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Article

Investigation of GaGG:Ce with TOFPET2 ASIC Readout for Applications in Gamma Imaging Systems

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Abstract: We investigated two detector modules, each consisting of a 8×8 matrix of GaGG:Ce crystals with a crystal size of $3 \times 3 \times 20 \text{ mm}^3$ and a 3.2 mm pitch. The light is collected by a 8×8 silicon photomultiplier array, with one silicon photomultiplier matching one crystal. The signals are read out and processed using the TOFPET2 ASIC. Performed laboratory tests of the detectors were performed using a ^{22}Na source, where energy and coincidence time resolution with different optical coupling were examined between the crystals and silicon photomultipliers, as well as under various operating voltages. The mean energy resolution of $9.8 \pm 0.6\%$ at 511 keV was observed and it was shown that the coincidence time resolution of $384 \pm 33 \text{ ps}$ could be achieved. The results reassure that the GaGG scintillator is a very promising candidate for the development of imaging systems, in particular, ones utilizing Compton scattering where the energy resolution plays a critical role and a moderate timing performance is acceptable.

Keywords: GaGG; silicon photomultiplier; positron emission tomography; Compton imaging; gamma polarization; TOFPET2

1. Introduction

Detectors applied in many modalities of medical imaging, such as Positron Emission Tomography (PET), are based on scintillation materials, most commonly inorganic crystals. The desirable properties of the crystals are a high light yield resulting in good energy resolution, high stopping power, and a fast decay time needed to achieve a good coincidence time resolution in Time-of-Flight PET (ToF-PET) systems [1]. One of the most commonly used materials in PET systems is Cerium doped Lutetium Yttrium Oxyorthosilicate, $\text{Lu}_{1.8}\text{Y}_{0.2}\text{SiO}_5:\text{Ce}$ (LYSO:Ce) [2], satisfying those prerequisites. Scintillation crystals that supersede LYSO in the light output, such as CeBr_3 [3] or LaBr_3 [4], offer better energy resolutions compared to LYSO [5], but their extremely high hygroscopicity and frailness make them difficult to handle since they often need expensive encapsulations.

Gadolinium Aluminum Gallium Garnet doped with Cerium (GaGG:Ce), $\text{Gd}_3\text{Al}_2\text{Ga}_3\text{O}_{12}:\text{Ce}$, is a relatively new material that first appeared less than 10 years ago, offering significantly higher light output than LYSO [6,7]. It has been investigated in various form-factors and readout configurations and the results also demonstrated an improved energy resolution [8–12]. It is also advantageous by not having intrinsic radioactivity, which makes it a good candidate for single arm (i.e., non-coincidence) applications. On the other hand, its scintillation has a longer decay time, resulting in a lower coincidence time resolution (CTR) compared to LYSO [13–15]. The properties of GaGG crystals of various form factors and from various producers, coupled to different light collection devices, are being actively investigated, since this scintillator might prove advantageous for imaging devices

where a high energy resolution is needed, such as medical imaging devices in PET. GaGG may be especially interesting for improving the detection capabilities of Compton imaging systems, which use Compton scattering kinematics to determine the direction of incoming gamma radiation [16–18].

Compton scattering may also be used to measure polarization correlations of 511 keV annihilation quanta and exploit them to improve signal to noise ratio and consequently the image contrast in PET [19–21]. A great way to measure the Compton scattering is to apply segmented scintillator detectors coupled to Silicon photomultipliers (SiPMs). Commonly, systems with two readout layers are used to detect the recoil electron and scattered photon, respectively. It has been demonstrated that single layer readout detectors may reconstruct the Compton scattering [22] and conceptual studies have been performed to evaluate their performance as Compton cameras [23]. Moreover it a prototype system has shown that using two modules of segmented Lutetium Fine Silicate (LFS) crystals read out by SiPMs, make it possible to observe azimuthal modulation related to polarization correlations of annihilation quanta [24] and the results suggested that a further improvement in the precision identification of polarization correlations would be desirable. One way to achieve this is to improve the energy resolution of the system, hence the GaGG:Ce scintillator might prove an interesting solution.

