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Positron range in PET imaging: an alternative approach for assessing and correcting the blurring

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Abstract

Background: Positron range impairs resolution in PET imaging, especially for high-energy emitters and for small-animal PET. De-blurring in image reconstruction is possible if the blurring distribution is known. Further, the percentage of annihilation events within a given distance from the point of positron emission is relevant for assessing statistical noise.

Aims: The paper aims to determine positron range distribution relevant for blurring for seven medically relevant PET isotopes, ¹⁸F, ¹¹C, ¹³N, ¹⁵O, ⁶⁸Ga, ⁶²Cu, and ⁸²Rb, and derive empirical formulas for the distributions. The paper focuses on allowed-decay isotopes.

Methods: It is argued that blurring at the detection level should not be described by positron range r, but instead the 2D-projected distance δ (equal to the closest distance between decay and line-of-response). To determine these 2D distributions, results from a dedicated positron track-structure Monte Carlo code, EPOTRAN, were used. Other materials than water were studied with PENELOPE. *Results*: The radial cumulative probability distribution $G_{2D}(\delta)$ and radial probability density distribution $g_{2D}(\delta)$ were determined. $G_{2D}(\delta)$ could be approximated by the empirical function $1 - \exp(-A\delta^2 - B\delta)$, where $A = 0.0266 (E_{mean})^{-1.716}$ and $B = 0.1119 (E_{mean})^{-1.934}$, with E_{mean} the mean

positron energy in MeV and δ in mm. Radial density distribution $g_{2D}(\delta)$ could be approximated by differentiation of $G_{2D}(\delta)$. Distributions in other media were very similar to water.

Conclusion: Positron range is important for improved resolution in PET imaging. Relevant distributions for positron range have been derived for seven isotopes. Distributions for other allowed-decay isotopes may be estimated with the above formulas.

Keywords: Monte Carlo simulation, positron range, spatial resolution, annihilation, PET, point spread function

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1. Introduction

Because of the improving spatial resolution of the physical detectors, the distance from positron emission to positron annihilation (positron range) is becoming a factor of increasing importance for the spatial resolution of positron emission tomography (PET). Positron range introduces uncertainty in the position of the positron emitter, resulting in blurring of the PET images. This effect is very clearly seen for high-energy emitters such as ⁸²Rb, but also for less energetic emitters in small-animal PET imaging where blurring on a millimetre scale is important.

While positron range in itself is a fundamental physical aspect of the slowing-down processes leading to annihilation, it is possible to correct for the blurring in the reconstructed images, if the reconstruction algorithm incorporates knowledge of spatial distribution of the blurring effect (Cal-González *et al* 2009;Rahmim *et al* 2008;Ruangma *et al* 2006). Therefore, the distribution of positron ranges and its effect on PET imaging is a subject of prime importance, especially as technical advances may further improve detector resolution.

A point source emitting positrons will give rise to a three-dimensional (3D) distribution of annihilation points, which will characterize the positron range effect in the reconstructed images. But since a PET scanner measures lines of response (LORs), not annihilation positions, the 3D distribution is hard to measure directly (this also holds for Time-of-Flight PET, which reduces the length of the LOR, but still does not determine a point position). To determine positron range, Derenzo (1979) made a series of often-cited measurements using material of very low density, in an experimental set-up where only a narrow plane was measured at a time. This corresponds to projecting the 3D annihilation distribution onto a single line, giving the 1D representation often used in papers on positron range, see e.g. (Blanco 2006;Champion *et al* 2007;Levin *et al* 1999).

However, considering a single detector pair (rather than a full detector ring), the logical projection will be 2D: The detector pair measures in the plane orthogonal to the LOR defined by the two detectors, and positron range influences not only axial resolution (1D) but also transversal resolution. Thus, for a reconstruction algorithm modelling positron range at the level of LORs, the distribution of the 2D projection of annihilation points will be very relevant.

