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Applications of amorphous track structure models for correction of ionization quenching in organic scintillators exposed to ion beams

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Abstract

The scintillation response of organic plastic scintillators irradiated with heavy ions is investigated with the open-source code ExcitonQuenching. The software relies on amorphous track structure theory to account for the radial energy deposition by secondary electrons (EDSE) in ion tracks. The kinematic Blanc model is applied to evaluate the ionization quenching for a given ion by taking the decay time, light yield, and density of the scintillator into account. ExcitonQuenching predicts the scintillation response without a priori knowledge of any measured response curves in contrast to other EDSE models, such as the correction method due to Birks, which rely on free fitting parameters for each ion. ExcitonQuenching is validated against published measurements of the Pilot-U scintillator exposed to several ions. The agreement with experimental data is between 5% and 9% for ions with atomic number \( z \leq 6 \) but deviates significantly for heavier ions.

Keywords: Ionization quenching, organic plastic scintillators, ion beam dosimetry, amorphous track structure theory, particle therapy

1. Introduction

Water equivalent organic plastic scintillator detectors coupled to optical fibers for remote read out (Beddar et al., 1992a,b) enable measurements of absorbed dose with minimal perturbation and a high degree of resolution in time (<1 ms) and space (sub-mm). While both the temperature variations (Buranurak et al., 2013) of the scintillator and the radiation induced stem-effect in the fibre (Archambault et al., 2005) are well-understood and may be corrected for, the scintillators suffer in line with other solid state detectors from a response that is nonproportional to the local energy deposition. The reduced scintillator response occurring at high ionization densities, termed ionization quenching or nonproportionality, remains a challenge. Several kinematic models have been proposed to correct ionization quenching in inorganic scintillators (Michaelian and Menchaca-Rocha, 1994; Vasil’ev, 2008; Williams et al., 2010) but few have been suggested for organic materials.

Birks (1951) suggested a semi-empirical model to correct the scintillator signal for ionization quenching which later was extended by Chou (1952) for particles with a high linear energy transfer (LET) with an additional free parameter. Nonetheless, neither of the models account for the local ionization density in ion tracks. The radial energy deposition by secondary electrons (EDSE) in the tracks of two ions with the same LET, but different atomic numbers, differ due to the ranges of the liberated secondaries. The differing ionization densities in the ion tracks thus cause the scintillation responses to differ as experimentally observed by Buenerd et al. (1976). Consequently, the Birks and Chou models require experimentally determined correction factors for each particle and each scintillator in question.

Michaelian and Menchaca-Rocha (1994) approached the ionization quenching issue with a model based on the EDSE and were as such capable of distinguishing between the quenching from different particles with the same LET. However, their model requires determination of several free quenching parameters by fitting the model to experimentally obtained data. The present work investigates the numerical framework ExcitonQuenching (Christensen and Andersen, 2018) for the case of plastic scintillators exposed to ions heavier than protons.

The software combines amorphous track structure theory and the decay time, light yield, and density of...
the scintillator to estimate the local ionization densities, upon which the scintillator response is evaluated through a kinematic model. ExcitonQuenching is in contrast to other EDSE models able to predict the quenched scintillator response without fitting parameters to specific response curves and can furthermore quantify the temporal structure of ionization quenching. The open-source software ExcitonQuenching is accessible for download\(^1\) to correct the ionization quenching in organic plastic scintillators exposed to ions.

2. Methods

2.1. Ionization quenching

The luminescence \(dL\) per unit length \(dx\) as a function of a particle’s linear energy transfer is often written as

\[
\frac{dL}{dx} = A \cdot \frac{dE}{dx} \cdot QCF, \quad QCF = 1 + kB \cdot \frac{dE}{dx},
\]

where \(A\) denotes the light yield of the scintillator in units of photons emitted per deposited energy, and the quenching correction factor \((QCF)\) is expressed as a linear function with slope \(kB\) in the Birks model. The scintillator response \(L\) for a particle with kinetic energy \(E\) may consequently be computed from eq. (1) as

\[
\frac{dL}{dE} = A \cdot QCF^{-1} = L = \int_{0}^{E} A \cdot QCF^{-1} \cdot dE
\]

for a thick scintillator. The QCF for a given ion and scintillator is a function of the kinetic energy and the EDSE which can be achieved through a track structure.