The goal of this investigation is to examine the energy resolution and coincidence time resolution of GaGG:Ce in a specific matrix configuration of 8×8 crystals of $3 \times 3 \times 20 \text{ mm}^3$, coupled to the matching SiPM arrays and read out by fast digital electronics optimized for PET systems.

2. Materials and Methods

2.1. Experimental Setup

The experimental setup consisted of two detector modules with GaGG:Ce scintillating pixels and Silicon Photomultiplier (SiPM) arrays which were read out by fast data acquisition and processing system, the details of which are given below.

2.2. GaGG:Ce Detector Modules

We investigated the properties of two detector modules, each consisting of a 8×8 matrix of GaGG:Ce scintillator crystals and a SiPM array. We obtained the GaGG:Ce matrices from Hilger Crystals Ltd (Margate, UK). Each contained 64 crystals with a size of $3 \text{ mm} \times 3 \text{ mm} \times 20 \text{ mm}$. All crystal sides were polished and assembled with a 0.2 mm reflector material, resulting in a matrix pitch of 3.2 mm. The reflector was an epoxy resin loaded with a specific TiO_2 -based pigment (details proprietary to Hilger Crystals), with the reflectivity of $>98\%$ at 520 nm. The matrix is shown in the Figure 1. The main properties of the material are listed in Table 1 along with the properties of the LYSO scintillator for comparison.

The scintillator matrices were coupled to SiPM arrays with matching pixel dimensions of $3 \text{ mm} \times 3 \text{ mm}$ and the same pitch of 3.2 mm (Hamamatsu Photonics, Hamamatsu, Japan, model S13361-0808AE). The SiPMs have a breakdown voltage at 51.4 V and they were operated at an overvoltage $V_{OV} = 2\text{--}5 \text{ V}$ to explore the impact on energy and timing resolution. Another thing we wanted to explore was the influence of optical coupling to the detector performance. Previously we reported that using optical grease may not be optimal with crystal matrices, since it tends to infiltrate between the crystals, changing the optical properties [25]. Therefore other optical coupling materials were used: The Module I was optically coupled using 1 mm of thick silicon pad (Eljen Technology, Sweetwater, TX, USA, EJ-560) and the Module II was optically coupled using optical cement (Eljen Technology, Sweetwater, TX, USA, EJ-500). The former optical material had a refraction index, $n = 1.43$ while the latter had $n = 1.57$ according to the manufacturer. A lightproof housing for the modules was 3D printed using 2 mm of thick black ABS material.

Table 1. The properties of GaGG(Ce) [26] and LYSO [2] according to the manufacturers. * We note that the upper limit of the emission spectral range of GaGG:Ce quoted in the literature is mostly between 650–750 nm [8,11,13].

Property	GaGG:Ce	LYSO:Ce
Decay constant [ns]	88	36
Density [g cm^{-3}]	6.7	7.1
Z_{eff}	54.4	66
Emission spectral range [nm]	475–800 *	380–480
Peak of the emission spectrum [nm]	520	420
Light yield [photons/MeV]	~55,000	33,200
Higroscopicity	No	No
Refractive index at emission max	1.90	1.81