In any homogeneous medium and in absence of magnetic field, the situation is spherically symmetrical, and the mathematical information in the 3D, 1D, and 2D distributions will be the same, but converting e.g. a 1D distribution to a 2D distribution is not a simple calculation. However, 2D projections seem to have been given only little attention in the literature. Derenzo (1979) computed curves for ¹¹C, ⁶⁸Ga, and ⁸²Rb (from a ⁸²Sr source, which decays to ⁸²Rb by electron capture) showing the "Fraction projected annihilations occurring within a circle of radius R", i.e., the cumulated 2D probability distribution. Sánchez-Crespo *et al* (2004) investigated ¹⁸F, ¹⁵O, and ⁸²Rb by projecting the 3D distribution onto a plane and reported point spread functions of this projection along a line. And recently, Cal-González *et al* (2010;2011) have considered 2D representations of the annihilation distribution and compared them with other representations. But little other work seems to have been conducted on 2D representations of the annihilation distribution.

The present work aims to determine in details the cumulative 2D distributions for a series of medically relevant PET isotopes by means of a previously published dedicated positron track-structure Monte Carlo

simulation (Champion *et al* 2006;Champion *et al* 2007). Furthermore, it is aimed to provide a systematic procedure for estimating the cumulative distribution for other allowed-decay PET isotopes given only the mean positron energy E_{mean} . (A beta decay is allowed if parity is unchanged and nuclear spin change is 0 or ±1 (Krane 1988); e.g. $1^+ \rightarrow 0^+$ as in the decay ${}^{18}\text{F} \rightarrow {}^{18}\text{O}$.) The focus on allowed-decay PET isotopes is practical. The energy distributions can be described by known, analytical functions, and very many of the medical PET isotopes decay by allowed decay. Besides, differential distributions will be provided by following the same methodology. The results from the present Monte Carlo code (EPOTRAN) is compared with results from the more common PENELOPE code, and the influence of other biological media than water is estimated.

2. Material and methods

2.1 Geometry and notation

The complete blurring of a point source in an actual measurement will depend not only on positron range but also on photon non-collinearity and the properties of the detection system (such as crystal size and depth of interaction). However, all these effects can be considered individually and afterwards combined (Levin *et al* 1999;Qi *et al* 1998;Rahmim *et al* 2008). In the present context we will assume point-like detectors and 180° relative angle of the two photons emitted from annihilations, i.e., focus on positron range.

Consider a point source located at (0,0,0) in a Cartesian coordinate system. Assuming spherical symmetry, the volume density f(x,y,z) of annihilatons is function of range: f(x, y, z) = f(r). Following the notation of Cal-González *et al* (2010), we will define the radial density as $g_{3D}(r) = 4\pi r^2 f(r) = 4\pi r^2 f(x,y,z)$.

The LORs measured parallel to a given direction will be cylindrically symmetrically distributed. As the distribution of LORs will be the same for all directions, we can treat the situation as if all LORs were parallel to the same direction. Taking the *z* axis as the (arbitrary) choice of direction, the projection is onto the *xy* plane. The distance from origin in 2D, $\delta = \sqrt{x^2 + y^2}$, corresponds to the distance between the point source and the LOR. Again we have symmetry: The 2D density distribution f(x,y) is a function of these distances, $f(x, y) = f(\delta)$, with radial density $g_{2D}(\delta) = 2\pi\delta f(\delta)$. The cumulative probability distribution $G_{2D}(\delta)$, *i.e.* the probability that a LOR will have distance from the point source no greater than δ , can be calculated by integration:

$$G_{2D}(\delta) = \int_{0}^{\delta} g_{2D}(\delta') d\delta'.$$
(1)

Our notation for g_{2D} and G_{2D} is consistent with that of Cal-González *et al* (2010), except that we have chosen letter δ for 2D radius (LOR distances from point source), reserving *r* for 3D radius (positron range).

