) which defaults to 1, as it is experimentally difficult to measure.

6.8.3 1-Exponential

The sequential steps for converting blood measurements to an input curve are explained in a dedicated section\(^{12}\). The current parent fraction\(^{12}\) model serves for converting a plasma activity curve into an input curve (authentic, unchanged tracer in plasma).
The 1-Exponential model assumes that the concentration ratio of authentic tracer to total tracer in plasma has been determined at certain times during the acquisition and loaded with the Load Fraction entry in the menu.

**Operational Model Curve**

The 1-Exponential model applied by Wu et al [1] for modeling of the parent fraction has the following functional form

\[
\begin{align*}
    f_{\text{parent}}(t) &= f_p \left\{ \begin{array}{ll}
        1 & \mbox{if } t \leq \text{Begin} \\
        A e^{-u t / T} + (1 - A) & \mbox{if } t > \text{Begin}, u = t - \text{Begin}
    \end{array} \right.
\end{align*}
\]

The calculated fraction is multiplied with \(f_p\), the free fraction of authentic tracer in plasma, i.e. the fraction of tracer not bound to plasma proteins.

The fraction calculation assumes negligible metabolites before a Begin time, and therefore remains constant at 1. From Begin on it decreases exponentially towards an equilibration fraction (1-A) with halftime \(T\).

**Parameter Fitting**

The free fraction \(f_p\) is an input parameter which has no impact on fitting the model curve to the fraction data. The default for \(f_p\) is 1 as it is experimentally difficult to measure.

The model supports the fitting of the parameters Begin, Scale (=A) and Halftime. All data samples are considered in the fitting process irrespective of Begin. The Scale parameter is restricted to the range [0,1] per default.

**Use without Measured Parent Fraction**

The 1-Exponential parent fraction model can be applied even if no parent fraction measurements were loaded. In this case the user has to specify a parameter set which establishes a representative metabolite correction for the used tracer.

**Reference**

6.8.4 2-Exponentials

The sequential steps for converting blood measurements to an input curve are explained in a dedicated section. The current parent fraction model serves for converting a plasma activity curve into an input curve (authentic, unchanged tracer in plasma).

The 2-Exponential model assumes that the concentration ratio of authentic tracer to total tracer in plasma has been determined at certain times during the acquisition and loaded with the Load Fraction entry in the menu.

Operational Model Curve

The 2-Exponential model applied by Wu et al [1] for modeling of the parent fraction has the following functional form

\[ f_{\text{parent}}(t) = \begin{cases} \frac{1}{f_p} & t \leq \text{Begin} \\ A e^{-\ln 2/t_1} + (1-A) e^{-\ln 2/t_2} & t > \text{Begin}, u = t - \text{Begin} \end{cases} \]

The calculated fraction is multiplied with \( f_p \), the free fraction of authentic tracer in plasma, i.e. the fraction of tracer not bound to plasma proteins.

The fraction calculation assumes negligible metabolites before a Begin time, and therefore remains constant at 1. From Begin on it decreases as a sum of two exponentials with halftimes \( T_1 \) and \( T_2 \). Note that because only one scale factor A is used, continuity at time Begin is ensured.

Parameter Fitting

The free fraction \( f_p \) is an input parameter which has no impact on fitting the model curve to the fraction data. The default for \( f_p \) is 1 as it is experimentally difficult to measure.

The model supports the fitting of the parameters Begin, Scale (=A), Halftime 1 and Halftime 2. All data samples are considered in the fitting process irrespective of Begin. The Scale parameter is restricted to the range \([0,1]\) per default.

Use without Measured Parent Fraction

The 2-Exponential parent fraction model can be applied even if no parent fraction measurements were loaded. In this case the user has to specify a parameter set which establishes a representative metabolite correction for the used tracer.
Reference

6.8.5 3-Exponentials

The sequential steps for converting blood measurements to an input curve are explained in a dedicated section. The current parent fraction model serves for converting a plasma activity curve into an input curve (authentic, unchanged tracer in plasma).

The 3-Exponential model assumes that the concentration ratio of authentic tracer to total tracer in plasma has been determined at certain times during the acquisition and loaded with the Load Fraction entry in the menu.

