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Citation: Applied Physics Letters 102, 161915 (2013); doi: 10.1063/1.4803440
View online: http://dx.doi.org/10.1063/1.4803440
View Table of Contents: http://scitation.aip.org/content/aip/journal/apl/102/16?ver=pdfcov
Published by the AIP Publishing
Improvement of γ-ray energy resolution of LaBr₃:Ce³⁺ scintillation detectors by Sr²⁺ and Ca²⁺ co-doping

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(Received 14 March 2013; accepted 14 April 2013; published online 26 April 2013)

Commercially available LaBr₃:5% Ce³⁺ scintillators show with photomultiplier tube readout about 2.7% energy resolution for the detection of 662 keV γ-rays. Here we will show that by co-doping LaBr₃:Ce³⁺ with Sr²⁺ or Ca²⁺ the resolution is improved to 2.0%. Such an improvement is attributed to a strong reduction of the scintillation light losses that are due to radiationless recombination of free electrons and holes during the earliest stages (1–10 ps) inside the high free charge carrier density parts of the ionization track. © 2013 AIP Publishing LLC. [http://dx.doi.org/10.1063/1.4803440]

Scintillating materials have been applied for more than a century, and they played a crucial role in the discovery of X-rays, β-particles, and α-particles and today in the quest for the Higgs boson. Not long after the development of the photomultiplier tube (PMT) around 1940, the very important NaI:Tl⁰⁰⁰⁺ scintillation crystal was discovered. Till today NaI:Tl accounts for 80% of all scintillator volume sold world-wide. Lu₂SiO₅:Ce³⁺ (LSO:Ce) combines a strong interaction with γ-rays because of its high density and the presence of the high atomic number Lu atoms, with a very fast scintillation decay time of 35 ns from the lanthanide Ce³⁺. It makes this material ideally suited for medical imaging in positron emission tomography (PET)-scanners. Modern day PET-scanners may contain tens of thousands of individual 2 × 2 × 20 mm³ LSO:Ce crystal pixels to generate a three dimensional image of a scanned patient. The electromagnetic calorimeter at CERN, Geneva, which is used for the quest for the Higgs boson, contains 110 000 23 cm long PbWO₄ scintillating crystals developed in the 1990s. Starting from the year 2000, we developed the Ce³⁺ activated La-halide scintillators, first LaCl₃:Ce (Ref. 6) and later LaBr₃:Ce. Particularly LaBr₃:Ce combines excellent properties, i.e., an extremely fast scintillation pulse (16 ns) and a record low energy resolution for the detection of γ-rays. A full width at half maximum (FWHM) resolution of 2.8% at 662 keV was unequalled at that time.

In this letter we present a study on Sr²⁺ and Ca²⁺ co-doped LaBr₃:5% Ce³⁺ and CeBr₃ scintillators. We will show that such co-doping improves the linearity of scintillation response with X-ray or γ-ray energy. As a result, the energy resolution improves, and record of low energy resolutions of 2.0% at 662 keV and 6.5% at 59.5 γ-ray detection will be demonstrated. One set of Ca²⁺ and Sr²⁺ co-doped samples used in this study were grown by the University of Bern. The starting materials LaBr₃, CeBr₃, SrBr₂, and CaBr₂ were obtained from oxides with the NH₄Br method. To avoid moistening, the starting materials and products were always handled and stored in a N₂ glove box. The starting materials were weighted in the corresponding stoichiometric ratios (total of 8 g) into silica ampoules and sealed under high vacuum. 5 mol. % of CeBr₃ and 0.5 mol. % of either SrBr₂ or CaBr₂ were added. The mixtures were molten in a horizontal tube furnace and homogenized. The ampoules were placed in vertical Bridgman furnaces, heated to approximately 10 °C above their melting points and then cooled to RT over a period of 14 days. Another independent set of Sr²⁺-co-doped samples were provided by the Saint-Gobain Company. They were synthesized with a propriety method also used for the commercially available BriLaNce380 standard LaBr₃:5%Ce scintillators. The Ce³⁺ concentration added to the starting material was 5 mol. %, and 0.35 mol. %–0.75 mol. % of SrBr₂ was added. An ICP (Inductively Coupled Plasma) analysis on the samples revealed that typically 200 ppm of Sr²⁺ is actually incorporated in the final single crystal.