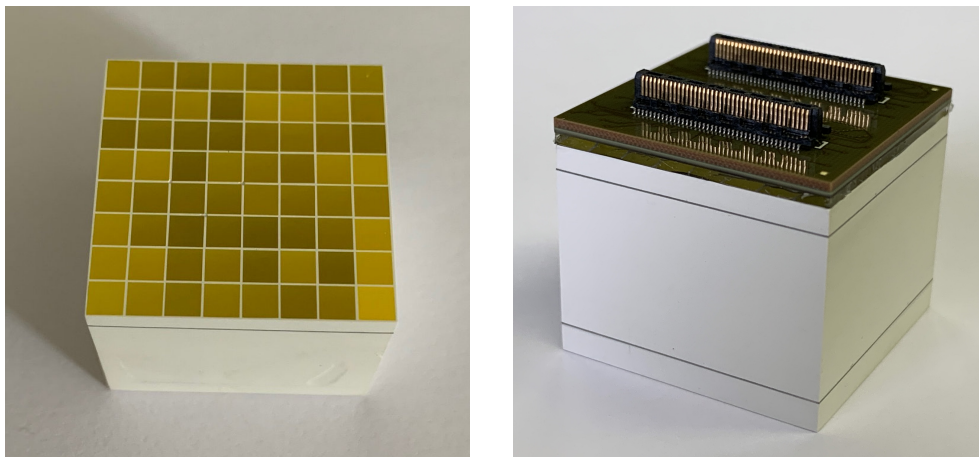


Figure 1. GaGG:Ce matrix containing 64 crystals with a size of $3 \text{ mm} \times 3 \text{ mm} \times 20 \text{ mm}$ in 8×8 configuration. All crystal sides were polished and casted in 0.2 mm of white epoxy (**left**). GaGG:Ce matrix along with the SiPM array optically coupled with a silicon optical interface (**right**).

2.3. SiPM Control and Data Acquisition System

The data acquisition (DAQ) and processing system TOFPET2 produced by PETsys Electronics—Medical PET Detectors, S.A. (Oeiras, Portugal), was used [27,28]. The system consists of a central DAQ module, which can accompany up to 8 front-end boards (FEB), where two detector modules fit on one board. Each FEB contains two powerful ASICs enabling full control, readout, and data processing from the SiPMs. In the basic setup this system can read up to 1024 channels, 64 per module. The system comes with an evaluation kit, containing two $3 \times 3 \times 3 \text{ mm}^3$ LYSO crystals coupled to SiPMs with an appropriate adapter board compatible with the FEB socket. A Peltier-cooled black box is also a part of the evaluation kit, and it was used for the test measurements reported in this paper. The on-board firmware enables bias voltage selection individually for each module, as well as temperature monitoring. The TOFPET2 can be programmed to trigger on single events or coincidences and in the latter case it acquires events in which two signals occur in detector modules belonging to different trigger regions. For the measurements reported here, we used the default settings of the TOFPET2 system, enabling a recording of all signals and selection of coincidences by offline filtering.

2.4. Energy Calibration and Non-Linearity Correction

The SiPMs have a limited number of cells that can fire within one event. In this case, each SiPM channel in the array had 3600 microcells. This leads to a known effect of signal saturation at higher energies and it is especially pronounced with bright scintillators such as GaGG:Ce. The effect of signal saturation is clearly visible in Figure 2 (left), where the ratio of the peak positions of 1275 keV gamma and 511 keV gamma is smaller than $1275/511 \simeq 2.5$.

The analog-to-digital converter on the ASIC also exhibits a small non-linearity according to manufacturer PETSys. In order to perform the non-linearity correction and energy calibration one may use several gamma sources with different energies. However the DAQ system producer recommended applying a specific correction function which linearizes the charge Q , measured by the charge-to-digital converter (QDC):

$$E [\text{a.u.}] = c_0 c_1^{Q^2} + c_3 Q - c_0 \quad (1)$$

with $c_0 = 8.00000$, $c_1 = 1.04676$, $c_2 = 1.02734$, and $c_3 = 0.31909$ [29]. This approach was tested to correct the non-linearity and calibrated the resulting spectrum using only the 511 keV peak. In Figure 2 (right) are the energy spectra from Module I and II after the non-linearity correction and calibration. The ^{22}Ne (1275 keV) gamma peaks can have a deviation of up to $\pm 3\%$ from linearity, however the deviation from linearity within the 511 keV peak, under study in this work, is negligible. The width of the 1275 keV peak measured by Module II in Figure 2 (right) is $6.3\% \pm 0.3\%$ (FWHM), compared to $9.9\% \pm 0.1\%$ at 511 keV, showing that the scaling of the energy resolution with $\sim 1/\sqrt{E}$ is preserved.