2.2 Positron energy spectra and Monte Carlo simulation

The following series of medically relevant PET isotopes were considered: ¹⁸F, ¹¹C, ¹³N, ¹⁵O, ⁶⁸Ga, ⁶²Cu, and ⁸²Rb. These are all allowed-decay isotopes whose energetic disintegration spectra are represented by (Venkataramaiah *et al* 1985)

$$N(E)dE = C \cdot F(Z, E) \cdot p \cdot E \cdot (E_{\max} - E)dE, \qquad (2)$$

with *E* and E_{max} the energy and maximal energy of β particles, *p* the corresponding momentum and *C* a normalization constant. *F*(*Z*,*E*) represents the Fermi factor used for taking into account the effect of the Coulomb field on the electron distribution.

The 3D distributions f(x, y, z) of annihilation points were determined by using the full-differential Monte Carlo track-structure code, called EPOTRAN (an acronym for Electron and POsitron TRANsport in liquid and gaseous water) developed by Champion and Le Loirec to study the decay of positron emitters of medical interest (Champion *et al* 2007;Le Loirec *et al* 2007). In brief, for each isotope considered, a large number (about 250 000) positron tracks were simulated until annihilation, including all the processes, namely, ionization, excitation, elastic scattering, and positronium (Ps) formation. For more details, we refer the interested reader to (Champion *et al* 2012). The distribution $g_{2D}(\delta)$ were determined by binning $\delta = \sqrt{x^2 + y^2}$ into 10 µm bins. Finally, the cumulative probability $G_{2D}(\delta)$ was determined by numerical integration of g_{2D} .

2.3 Generalization of distributions

The resultant functions $G_{2D}(\delta)$ were compared among isotopes in two ways, namely, either by plotting $G_{2D}(\delta)$ as a function of δ , or by plotting $G_{2D}(\delta)$ as a function of δ/R_{mean} , where R_{mean} is the mean range of positrons for the isotope under consideration:

$$R_{mean} = \iiint r \cdot f(x, y, z) dx dy dz = \int_{0}^{\infty} r \cdot g_{3D}(r) dr$$
(3)

An empirical formula for $G_{2D}(\delta)$ was then derived for each allowed-decay isotope, based on two adjustable parameters. The dependence of the parameters on mean positron energy E_{mean} was also investigated. To estimate differential distributions $g_{2D}(\delta)$, the empirical formula for $G_{2D}(\delta)$ was differentiated, and the results compared with distributions of $g_{2D}(\delta)$ determined from the Monte Carlo data.

2.4 Effects of variation in biological tissues

For a medium of a given composition, the range and thereby δ will scale inversely with density. But the shape of the positron range distribution may change with different composition, see e.g. (Lehnert *et al* 2011).

Presently, EPOTRAN is not able to simulate positron track structure in a biological medium other than water. We have thus used the Monte Carlo code PENELOPE (2006 release) (Salvat *et al* 2006) first to compare density-scaled range of positrons in biologically relevant materials with the range in water and secondly to study the decay of ¹⁸F in water, lung and bone.

PENELOPE is a Monte Carlo code of general purpose that permits us to simulate the coupled transport of photons, electrons and positrons in matter with a very good accuracy at low energies. Photons are transported within the conventional detailed method whereas electrons and positrons are simulated using a mixed scheme where interactions are classified into hard and soft. Hard events are simulated step by step and involve angular deflection or energy losses above user-defined cutoffs whereas soft events occurring between 2 hard collisions are simulated by means of an artificial single event. Simulation is controlled by seven parameters, fixed for the various materials used in the geometry:

(*i*) The absorption energies E_{abs} is the energy at which the track evolution is stopped. The kinetic energy of the particle is then locally deposited.

(ii) C_1 is the average angular deflection between 2 hard elastic collisions.