2.2. Amorphous track structures

Amorphous track structure theory models the EDSE as a continuous radial function and thus predicts the local ionization density based on the kinetic energy of the primary particle and the material composition of the medium. The present work focuses on the Chatterjee-Schaefer track structure model (Chatterjee and Schaefer, 1976), which consists of a dense core region of enormous ionization density and a penumbral region with the characteristic \(r^{-2}\) radial decrease. The Chatterjee-Schaefer track structure model is defined in eq. (A.1) and illustrated in figure 1 for different ions with the same LET. As the ion core regions may exhibit huge excitation densities for slow, heavy ions, the Chatterjee-Schaefer results are compared to the Scholz-Kraft track structure model (Scholz and Kraft, 1996) to evaluate the influence of the applied track structure model. The Scholz-Kraft track structure model is outlined in eq. (A.2).

\footnote{\url{https://github.com/jbrage/ExcitonQuenching}}

2.3. Kinematics of excited states

An ion depositing an energy \(dE\) per unit length \(dx\) in a scintillator gives rise to a linear density of \(N = dE/dx \cdot A\) of excited states (excitons). The exciton distribution immediately after an ion traversed the material slab of unit length \(dx\) is radially distributed according to the Chatterjee-Schaefer track structure.

The excitons of density \(n\) in the ion track are in a kinematic model suggested by Blanc et al. (1962, 1964) allowed to diffuse, fluoresce, and quench according to

\[
\frac{\partial n}{\partial t} = D \nabla^2 n - (p + k)n - an^2,
\]

where the diffusion constant \(D\) and bimolecular quenching parameter \(\alpha\) are estimated from fits in Christensen and Andersen (2018) and listed in table 1. The unimolecular quenching parameter \(k\) and the rate of fluorescence emission \(p\) are related to the decay time \(\tau\) of the scintillator via \(\tau^{-1} = p + k\) (Birks, 1964) and weighted equally. Eq. (3) models the fluorescence emission from the first excited singlet state to the ground state (Blanc et al., 1962).

Table 1: The generic model parameters are calculated in Christensen and Andersen (2018) whereas the decay time \(\tau\), light yield \(A\) (in percent relative to anthracene), and density \(\rho\) for the Pilot-U plastic scintillator are given by the manufacturer (Eljen Technology, USA).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Scintillator parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diffusion (D)</td>
<td>(1.0 \times 10^{-4})</td>
<td>(5.9 \times 10^{-4})</td>
</tr>
<tr>
<td>Decay (\alpha)</td>
<td>(1.0 \times 10^{-9})</td>
<td>(1.0 \times 10^{-9})</td>
</tr>
<tr>
<td>Decay (p)</td>
<td>(1.4)</td>
<td>(1.4)</td>
</tr>
<tr>
<td>Decay (\tau)</td>
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<td>(30)</td>
</tr>
<tr>
<td>Light yield (A)</td>
<td>(67)</td>
<td>(67)</td>
</tr>
<tr>
<td>Density (\rho)</td>
<td>(1.023)</td>
<td>(1.023)</td>
</tr>
</tbody>
</table>

2.4. Quenching correction factors

The kinetic eq. (3) governs the exciton interactions where the initial exciton distribution is given by
the track structure model and scintillator light yield and material composition. The emitted light is calculated by solving eq. (3) numerically (see Christensen and Andersen (2018) for details) while integrating over the fluorescence term $p_\text{f}$. Solving eq. (3) with $\alpha = 0$ gives the light emission in the absence of ionization quenching while solving the same equation with $\alpha \neq 0$ gives the quenched scintillator response. The corresponding QCF can consequently be obtained as the ratio of the unquenched signal to the quenched signal.

The following section applies ExcitonQuenching to investigate the ionization quenching in a plastic scintillator (Pilot-U, Eljen Technology, USA) which was subject to irradiation with several ions heavier than protons by Buenerd et al. (1976). The results are obtained as predictions without fitting free parameters to experimentally determined scintillator responses for the Pilot-U scintillator in contrast to the Birks model and most EDSE models.