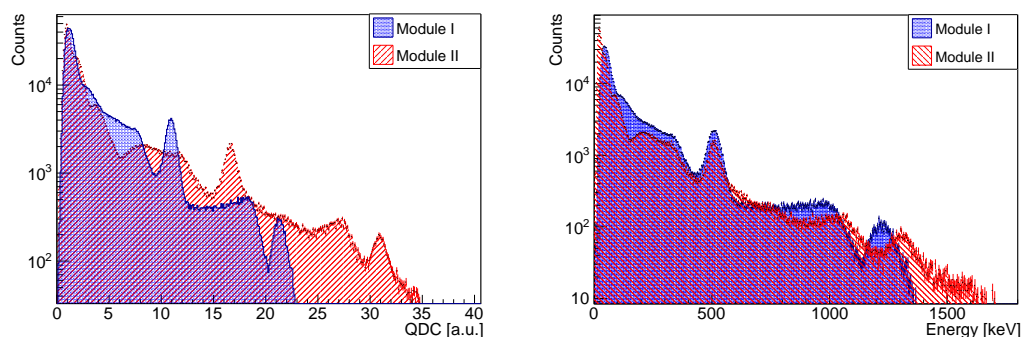


Figure 2. The ^{22}Na spectra before non-linearity correction and calibration measured by a crystal from Module I and Module II at $V_{OV} = 3$ V, respectively (**left**). The energy spectra of ^{22}Na after applying the non-linearity correction and calibration of the 511 keV peak (**right**).

2.5. Reference Measurements

To test the performance of the SiPM control and DAQ system, two LYSO pixel detectors of $3 \times 3 \times 3$ mm³ (produced by Epic-Crystal, Kunshan, China) were used, coupled to SiPM (KETEK GmbH, München, Germany, model PM3325-WB-D0), that had been provided as a part of the evaluation kit for reference measurements.

The energy spectrum of coincident events originating from ^{22}Na source was formed by plotting only the energies of the events where both opposing crystals fired. An example is shown in Figure 3 (left). At the recommended overvoltage value $V_{OV} = 4$ V, the energy resolution $\Delta E/E = 12.4 \pm 0.4\%$ at 511 keV was obtained.

The coincidence time spectrum was constructed by the difference of the two detector trigger times, taking into account only the signals within the energy range $511 \text{ keV} \pm 1\sigma$, where σ is the standard deviation of the Gaussian fit to the photo-peak (Figure 3). At $V_{OV} = 4$ V, we obtained the coincidence time resolution of 235 ± 7 ps. Since the system consists of two same detectors, one may consider that the contribution of each detector to the time resolution is $\text{CTR}/\sqrt{2}$, hence:

$$\delta t_{\text{LYSO}} \simeq 165 \pm 5 \text{ ps.} \quad (2)$$

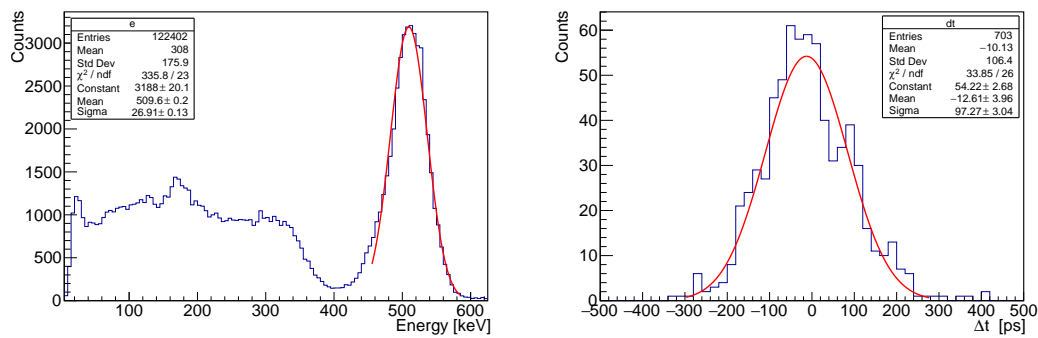


Figure 3. The 511 keV gamma spectrum obtained by $3 \times 3 \times 3$ mm³ LYSO crystal with KETEK SiPM at 4 V overvoltage (left). Coincidence time spectrum of annihilation gammas from ²²Na, detected by two $3 \times 3 \times 3$ mm³ LYSO crystals with KETEK SiPM at 4V overvoltage (right). Events with energy within 511 keV $\pm 1 \sigma$ were selected.

3. Results

The gamma source throughout the measurements was a point-like ²²Na. All measurements were carried out in controlled temperature conditions in the temperature range of 18.5 ± 0.5 °C. The breakdown voltages of both SiPM arrays were $V_{BD} = 51.4$ V and measurements were performed at voltages $V_{OV} = 2$ –5 V above the breakdown value. The TOFPET2 ASIC was used with the default (factory) settings and events where two detectors fired in coincidence were filtered out.