- (*iii*) C_2 is the maximal fractional energy loss between 2 hard elastic collisions.
- (iv) W_{CC} is the cutoff energy for hard inelastic interactions.
- (v) W_{CR} is the cutoff energy for hard bremsstrahlung emission.

To be as close as possible as the simulations performed with EPOTRAN, we have chosen for water, lung and bone the same set of parameters which corresponds to a detailed simulation, thus $E_{abs} = 100$ keV for photons, electrons and positrons and C_1 , C_2 , W_{CC} and W_{CR} are equal to zero.

Tracking particles in PENELOPE is not obvious. It is however possible to determine the point of creation of particles, and in our case of annihilation photons. We have thus used a geometry consisting in concentric cylinders separated by 10 µm, which is the bin used for the distribution $g_{2D}(\delta)$. The cylinders are centred on (0;0;0) which is the point of emission of positrons and they are full of water, bone or lung depending on the tissue under investigation. The simulation runs until the photons are detected on a spherical phase-space file (PSF) which stores a lot of information concerning the particles and especially the body (the cylinder in our case) where they have been created. Thanks to these data, we can thus determine directly the 2D distribution of annihilation points and obtain the cumulative probability $G_{2D}(\delta)$.

3. Results

3.1 Ranges and cumulative probability distributions

The investigated radio-isotopes are listed in Table 1 along with mean and maximum values for energy and range for the emitted positrons; for more details see (Champion *et al* 2007;Le Loirec *et al* 2007). Compared to other results from the literature (see Table 1 and figure 1) we obtain somewhat larger values for R_{mean} . This difference is due to the inclusion of Ps formation in the simulation, as the electrically neutral Ps loses energy more slowly than the charged e⁺ particle (Champion 2003;Champion *et al* 2006).

The literature results for R_{mean} do not include experimental data. Experimental results on R_{mean} are, unfortunately, scarce. Cho *et al* (1975) measured on a line source and determined an "effective range" from

the FW(0.1)M on this distribution, and Derenzo (1979;1986) determined the root-mean-square (rms) spread for the 1D distribution. While such data can be used as indicators for R_{mean} , we are not aware of experimental results that aim at determining R_{mean} as such.

It should be mentioned that while the Monte Carlo code used here assumes the neutral Positronium (Ps) to lose its kinetic energy mainly through phonon interactions at all energies, recent experimental results (Brawley *et al* 2010a;Brawley *et al* 2010b) found unexpectedly large Ps interactions at sub-keV energies. While these results are in themselves very interesting, a discrepancy in Ps cross section at very low energy will only have marginal influence on our results, since most of the path travelled by Ps will be with much higher kinetic energy.

The calculated cumulative probabilities $G_{2D}(\delta)$ for closest distance δ are shown in figure 2. Not surprisingly, the distance δ that includes a given fraction of the LORs, depends on the energy spectrum of the emitted positrons. However, when the same data are plotted as a function of δR_{mean} , *i.e.* scaled by the isotope-dependent mean range, the results are rather similar among isotopes (see figure 3).

 Table 1: Mean and maximum values for energy and range for the emitted positrons (Champion *et al* 2007;Le

 Loirec *et al* 2007).

Isotopes	E _{mean} (keV)	E _{max} (keV)	<i>R_{mean}</i> (mm)	<i>Literature</i> <i>R_{mean} (mm) ^a</i>	<i>R_{max}</i> (mm)
¹⁸ F	252	635	0.660	0.6	2.633
¹¹ C	390	970	1.266	1.1	4.456
¹³ N	488	1190	1.730	1.5	5.572
¹⁵ O	730	1720	2.965	2.5	9.132
⁶⁸ Ga	844	1899	3.559	2.9	10.273
⁶² Cu	1280.6	2926	6.077		16.149
⁸² Rb	1551	3378	7.491	5.9	18.603

^a (Partridge *et al* 2006)



Figure 1: Cumulative probabilities for ¹⁸F obtained with EPOTRAN and PENELOPE.