Operational Model Curve

The general 3-Exponential model for the parent fraction has the following functional form

\[ f_{\text{parent}}(t) = f_p \begin{cases} A_1 e^{-\ln 2 \cdot T_1} & t \leq \text{Begin} \\ A_2 e^{-\ln 2 \cdot T_2} + A_3 e^{-\ln 2 \cdot T_3} & t > \text{Begin}, u = t - \text{Begin} \end{cases} \]

The calculated fraction is multiplied with \( f_p \), the free fraction of authentic tracer in plasma, i.e. the fraction of tracer not bound to plasma proteins.

The fraction calculation assumes negligible metabolites before a Begin time, and therefore remains constant at 1. From Begin on it decreases as a sum of three exponentials with halftimes \( T_1, T_2 \) and \( T_3 \). Note that the calculated fractions are restricted to the physiologic \([0,1]\) value range.

Parameter Fitting

The free fraction \( f_p \) is an input parameter which has no impact on fitting the model curve to the fraction data. The default for \( f_p \) is 1 as it is experimentally difficult to measure.

The model supports the fitting of the parameters Begin, Amplitude 1, 2, 3 (=A_1, A_2, A_3) and Halftime 1, 2, 3 (=T_1, T_2, T_3). All data samples are considered in the fitting process irrespective of Begin. The three Amplitude parameters are restricted to the range \([0,1]\) per default.

Use without Measured Parent Fraction
The 3-Exponential parent fraction model can be applied even if no parent fraction measurements were loaded. In this case the user has to specify a parameter set which establishes a representative metabolite correction for the used tracer.

### 6.8.6 Hill Function

The sequential steps for converting blood measurements to an input curve are explained in a dedicated section. The current parent fraction model serves for converting a plasma activity curve into an input curve (authentic, unchanged tracer in plasma).

The Hill model assumes that the concentration ratio of authentic tracer to total tracer in plasma has been determined at certain times during the acquisition and loaded with the Load Fraction entry in the menu.

**Operational Model Curve**

The Hill model applied by Gunn et al [1] for modeling of the parent fraction has the following functional form

\[
 f_{\text{parent}}(t) = \begin{cases} 
 f_p & t \leq \text{Begin} \\
 1 - \frac{A t^B}{t^B + C} & t > \text{Begin}, t = t - \text{Begin} 
\end{cases}
\]

whereby the calculated fraction is multiplied with \( f_p \), the free fraction of authentic tracer in plasma, i.e. the fraction of tracer not bound to plasma proteins.

The fraction calculation assumes negligible metabolites before a Begin time, and therefore remains constant at 1. From Begin on it decreases smoothly.

**Parameter Fitting**

The free fraction \( f_p \) is an input parameter which has no impact on fitting the Hill curve to the fraction data. The default for \( f_p \) is 1 as it is experimentally difficult to measure.

The model supports the fitting of the parameters Begin, A, B and C. All data samples are considered in the fitting process irrespective of Begin. The scaling parameter A is restricted to the range [0,1] per default.

**Use without Measured Parent Fraction**
The Hill parent fraction model can be applied even if no parent fraction measurements were loaded. In this case the user has to specify a parameter set which establishes a representative metabolite correction for the used tracer.

Reference

6.8.7 Watabe Function

The sequential steps for converting blood measurements to an input curve are explained in a dedicated section. The current parent fraction model serves for converting a plasma activity curve into an input curve (authentic, unchanged tracer in plasma).

The Watabe model assumes that the concentration ratio of authentic tracer to total tracer in plasma has been determined at certain times during the acquisition and loaded with the Load Fraction entry in the menu.

Operational Model Curve

The Watabe model applied by Watabe et al [1] for modeling of the parent fraction has the following functional form

\[ f_{\text{parent}}(t) = f_p \begin{cases} \frac{1}{1 + \left(\frac{A_u}{B}\right)^8} & t \leq \text{Begin} \\ \frac{u - \text{Begin}}{t > \text{Begin}, u = t - \text{Begin}} \end{cases} \]

with B set to 2. The calculated fraction is multiplied with \( f_p \), the free fraction of authentic tracer in plasma, i.e. the fraction of tracer not bound to plasma proteins.