3.1. Energy Resolution

Examples of the measured spectra of 511 keV gamma are shown in Figure 4, where the one on the left corresponds to a crystal in Module I and the one on the right to a crystal in Module II. To determine the energy resolution $\Delta E / E$, the Gaussian function was fitted with the linear background:

$$f(x) = p_0 e^{-\left(\frac{x-p_1}{2p_2}\right)^2} + xp_3 + p_4. \quad (3)$$

The mean energy resolutions are obtained from the measured resolutions of the fired crystals in a module. They are summarized in Table 2. The uncertainties are the standard deviations of the measured values in each configuration and reflect the (in)homogeneity within a module. The best energy resolution of a single crystal was 9.2% for a pixel in Module II at $V_{OV} = 3$ V, shown in Figure 4 (right).

The results clearly indicate that better resolution is achieved with the Module II at all bias voltages. Before gluing it with the optical cement, we tested the GaGG matrix and the SiPM of the Module II using silicon optical pad. The mean energy resolution of the fired pixels was $12.4\% \pm 1.0\%$ at $V_{OV} = 3$ V and it was $11.3\% \pm 1.2\%$ at $V_{OV} = 4$ V. Within the uncertainties, these results are consistent with the ones of the Module I quoted in Table 2 for the respective bias voltages, suggesting that the difference in the performance between the modules is driven by the optical coupling.

Table 2. The measured energy resolution $\Delta E / E$ at 511 keV. The quoted value are mean resolutions of fired pixels in the module. The uncertainties are the standard deviations of the measured values in each configuration.

V_{OV} [V]	Module I	Module II
2	$14.3\% \pm 1.4\%$	$10.6\% \pm 0.3\%$
3	$12.4\% \pm 0.8\%$	$9.8\% \pm 0.6\%$
4	$12.2\% \pm 0.8\%$	$10.1\% \pm 0.4\%$
5	$13.1\% \pm 0.7\%$	$10.5\% \pm 0.7\%$

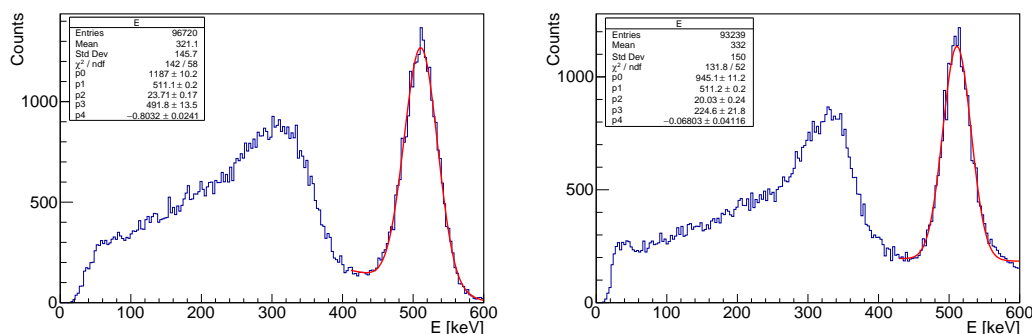


Figure 4. The spectra of 511 keV gammas from ^{22}Na measured by a crystal from Module I at $V_{OV} = 4$ V (left) and from Module II at $V_{OV} = 3$ V (right). The fit function is the one given in Equation (3). The spectra with the best obtained energy resolutions are shown: $\Delta E/E = 10.9\%$ for the Module I and $\Delta E/E = 9.2\%$ for the Module II.

3.2. Coincidence Time Resolution

Coincidence time resolution was measured for the coincident detection of the 511 keV annihilation quanta. First, we selected the events where both oppositely facing pixels had fired. Then, the coincidence time spectrum was constructed by taking the trigger time difference, $\Delta t = t_2 - t_1$, of the two signals which are both within the energy range of $511 \text{ keV} \pm 1 \sigma$ in their respective energy spectra.