Small samples of about 3 × 3 × 1 mm³ size were cut from the original larger single crystal boule and mounted on top of the entrance window of a standard bialkali Hamamatsu R1791 PMT or a super bialkali Hamamatsu R6231-100 PMT. The sample was covered with several layers of white reflecting Teflon tape to ensure optimal photon collection on the PMT window. Scintillation pulse height spectra of ¹³⁷Cs and ²⁴¹Am radioactive sources were recorded with conventional techniques using an electronic shaping time of 10 µs. Since the samples are hygroscopic all experiments were performed in an M-Braun UNILAB dry box with moisture content less than 1 ppm. The scintillation yield, or number of detected photons by the PMT, was obtained from the ratio of the scintillation charge output pulse and the charge output pulse generated by the detection of one single photon as outlined in Refs. 8 and 9. The method and equipment used for avalanche photodiode (APD) readout was the same as reported in Ref. 8. An electronic shaping time of 2 µs was used. Scintillator response studies were also performed at the X1 X-ray station of Hasylab in Hamburg, Germany. In these studies the sample was sealed in a silica ampoule and mounted on a PMT placed directly in front of the output slit of a tunable double Bragg reflection monochromator providing an X-ray resolution of 1 eV at 9 keV rising to 20 eV at 100 keV. A 50 × 50 µm slit...
area was used assuring that always the same small volume of the sample was irradiated.

One may distinguish three contributions to the energy resolution $R$, and assuming that they are independent from one another

$$R^2 = \frac{2.35^2}{N_{ndp}} + R_{intr}^2 + R_{det}^2$$

(1)

applies. The first term is from the Poisson statistics in the number of detected photons $N_{ndp}$, the second term is an intrinsic contribution from the scintillator. The third term contains the contributions from variations in crystal quality throughout the bulk, variations in photon collection efficiency on the entrance window of the photon detector, and a noise contribution from the photon detector. By means of proper crystal growth and packaging technology this contribution needs to be minimized. Figure 1 shows the energy resolution $R$ at 662 keV as observed for current day scintillators that are readout with an ordinary bialkali PMT against the fundamental limit $R_{ndp} = 2.35/\sqrt{N_{ndp}}$. The best resolution of 2.65% is observed for LaBr$_3$:Ce. Only SrI$_2$:Eu approaches similar energy resolution$^{10,11}$ when small samples are used.

Scintillator research always aimed to increase $N_{ndp}$ in order to minimize $R_{ndp}$. Such strategy directed research towards smaller bandgap materials; first chlorides, then bromides, and nowadays the iodides are intensively investigated$^{12}$. The importance of the intrinsic contribution has always been neglected in scintillation research and development, probably because its true physical origin was not clear and also not measurable. Yet, the large deviation between $R$ and $R_{ndp}$ for the widely applied LSO:Ce, NaI:Tl, and CsI:Tl scintillators in Fig. 1 must be attributed to such intrinsic contribution. It is caused by a scintillator response that is not proportional to the amount of ionization energy deposited in the scintillator.$^{13,14}$ Figure 2 compares the response curves for NaI:Tl,$^{15}$ Lu$_2$SiO$_5$:Ce,$^{16}$ and LaBr$_3$:5% Ce (Ref. 17) as has been determined with monochromatic synchrotron X-ray excitation. Here the scintillation response is defined as the ratio of the photon yield/MeV observed at the X-ray energy set by the monochromator to the photon yield/MeV observed at 662 keV, and ideally it is 100% at all energies. The response of NaI:Tl at 20 keV is 17% higher than at the reference energy of 662 keV, whereas LSO:Ce is almost 35% less efficient. LaBr$_3$:5%Ce approaches best the ideal response, and indeed it shows the lowest energy resolution in Fig. 1. Recently we found that the shape of the response curve of LaBr$_3$:Ce depends on the temperature$^{17}$ and the Ce$^{3+}$ concentration,$^{18}$ which demonstrates that the shape is not a truly fundamental property of a scintillator and that it can be altered.