Figure 2: Cumulative probabilities $G_{2D}(\delta)$ obtained for the radio-isotopes investigated



Figure 3: Cumulative probabilities $G_{2D}(\delta)$ versus δ/R_{mean} .

3.2 Empirical estimation of cumulative probability distributions

Despite their similarity, the plots in figure 3 cannot be described by a single curve. Instead we have for each isotope fitted the cumulative probability curves reported in figure 2 by the following empirical function:

$$\zeta(\delta) = 1 - \exp(-A\delta^2 - B\delta). \tag{4}$$

This form is chosen such that for any positive values of *A* and *B*, it will correctly rise monotonously from $\zeta(0) = 0$ towards 1, $\zeta(\infty) = 1$. Furthermore, for any probability $0 \le \zeta < 1$ considered, the corresponding value of δ can be found as the positive solution to the equation

$$A\delta^{2} + B\delta + \ln(1 - \zeta) = 0.$$
⁽⁵⁾

The values obtained for the parameters A and B are reported in Table 2.

Isotopes	A (mm ⁻²)	<i>B</i> (mm ⁻¹)
¹⁸ F	0.2951 (± 3.6%)	1.5090 (± 0.57 %)
¹¹ C	0.1262 (± 1.7 %)	0.7010 (± 0.45 %)
¹³ N	0.0885 (± 1.0 %)	0.4681 (± 0.38 %)
¹⁵ O	0.0436 (± 0.54 %)	0.2266 (± 0.33 %)
⁶⁸ Ga	0.0414 (± 0.35 %)	0.1499 (± 0.34 %)
⁶² Cu	0.0164 (± 0.28 %)	0.0671 (± 0.40 %)
⁸² Rb	0.0126 (± 0.23 %)	0.0468 (± 0.42 %)

Table 2: Fitting parameters for the empirical cumulative probability given by (4). The standard errors are indicated into brackets for each isotope.

The parameters *A* and *B* obviously depend on isotope. In figure 3 we scaled the curves by the isotopedependent R_{mean} . For the parameters *A* and *B* we investigated instead dependence of the more easily determined value E_{mean} and found that they could be described as power functions of E_{mean} :

$$A = 0.0266 \cdot (E_{mean})^{-1.716} \text{ and } B = 0.1119 \cdot (E_{mean})^{-1.934}, \tag{6}$$

where E_{mean} is given in MeV, and the units of *A* and *B* are mm⁻² and mm⁻¹, respectively. The variations of *A* and *B* with E_{mean} as well as the interpolated power functions are reported in figure 4.



Figure 4: The fitting parameters *A* and *B* (open and solid squares, respectively) versus the mean energy of the isotope under consideration.

As a simple check of these expressions, we study also the case of ⁵²Fe, a radio-metal generatorproduced isotope which emits positrons of relatively low energy, $E_{max} = 803.6$ keV and $E_{mean} = 344$ keV = 0.344 MeV. Clinically, ⁵²Fe has been used in haematological studies to follow iron kinetics, but such studies are complicated by the fact that ⁵²Fe decays to another PET isotope, ^{52m}Mn (Haddad *et al* 2008;Lubberink *et al* 1999). By using (6), we have estimated the fitting parameters as A = 0.1660 mm⁻² and B = 0.8813 mm⁻¹. Figure 5 shows the calculated $G_{2D}(\delta)$ for ⁵²Fe along with the empirical function $\zeta(\delta)$ described by these values of A and B.



Figure 5: Calculated function $G_{2D}(\delta)$ along with empirical function $\zeta(\delta)$ from estimated values of *A* and *B* for ⁵²Fe (solid and dashed line, respectively). A direct fit to the data is shown as a dot-dash-dot line almost coinciding with the full line.