We carried out three sets of measurements. In the first set, the system consisted of a single LYSO pixel detector coupled to SiPM PM3325-WB-D0 (KETEK GmbH, München, Germany) and the GaGG Module I, which used 1 mm of thick silicon pad (Eljen Technology, Sweetwater, TX, USA, EJ-560) for optical coupling between the crystal matrices and the SiPM array. In the second set of measurements, it was replaced by Module II, where optical cement (Eljen Technology, Sweetwater, TX, USA, EJ-500) was used for optical coupling. In the third set of measurement, Module I and Module II were tested in coincidence. In measurements with the LYSO crystal, the detector was always kept at the same bias voltage of $V_{OV} = 4$ V. Examples of obtained coincidence time spectra are shown in Figure 5–7, respectively for the three sets of measurements. The results are summarized in Table 3.

Table 3. The measured coincidence time resolution: Each value is the mean coincidence time resolution (CTR) of multiple pixel pairs in that configuration and the uncertainty is the standard deviation of the measured values.

V_{OV} [V]	Set 1	Set 2	Set 3
	Module I vs. LYSO	Module II vs. LYSO	Module I vs. Module II
2	560 ± 33 ps	387 ± 21 ps	625 ± 26 ps
3	479 ± 22 ps	359 ± 18 ps	554 ± 39 ps
4	453 ± 21 ps	318 ± 28 ps	511 ± 27 ps
5	448 ± 24 ps	345 ± 15 ps	533 ± 29 ps

From the measured CTR in Sets 1 and 2, and the extracted contribution of the LYSO detector (Equation (2)), we can derive the contribution of the GaGG detector as:

$$\delta t_{\text{GaGG}} = \sqrt{\text{CTR}_{\text{GaGG,LYSO}}^2 - \delta t_{\text{LYSO}}^2} \quad (4)$$

For Module I at $V_{OV} = 4$ V, this yields $\delta t_{\text{GaGG,I}} = 422 \pm 23$ ps and for Module II at $V_{OV} = 4$ V, it is $\delta t_{\text{GaGG,II}} = 272 \pm 33$ ps. Hence one can deduce the system's CTR of consisting of a pair of GaGG crystals from modules I and II:

$$\text{CTR}_{\text{GaGGI,II}} = \sqrt{\delta t_{\text{GaGG,I}}^2 + \delta t_{\text{GaGG,II}}^2} \quad (5)$$

According to the above equation, a system consisting of Module I and Module II would have the $CTR = 502 \pm 26$ ps, which is consistent with the measured value 511 ± 27 ps at $V_{OV} = 4$ V (Table 3, Set 3). Hence, we can use the measured data to deduce the CTR of a system consisting of two GaGG modules coupled by optical cement according to:

$$CTR_{\text{GaGG}} = \sqrt{2 \times \delta t_{\text{GaGG,II}}^2} \quad (6)$$

The resulting coincidence time resolution at $V_{OV} = 4$ V is $CTR_{\text{GaGG}} = 384 \pm 33$ ps.

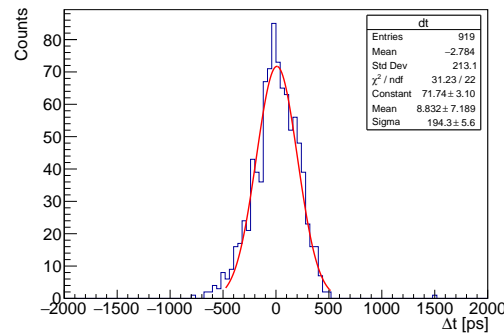


Figure 5. Coincidence time spectrum of annihilation gammas from ^{22}Na , detected by a $3 \times 3 \times 3$ mm³ LYSO crystal and a crystal from GaGG Module I operating at $V_{OV} = 4$ V.

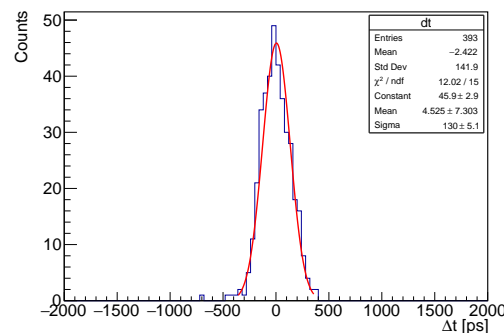


Figure 6. Coincidence time spectrum of annihilation gammas from ^{22}Na , detected by a $3 \times 3 \times 3$ mm³ LYSO crystal and a crystal from GaGG Module II operating at $V_{OV} = 4$ V.

