

An active electron polarized scintillating GSO target for neutrino physics

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Abstract

The feasibility of an electron-polarized, active target to be used as detector in neutrino scattering experiments, suggested by several theoretical papers, has been investigated. We report on the properties of the paramagnetic crystal Gd_2SiO_5 (GSO), in which 7.7 % of the total number of electrons present can be polarized by lowering the temperature and applying an intense external magnetic field.

The material magnetic susceptibility has been measured down to cryogenic temperatures showing that for $H = 5$ T and $T = 4$ K about 80% of the maximum allowed magnetization can be attained. Also the spectral and time response of the crystal have been characterized and the scintillation process has been studied using a photomultiplier to measure the response to gamma rays irradiation and cosmic rays operating the GSO crystal at 13.5 K. An avalanche photodiode (APD) readout of the scintillation signal from the GSO crystal has also been performed, since the magnetic field-independent response of this device allows it to be placed close to the crystal in the cryogenic environment.

Key words: neutrino detectors, APD, scintillators, magnetization measurements, paramagnetic material, SQUID.

1. Introduction

The standard model of the electro-weak interaction [1] is based on a Lorentz structure with exchange of vector-axial (V-A) current [2, 3], where left-handed neutrinos can interact mainly with left-handed electrons (or nuclei). When a detecting material is prepared in a spin polarized state, this feature provides a mean to control the contribution of the weak interaction to the total cross section of the process. If the target is polarized by an externally applied magnetic field, it is possible to change the rate of the interaction by inverting the field direction.

A new generation of neutrino experiments could stem using detectors based on this mechanism, as suggested in several theoretical papers [4, 5, 6]. It has been pointed out that investigation of the flavor composition of a (anti)neutrino beam could be possible by studying their scattering from polarized electrons in a polarized target [7]. A polarized electron target was suggested also to improve the sensitivity of the search for

the neutrino magnetic moment in the scattering process (anti)neutrino - electron [8]. Compared to traditional detection techniques, in this approach - that makes use of polarizable material - the electromagnetic background limit should not be relevant, since it could be subtracted from the spectra as a measurable noise contribution. This would be important especially in the detection of low energy neutrinos. Besides, switching the direction of the field will permit discrimination against cosmic-ray induced detector response, removing much of the noise limiting the currently used detectors. A neutrino telescope has been proposed in which the active detecting material is based on *nuclei* originally polarized with spin antiparallel to the spin of the incoming neutrinos [9]; absorption of neutrinos from the In^{115} nucleus is suggested, and the fully polarized detector is described as being twice as sensitive as the unpolarized detector.

The research of polarized matter may be of interest also for other research subjects such as axions [10], CP violation through the study of the electric dipole moment [11], and spin-spin interaction in gravitation [12].

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In this paper we present measurements that confirm the possibility to realize an active polarized target crystal of Gd_2SiO_5 (GSO) doped with Cerium (GSO:Ce), as proposed in [13]. Within the herewith described approach, the detected output signal carrying information about interactions is scintillation. It is worth noticing, however, that it has also been proposed to use paramagnetic materials to build a magnetic adiabatic calorimeter [14], which would open the way to the use of several different types of materials, also in powder form, and likely characterized by higher degree of attainable electron polarization. Investigations on this calorimetric approach are under way in our group and will be the subject of a forthcoming paper.

GSO:Ce is a well known scintillator and its properties have been extensively studied (see for instance [15] and references therein). It has a density of 6.71 g/cm^3 and, at room temperature, its light yield is about 9000 photons/MeV. Its recombination time, measured at room temperature [15] is of approximately 60 ns for excitation energies in excess of 4.51 eV (Gd^{3+} levels), while it is about 26 ns in the UV region (Ce absorption).

The magnetic properties of $\text{Gd}_2\text{SiO}_5\text{:Ce}$ are due to the rare earth element Gd in the oxidation state Gd^{3+} ; the electronic configuration of trivalent Gd is $4f^7$, with seven localized unpaired electrons, responsible for the Curie paramagnetic behavior. The magnetization of a Curie paramagnet at a given temperature T and applied field H is described by the Brillouin function [16], according to which a high degree of magnetization is obtained at low temperature and high magnetic field. It is worth noticing that even when all the unpaired $4f^7$ electrons have a spin parallel to the applied magnetic field, only 7.7 % of total number of electrons present in the target are polarized.

Taking into account the behavior of Gd^{3+} paramagnetic salts reported in reference [17], the magnetic polarization of Gd within GSO can be as high as 6.3 Bohr magnetons/Gd atom for a unitary ratio between the applied magnetic field H and the temperature T . Therefore, a magnetic field of about 4 T should be sufficient to fully polarize the material at a temperature of 4.2 K.

It is the aim of the present work to check the possibility of obtaining the desired high degree of polarization in GSO crystals for ordinary laboratory values of H and T and to characterize the scintillating response under these conditions.