3.3 Radial probability density distributions

For determination of point spread functions (PSFs), the probability density distribution should be known, either as area density, $f(x, y) = f(\delta)$, or as radial density, $g_{2D}(\delta)$. We here focus on the radial probability density, which can be compared to the result of differentiation of the estimate for cumulated probability, $\zeta(\delta)$ from (4):

$$g_{2D}(\delta) = \frac{d}{d\delta} G_{2D} \approx \frac{d\zeta}{d\delta} = (2A\delta + B) \cdot \exp(-A\delta^2 - B\delta).$$
(7)

It will be noted that $d\zeta/d\delta$ is normalized to total area 1, since $\zeta(0) = 0$ and $\zeta(\infty) = 1$. The Monte Carlo data are compared with the result of (7) in figure 6.



Figure 6: Functions $g_{2D}(\delta)$ taken from Monte Carlo calculations (broken lines) compared with the expression from (7) (smooth lines), using values of *A* and *B* from table 2. For ⁵²Fe, the values of *A* and *B* from (6) were used. Monte Carlo data are reported with a 10 µm bin for ¹⁸F, ¹¹C, ¹³N, and ⁵²Fe, and with a 30 µm bin for ¹⁵O, ⁶⁸Ga, ⁶²Cu, and ⁸²Rb.

3.4 Effects of variation in biological tissues

The PENELOPE results for density-scaled positron range in different media are given in Table 3. It is seen that soft tissue has only very small differences from water, while bone tissue deviates by about 13%.

Table 3: Approximate mean range for two isotopes (chosen to represent low and high energy) calculated with PENELOPE in the continuing slowing down approximation (CSDA). For soft and lung tissue, the largest difference from water is 1.2% whereas it is up to 13% for bone tissue.

Biological mate	erials (ICRP compositions)	<i>range</i> \cdot <i>density</i> (g \cdot cm ⁻²)	
Name	<i>Name</i> Density ρ (g·cm ⁻³)		⁶² Cu
Water	1.00	0.0622	0.595
Lung	0.26	0.0634	0.597
Adipose	0.98	0.0613	0.591
Muscle	1.07	0.0635	0.602
Bone	1.92	0.0702	0.666

In figure 7, we have reported the cumulative 2D probabilities obtained in PENELOPE for ¹⁸F in water, bone and lung. The data are plotted as a function of $\delta \rho$ with ρ the density of the biological tissue. The tendency for bone-data to have larger distances can be seen as the bone curve being slightly below the other two curves in the upper half of the figure, but overall the density-corrected distributions are very similar for all three media.



Figure 7: Cumulative probability for water (solid line), bone (dash line) and lung (dotted line) as a function of $\delta \cdot \rho$. All three curves are based on PENELOPE results.

4. Discussion

4.1 Cumulative and differential 2D probability distributions

As noted in the introduction, the choice of 2D distributions is very relevant when considering measurements by individual detector pairs. The cumulative distribution $G_{2D}(\delta)$ shows us that for all the investigated isotopes, almost 2/3 of the annihilations take place within $\delta \leq R_{mean}$ (see figure 3), meaning that R_{mean} can be used as rough estimate of the expected broadening due to positron range.

When based on Monte Carlo data, rather than mathematical functions, the cumulative (integral) distribution is more robust than the differential distribution. But a reconstruction algorithm involving modelling of positron range will typically need the differential distribution, either as area density $f(\delta)$ or radial density $g_{2D}(\delta)$. As demonstrated in figure 6, the radial distributions can be well approximated by use of (7) with the values of *A* and *B* from Table 2. If *A* and *B* are determined from the simple, empirical relations (6) then the distribution will of course be less precise, but the curves for ⁵²Fe in Figures 5 and 6 indicates that such distributions can be used as a first approximation for other allowed-decay isotopes than those in Table 2.