3. Results

3.1. Ionization quenching

The QCF is calculated with ExcitonQuenching for $^4\text{He}$, $^6\text{Li}$, $^9\text{Be}$, $^{12}\text{C}$, and $^{16}\text{O}$ ions for a wide range of kinetic energies in the Pilot-U scintillator, where the LET as a function of energy for a given ion and material composition is calculated as suggested by Bird and Williams (1990). The QCFs as a function of LET is shown in figure 2 where the Birks model in eq. (1) has been fitted to each set of calculated QCFs.

Figure 2: The quenching correction factor as a function of LET for five ions is calculated with ExcitonQuenching plotted with symbols. The Birks model is fitted to each set of data shown with solid lines.

The scintillation efficiency $dL/dE$ for the five ions are calculated according to eq. (2) as a function of the QCF which in turn depends on the kinetic energy of the particle. The results are shown in figure 3 as a function of kinetic energy.

3.1.1. Light emission for isotopes

The scintillation efficiency for three isotopes of hydrogen, helium, and lithium are in figure 4 calculated as a function of velocity $\beta = v/c$ and velocity per atomic number $\beta/z$, respectively. The scintillation efficiency converges in all cases towards a quenching-free level for $\beta \to 1$, corresponding to QCF $\to 1$ in eq. (2).

Figure 3: Theoretical light emission per energy as a function of kinetic energy in eq. (2) for the Pilot-U scintillator irradiated with different ions.

3.2. Light output

The luminescence response from the plastic scintillator induced by a stopped ion of kinetic energy $E$ is calculated by integrating $dL/dE$ over the kinetic energy as given by eq. (2), where the QCF as a function of kinetic energy is interpolated for a continuous integration. The theoretical results are compared to the experimental results obtained by Buenerd et al. (1976) in figure 5.
where the results are collectively scaled to the experimental data with a single value. The light output is calculated for all ions using the Chatterjee-Schaefer and Scholz-Kraft track structures where the isotopes $^7\text{Li}$ and $^9\text{Be}$ are shown for reference.

![Graph](image-url)

Figure 5: The light emission as a function of kinetic energy for ions stopping in the plastic scintillator. The calculated ExcitonQuenching results are shown with solid lines (Chatterjee-Schaefer track structure) and dashed lines (Scholz-Kraft track structure), whereas the experimental data points from Buenerd et al. (1976) with symbols as given by the legend. The luminescence is shown for three isotopes of lithium for comparison. The ratios between the calculated and experimentally measured scintillation responses are shown below where the dashed horizontal lines show ±10% discrepancies.

4. Discussion

4.1. Track structure theory

The ion core densities for the five different ions in figure 1 with the same $\text{dE/dx} = 30\text{keV}\mu\text{m}^{-1}$ span almost two orders of magnitude and illustrate the importance of the track structure models; the oxygen ion ($E \approx 190\text{MeV/A}$) liberates secondaries with significantly larger kinetic energies than the $\alpha$ particle ($E \approx 5.5\text{MeV/A}$). The core and penumbra radii in the oxygen ion track are consequently broader which in turn reduce the ionization density and quenching. The variation of the QCFs as a function of $\text{dE/dx}$ in figure 2 illustrates how the ionization quenching varies with the atomic number and $\text{dE/dx}$ as experimentally confirmed by Newman and Steigert (1960).

4.2. Scintillation efficiency

The scintillation efficiency in figure 3 as a function of energy shows how lighter ions more efficiently convert energy into luminescence for a given energy. The scintillation efficiency for each ion slowly converges as the kinetic energy increases in turn reducing the ionization density of core region and quenching, which also was observed experimentally (Murray and Meyer, 1961).

The luminescence responses for the isotopes of a given element coincide as a function of velocity in figure 4, which also was experimentally confirmed by Avdeichikov et al. (2002) for the case of BGO exposed to light ions. The scintillation efficiency as a function of velocity per atomic number is approximately the same for all ions as explained and measured by Sibczynski et al. (2018) for GAGG:Ce excited by alpha particles.

4.3. Light emission

The scintillation response curves in figure 5, as ions are stopped in the plastic scintillator, illustrate the variation of luminescence efficiency with atomic number. The emitted light as a function of kinetic energy differs between isotopes of the same element, as illustrated in the figure with the 3 isotopes of lithium, which is in line with the experimentally measured responses reported by Avdeichikov et al. (2002) and figure 4.