A main cause for non-ideal response at low X-ray energy is radiationless electron-hole pair recombination that takes place in the high ionization density parts of the ionization track. That means close (<5 nm) to the track, and since the energy loss $dE/dx$ of the ionizing primary electron increases with decreasing electron velocity (or energy $E$), the recombination losses are highest at the end of the main track.$^{19}$ It is also high in the side tracks of secondary electrons created by “head-on collisions” of the primary electron with electrons of the scintillating medium. The stochastic nature of side track formation is the origin of $R_{intr}$. Only when the scintillation yield is independent on the electron energy, $R_{intr}$ will vanish and the fundamental limiting energy resolution $R_{ndp}$ comes within reach.

With the idea in mind that the shape of the response curve can be altered we decided to study the effect of Sr$^{2+}$ and Ca$^{2+}$ co-dopants on the performance of standard LaBr$_3$:5% Ce scintillators. In Fig. 2 the response curve for a Sr$^{2+}$ co-doped LaBr$_3$:5% Ce scintillator has been added. A large improvement as compared to standard commercial LaBr$_3$:5%Ce scintillators is observed, and the response is much closer to the ideal one.$^{20}$ The samples grown at the University of Bern and those at Saint-Gobain showed similar response curves. We also studied the response curve for CeBr$_3$, which appears poorer than that of standard.
LaBr₃:5%Ce as shown in Fig. 3. This is consistent with an observed poorer energy resolution of about 3.8% at 662 keV. Like in LaBr₃:5%Ce, co-doping with Sr²⁺ improves the response curve of CeBr₃ significantly. It is not yet as ideal as in LaBr₃:Ce,Sr, but we expect that by changing the co-doping concentration or by using different type of co-dopants the response can be further engineered towards the ideal one.

The improved response in the 10–100 keV region has consequences for the light yield and energy resolution at 662 keV. With a standard bialkali PMT, about 17 200 photons are detected from a standard LaBr₃:5%Ce scintillation crystal at 662 keV γ-ray energy. By using a Hamamatsu R6231-100 super bialkali PMT or a cooled APD that are higher quantum efficiency photon detectors, the number of photons increases to 24 000 and to 42 700, respectively. Resolution improves from 2.7% to 2.5% and to 2.45%. With Eq. (1) one then obtains $R_{\text{intr}} \approx 1.9\%$, and evidently $R_{\text{intr}}$ provides the dominant contribution to the observed resolution. The star data symbol in Fig. 1 shows that the energy resolution with a standard bialkali PMT for a Sr co-doped sample improves to 2.35%. With the super bialkali PMT the improvement is more impressive as shown with the 137Cs source pulse height spectrum in Fig. 4. The total absorption peak at 662 keV shows a record low resolution of 2.0%. The peak due to 33 keV La-Kα X-ray fluorescence escape is fully separated from the main peak, and also the Ba-Kα and Ba-Kβ X-ray fluorescence from the 137Cs source can be very well distinguished (see the inset of Fig. 4). We now calculate that for the Sr co-doped scintillator $R_{\text{intr}}$ has lowered to $\approx 1.3\%$ which is clearly connected with the more ideal response curve in Fig. 2. We predict that with well cooled APD readout, when $R_{\text{det}}$ becomes insignificant, a resolution of 1.8% should be feasible. First tests at 8°C gave 2.0% energy resolution, and improvements are to be expected with further cooling.

Fig. 5 compares the pulse height spectrum of a 241Am source measured with a standard and a Sr-co-doped LaBr₃ scintillation crystal. Resolution at 59.5 keV improves from 9.4% to 6.5%. Also, the peaks between 10 and 30 keV are much better resolved in the Sr co-doped sample.