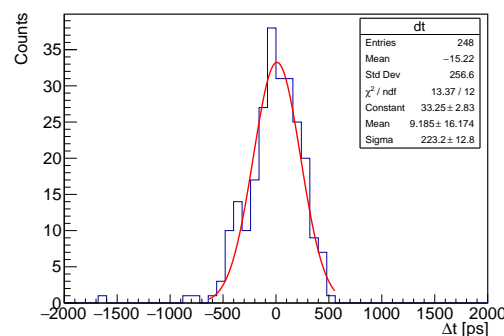


Figure 7. Coincidence time spectrum of annihilation gammas from ^{22}Na , detected by crystals from GaGG Modules I and II, both at $V_{OV} = 4$ V.

4. Discussion

The energy resolution results presented here are consistent with the measurement carried out using similar setups [10,12]. The observed dependence of the energy resolution on over-voltage, shows that dark counts and cross talk effects in the SiPM start playing a role, which negatively influences the resolution at high overvoltages. The observed energy and timing resolutions both suggest that coupling by optical cement results in a superior performance than the silicon interface. In fact the deduced coincidence time resolution of GaGG, $CTR_{\text{GaGG}} = 384 \pm 33$ ps, is according to our knowledge, one of the best yet observed for this crystal. The difference in the performance with the two optical coupling materials could have several reasons. First, we note that the optical cement has an index of refraction closer to the one of GaGG:Ce, which may result in a higher light transfer efficiency, as observed in [15]. Second, the optical cement coupling is much thinner (~ 100 μm) compared to the silicon's 1 mm, where photons incoming at large angles may be directed outside of the matching SiPM's active area, hence reducing the signal. Finally, the manufacturer quotes a ~ 94 transmittivity (for 2 mm thick silicon), while they quote $>99\%$ for the cement [30]. It remains a question for further investigations whether these factors can fully absorb the observed difference in the performance.

We can also compare the obtained results with a previously performed investigation of single $3 \times 3 \times 3$ mm³ GaGG:Ce crystals with a DRS4-based setup [31]. The setup consisted of two detectors, each containing of a single $3 \times 3 \times 3$ mm³ GaGG:Ce crystal (by Hilger Crystals) coupled to a SiPM produced by Hamamatsu Photonics (S13360-3050CS). Each crystal was wrapped in 0.2 mm of thick Teflon and coupled to the SiPM using optical grease. Signals were sampled at 1 GS/s by DRS4 evaluation board and stored for offline analysis. Temperature during measurement was 20 ± 1 °C. At an operating voltage $V_{BD} + 3$ V, we obtained the energy resolution of $11.9\% \pm 0.5\%$ and a coincidence time resolution, $CTR = 550 \pm 15$ ps [32]. Although comparing two different readout systems and different crystal geometries may be hard to interpret, we may conclude that the TOFPET2-based setup with optical cement coupling yielded comparable or better results, in spite of having longer crystals (and matrices), which tend to worsen the time resolution [14].

5. Conclusions

This research showed that good energy resolution and an acceptable coincidence time resolution could be achieved using GaGG:Ce scintillators. The coincidence time resolution that could be achieved with the setup was $CTR_{\text{GaGG}} = 384 \pm 33$ ps, which is comparable to resolutions of the most commercial PET systems.

The best mean energy resolution of the crystals in a module was $\Delta E/E = 9.8\% \pm 0.6\%$ at 511 keV, while the best observed resolution of a single crystal in the matrix was 9.2%. One has to bear in mind that the GaGG emission spectrum peaks at 520 nm, which is ~ 70 nm above the SiPM quantum efficiency maximum. Hence an optimal SiPM, in terms of crystal emission/SiPM efficiency spectra matching, might bring even higher energy resolutions.

Given the obtained results, we believe GaGG:Ce crystal is a great candidate for further developments of imaging systems, in particular ones based on Compton imaging, where the performance critically depends on the energy resolution.

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