The fraction calculation assumes negligible metabolites before a Begin time, and therefore remains constant at 1. From Begin on it decreases smoothly.

Parameter Fitting

The free fraction \( f_p \) is an input parameter which has no impact on fitting the model curve to the fraction data. The default for \( f_p \) is 1 as it is experimentally difficult to measure.
The model supports the fitting of the parameters $\text{Begin}$, $A$, $B$ and $C$. All data samples are considered in the fitting process irrespective of $\text{Begin}$. The scaling parameter $A$ is restricted to the range $[0,1]$ per default.

**Use without Measured Parent Fraction**

The **Watabe** parent fraction model can be applied even if no parent fraction measurements were loaded. In this case the user has to specify a parameter set which establishes a representative metabolite correction for the used tracer.

**Reference**


### 6.8.8 Power-Damped Exponentials

The sequential steps for converting blood measurements to an input curve are explained in a dedicated section. The current parent fraction model serves for converting a plasma activity curve into an input curve (authentic, unchanged tracer in plasma).

The **Power-Damped Exp** model assumes that the concentration ratio of authentic tracer to total tracer in plasma has been determined at certain times during the acquisition and loaded with the **Load Fraction** entry in the menu.

**Operational Model Curve**

The general **Power-Damped Exp** model for the parent fraction handles the situation where the parent fraction starts at a low value and then increases, before dropping. It has the following functional form

$$f_{\text{parent}}(t) = f_p \cdot \begin{cases} 0 & t \leq 0 \\ A_1 e^{-\ln 2/T_1} + A_2 e^{-\ln 2/T_2} + A_3 e^{-\ln 2/T_3} & t > 0 \end{cases}$$

The calculated fraction is multiplied with $f_p$, the free fraction of authentic tracer in plasma, i.e. the fraction of tracer not bound to plasma proteins.

**Parameter Fitting**
The free fraction $f_p$ is an input parameter which has no impact on fitting the model curve to the fraction data. The default for $f_p$ is 1 as it is experimentally difficult to measure.

The model supports the fitting of the parameters $\alpha$, Amplitude 1, 2, 3 ($=A_1$, $A_2$, $A_3$) and Halftime 1, 2, 3 ($=T_1$, $T_2$, $T_3$). The three Amplitude parameters are restricted to the range [0,1] per default. However, the calculated fractions are not restricted to a certain value range.

**Use without Measured Parent Fraction**

The Power-Damped Exp parent fraction model can be applied even if no parent fraction measurements were loaded. In this case the user has to specify a parameter set which establishes a representative metabolite correction for the used tracer.

**Reference**


### 6.8.9 Sigmoid Function

The sequential steps for converting blood measurements to an input curve are explained in a dedicated section. The current parent fraction model serves for converting a plasma activity curve into an input curve (authentic, unchanged tracer in plasma).

The Sigmoid model assumes that the concentration ratio of authentic tracer to total tracer in plasma has been determined at certain times during the acquisition and loaded with the Load Fraction entry in the menu.

![Sigmoid Function Diagram](image)

**Operational Model Curve**

The general Sigmoid function for the parent fraction has been applied for [11C]PBR28 and has the following functional form [1]

$$f_{\text{parent}}(t) = \begin{cases} f_p & \text{if } t \leq \text{Begin} \\ \left(1 - \frac{u^3}{u^3 + 10^4}\right)^b (1 + C) & \text{if } t > \text{Begin}, u = t - \text{Begin} \end{cases}$$

The calculated fraction is multiplied with $f_p$, the free fraction of authentic tracer in plasma, i.e. the fraction of tracer not bound to plasma proteins.
Parameter Fitting

The free fraction $fp$ is an input parameter which has no impact on fitting the model curve to the fraction data. The default for $fp$ is 1 as it is experimentally difficult to measure. The model supports the fitting of the parameters $A$, $B$ and $C$.

Use without Measured Parent Fraction

The Sigmoid parent fraction model can be applied even if no parent fraction measurements were loaded. In this case the user has to specify a parameter set which establishes a representative metabolite correction for the used tracer.

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7 References


References


References


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