The agreement between the ExcitonQuenching calculated and experimentally measured scintillation response curves for helium, lithium, and carbon ions confirms the application of track structure theory and the kinematic Blanc model to compute the ionization quenching for $z \leq 6$ ion beams. However, the deviation between the predicted and measured response for oxygen ions (as much as ±25%) in figure 5 exhibits a structure where the response is underestimated for low energies and overestimated for high energies. The magnitude and structure of the discrepancy between the two track structure models and the data are similar, implying that the deviation between theory and experimental data is related to the model itself and not the particular amorphous track structure model in question.

The kinematic Blanc model in eq. (3) governs the fluorescence emission from de-excitation of the first singlet state but does not account for triplet state interactions. Two triplet states may interact and give rise to one molecule in the ground state and one molecule in the singlet state and as such leading to fluorescence emission at a slower time scale than normally observed (Michaelian and Menchaca-Rocha, 1994; Jain et al., 2009). Thus, the probability for a bimolecular interaction between two triplet states and subsequent fluorescence emission increases with the local ionization density and occurs more frequently in carbon and oxygen ion tracks than in lighter ion tracks as helium and lithium. The possibility of extending the kinematic
Blanc model to include triplet state interactions was suggested by Blanc et al. (1964) but is out of the scope of the present work.

Another explanation of the deviation may be found in the current equation itself, where higher-order density terms may be required to balance the high-LET ions. Such a conclusion was in line with Torrisi (2000) where the Birks model with an additional free parameter, i.e., the Chou model, was concluded to provide a better fit to the data for high-LET.

Conversely, the good agreement between data and ExcitonQuenching for \( z \leq 6 \) ions matches the results in Christensen and Andersen (2018), where the software accurately predicted the scintillation response for another plastic scintillator (BCF-60, Saint-Gobain, France) irradiated with protons.

5. Conclusion

The open-source software ExcitonQuenching predicts the quenched scintillation response in plastic scintillators exposed to ion beams based on amorphous track structure theory and a kinematic exciton model. The agreement between the software and experimental data for helium, lithium and carbon ions is, except for one outlier, better than 9%, and as such in line with traditionally reported experimental uncertainties. The large discrepancy between the model and data for oxygen ions indicates that an additional model parameter, along with the inclusion of triplet state interactions, are required for high-LET ions.

Nonetheless, ExcitonQuenching accurately predicts the ionization quenching in the organic plastic scintillator exposed to ions with \( z \leq 6 \) based on the decay time, light yield, and density of the scintillator without any fit to the data. Such a predictive capability is in sharp contrast to most available EDSE models which contain free parameters requiring fits to each set of experimental data.

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Appendix A. Amorphous track structure models

Chatterjee-Schaefier. The track structure model due to Chatterjee and Schaefer (1976) consists of a dense core with radius defined as

\[
\begin{align*}
    r_{\text{min}} &= \beta_{\text{ion}} \times 11.6 \text{ nm}, \\
    r_{\text{max}} &= 0.768E - 1.925 \sqrt{E} + 1.257 \text{ [\mu m]} \\
    C_1 &= \log \left( \frac{r_{\text{max}}}{r_{\text{min}}} \right) - 1 \\
    N &= \frac{dE}{dx} \cdot A.
\end{align*}
\]

Scholz-Kraft. With core and penumbral radii defined as

\[
    r_{\text{max}} = 0.05 \ E^{1.7} \text{ [\mu m]}, \quad r_{\text{min}} = 0.01 \text{ [\mu m]},
\]

where \( E \) [MeV/nucleon] is the kinetic energy of the projectile, the Scholz-Kraft track structure model governs a radial exciton density of

\[
    n(r) = \begin{cases} 
    \frac{C_2}{r_{\text{min}}} \frac{2 + C_1}{r_{\text{min}}^2} & \text{for } r < r_{\text{min}} \\
    \frac{C_2}{r_{\text{max}}} & \text{for } r_{\text{min}} \leq r \leq r_{\text{max}} \\
    0 & \text{for } r > r_{\text{max}}
    \end{cases}
\]

for

\[
    C_2 = N \left( 1 + 2 \ln \frac{r_{\text{max}}}{r_{\text{min}}} \right)^{-1}.
\]
References


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