Figures 2 and 3 show a strong drop in the response curves when passing the K-shell or sometimes L-shell electron binding energies of the atoms in the scintillator material. The drop at the threshold is caused by a re-distribution of the available X-ray energy over a set of secondary electrons with relatively low energy that each creates ionization tracks. A better representation of the response of a scintillator is obtained by measuring the response as a function of electron energy instead of as function of X-ray or γ-ray energy. We recently developed a method to obtain such a response down to very low electron energy of 100 eV. By tuning the X-ray energy just above the K-shell electron binding energy $E_K$ of the most heavy element in the scintillator, a K-shell photoelectron will be ejected. The energy of this electron can be controlled within 10 eV accuracy simply by tuning the X-ray energy. One thus creates an internal electron source that is tunable from 100 eV to, say, 60 keV. The excess scintillation light produced by that photoelectron can be derived from the...
data. With this so-called K-dip spectroscopy method, response curves as a function of K-shell photoelectron energy were obtained as shown in Fig. 6. It reveals that $\text{Ca}^{2+}$ and $\text{Sr}^{2+}$ co-doping alter the response curve of standard LaBr$_3$:5%Ce in a similar fashion. The point of 10% recombination loss is at 4.1 keV in the standard LaBr$_3$:5%Ce scintillator, and it shifts towards 1.3 keV in the $\text{Sr}^{2+}$ and $\text{Ca}^{2+}$ co-doped scintillators. Since energy loss $-dE/dx$ of a track creating electron increases with $\approx 1/E$, this implies that the ionization density at 1.3 keV is three times higher than at 4.1 keV. Apparently, the scintillator has become a factor of three more tolerant towards high ionization density recombination losses by the co-doping with either $\text{Sr}^{2+}$ or $\text{Ca}^{2+}$.

The charge carrier dynamics taking place on the nm length scale around the track and in the ps time scale are extremely complicated and at this stage not fully understood. The current idea is that the nonradiative recombination rate $\Gamma_{nr}$ of free electrons and holes competes with the trapping rate $\Gamma_{tr}$ of those charge carriers by Ce$^{3+}$. Once trapped by Ce the desired radiative recombination will follow. $\Gamma_{nr}$ increases with ionization density, and it is therefore highest at the end points of the main and side tracks. When carrier mobility is high the free charge carriers may escape from the high ionization density parts of the track to survive the quenching phase, and one may define an escape rate $\Gamma_{esc}$. Since temperature affects the carrier mobility and Ce$^{3+}$ concentration affects the carrier trapping rate, $\Gamma_{tr}$ the co-doping either reduces $\Gamma_{nr}$ or increases $\Gamma_{esc}$ or $\Gamma_{tr}$ in order to explain the improved proportionality.

In conclusion, we demonstrated that the response curve of LaBr$_3$:5%Ce scintillators can be improved by $\text{Sr}^{2+}$ co-doping. The intrinsic resolution $R_{inst}$ at 662 keV is then reduced from 1.9% to 1.3% which results in 2.0% energy resolution when scintillation readout is done by a super bialkali PMT. We anticipate that a resolution of 1.8% should be possible with well cooled APD read-out. We found that Ca$^{2+}$ co-doping has similar effect as $\text{Sr}^{2+}$ co-doping, and both co-dopants also improve the response curve of CeBr$_3$ scintillators. The implications of the improved energy resolution of LaBr$_3$:5%Ce are immediate. Particularly for applications where energy resolution is crucial, e.g., in nuclear isotope identifiers, the benefits of co-doping are evident. With a better resolution, different isotopes are more reliably detected which is of particular importance in systems for homeland security inspections and for nuclear non-proliferation.

This work was funded by the Dutch Technology Foundation (STW) and supported by Saint-Gobain Crystals, France and by the European Community’s Seventh Framework Program (FP7/2007-2013) under Grant Agreement No. 226716. We sincerely thank Dr. A. Owens and Dr. F. Quarati for sharing their beamtime at the synchrotron with us and their assistance in some of the experiments.

Fig. 6. The K-shell photo-electron response curves for standard LaBr$_3$:5%Ce and $\text{Sr}^{2+}$ and $\text{Ca}^{2+}$ co-doped samples.