Our focus on radial density has the advantage of avoiding the very peaked (cusp-shaped) distributions found for both 1D (Derenzo 1979) and 2D (Sánchez-Crespo *et al* 2004) non-radial distributions. These peaks are so sharp that the peak height becomes dependent on the details of the calculations (e.g., bin size), meaning that measures dependent on peak height (such as FWHM) do not have a strict meaning (Levin *et al* 1999).

In fact, our own results indicate that the radial distribution $g_{2D}(\delta)$ do not necessarily go to zero for $\delta \rightarrow 0$ (see figure 6), i.e. the area distribution $f(\delta) = g_{2D}(\delta) / (2\pi\delta)$ may diverge for $\delta \rightarrow 0$, in which case FWHM is meaningless. Note that such a divergence is not unphysical as long as the area under the singularity is finite. For example, a simple harmonic oscillator, $x = A \sin(\omega t)$, will have a density distribution for x which diverges at $x = \pm A$, because velocity is zero in these positions. In our case, g_{2D} can be integrated into G_{2D} , with the (physically correct) implication that only a finite number of positrons will be detected near $\delta = 0$.

It can be argued against the form (4) for the cumulative distribution $G_{2D}(\delta)_{\text{estimate}} = \zeta(\delta)$ that it never reaches 1, and therefore the differential distributions $g_{2D}(\delta)_{\text{estimate}} = d\zeta/d\delta$ never fall to 0, despite positrons having a maximum range. The effect of this can be seen on figure 6, where the final part of the curves $g_{2D}(\delta)_{\text{estimate}}$ tend to be slightly above the Monte Carlo data, especially for the high-energy isotopes. Similar objections could however be raised against describing positron range distributions by exponentials (Derenzo 1979) or Gaussian functions (Palmer *et al* 1992). Mathematically, it can be seen that $d\zeta/d\delta$ will eventually approach the zero value faster than a Gaussian function: $d\zeta/d\delta$ can be considered the product of a Gaussian, $\exp(-A\delta^2)$, and a factor that goes towards zero for large distances, $(2A\delta + B) \cdot \exp(-B\delta)$, possibly after an initial rise. If a cut-off distance $\delta_{cut-off}$ is introduced, setting $g_{2D}(\delta)_{\text{estimate}} = 0$ for $\delta > \delta_{cut-off}$, then the area under the curve will then be slightly less than 1, namely $\zeta(\delta_{cut-off})$. A possible choice for $\delta_{cut-off}$ can be R_{max} which can be estimated from E_{mean} as a linear function, based on Table 1 (fit not shown).

In short, our formulation based on A and B is suitable for applications dependent on the whole distribution (such a de-blurring), while it is less suitable or unsuitable for applications sensitive to the behaviour of a small fraction of the positrons (such as the determination of maximal distances). Figure 7 indicates that the same values of A and B can be used for different media if distances are scaled by density, as it has already been proposed by Lehnert *et al* (2011) for a 1D distribution model.

4.2 Limitations

The current study was made only for allowed-decay isotopes. Non-allowed-decay isotopes can have other relations between E_{mean} and spectrum shape, meaning that the empirical relation (6) cannot be expected to give usable results for non-allowed-decay isotopes. As an example we tried it for ¹²⁴I (data not shown), which is non-allowed-decay, and found large discrepancies from expected $G_{2D}(\delta)$.

5. Conclusion

Nowadays, it is well-known that when high-energy positron emitters are used in PET scanning, the positron range adds to resolution degradation. In the major part of existing studies this degradation is related to the positron range (r). In the present work, we have proposed to use not the positron range but instead the shortest distance (δ) between point of emission and the LOR. Annihilation point radial 2D density distributions as well as cumulated probabilities for a series of medically relevant PET isotopes were provided by means of Monte Carlo simulation. Empirical expressions of these distributions were then proposed for the allowed-decay PET isotopes here investigated.

These results allow easy estimation of the percentage of annihilations happening within a given distance and may give approximate density distributions for use in reconstruction algorithms taking positron range into account.

Conflicts of interest: none

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