This paper is divided in two parts. The first is devoted to characterize the magnetic behavior of GSO, through the study of the temperature dependence on the magnetic susceptibility $\chi = M/H$ and of the field dependence of the magnetization. Measurements of the opti-

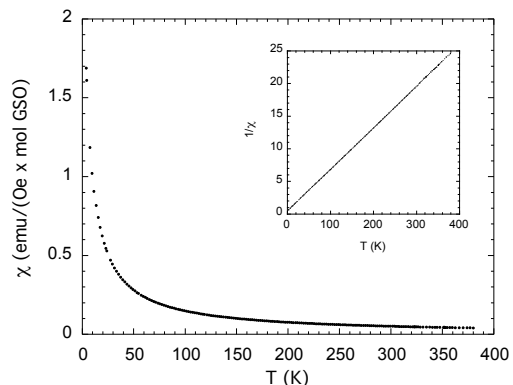


Figure 1: Magnetic susceptibility of a cube shaped GSO single crystal measured with $H = 100 \text{ Oe}$ applied along the three cube edges. The insert shows the inverse susceptibility as a function of T , showing the linear Curie-Weiss behavior.

cal response to short laser pulses in the UV range, conducted with the material kept at cryogenic temperatures and with an external magnetic field, are also reported.

In the second part of the paper the scintillation properties of magnetically polarized GSO under gamma ray irradiation are discussed. For this characterization, two different methods were used, one using a photomultiplier tube (PMT) for the read-out and one using a silicon avalanche photodiode (APD) operated at cryogenic temperatures.

2. Material Characterization: Susceptibility, Spectral Response and Time Response

2.1. Magnetic characterization

The DC (i.e., static field) magnetization measurements were performed using a SQUID-based Magnetic Property Measurement System (Model MPMS-XL5 by Quantum Design Inc.). The SQUID magnetometer has a sensitivity of 10^{-8} emu ; the system is equipped with a superconducting magnet producing fields up to 50 kOe and calibrated using a Pd standard. The residual field during the zero field cooling was less than $5 \cdot 10^{-2} \text{ Oe}$.

A small, cube shaped GSO crystal was cut from a prismatic single crystal and the magnetic susceptibility was measured in the field cooling mode for $H = 100 \text{ Oe}$ applied along the three cube edges. Since no differences were detected among the three measurements, Fig. 1 reports one of the three obtained $\chi(T)$ curves.

As shown in Fig. 1, the susceptibility follows a Curie-Weiss law behavior $\chi = C/(T - \theta)$, with $C = p_{eff}^2 N_a B^2 / 3k_B$, in which N_a is the Avogadro number, μ_B

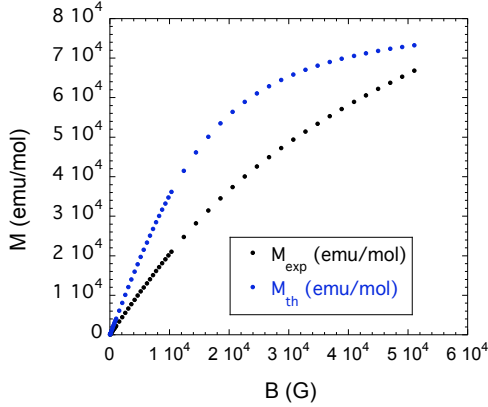


Figure 2: $M(B)$ curve at $T = 4$ K for GSO. The theoretical curve (upper one) is calculated by the Brillouin function for the $4f^7$ electronic configuration of free Gd^{3+} ions. This curve asymptotically saturates to the value of $7 \mu_B/Gd$ atom.

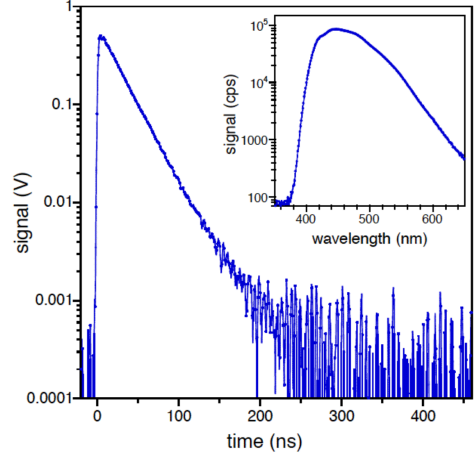


Figure 3: Scintillation signal and emission spectra (upper insert) at room temperature and $B = 0.9$ T.

125 is the Bohr magneton, k_B is the Boltzmann constant and
 126 p_{eff} is the effective number of Bohr magnetons for each
 127 magnetic ion. A linear fit of the $\chi^{-1}(T)$ plot is displayed
 128 as insert to Fig. 1. The fit gives $\theta = -8.9(2)$ K and
 129 $C = 16.13(7)$ emu K/(Oe mol), from which we obtain
 130 for p_{eff} , the effective number of Bohr magnetons for Gd
 131 atom, the value of $8.02(2)\mu_B/Gd$ atom, in good agree-
 132 ment with what reported in literature for Gd^{3+} species.
 133 The negative sign of the Weiss temperature indicates anti-
 134 ferromagnetic correlations between Gd^{3+} ions in GSO.
 135 However, no sign of transition towards a magnetically
 136 ordered phase was detected down to the lowest mea-
 137 sured temperature of 2 K.

138 The field dependence of the magnetization, measured
 139 after zero field cooling to 4 K is shown in Fig. 2, which
 140 displays M as a function of the internal field B acting
 141 on the Gd magnetic moments, obtained from the applied
 142 field H (corrected for the normalized magnetization
 143 factor N taken from [19]) and the GSO magnetization
 144 M (magnetic dipole moment per unit volume) as
 145 $B = H + 4\pi(1 - N)M$ in CGS units.

146 The upper curve in Fig. 2 is the theoretical one, calcu-
 147 lated by the Brillouin function, which describes the
 148 behavior of a collection of $4f^7$ independent ions, i.e.
 149 in absence of exchange interactions. It is seen that
 150 the experimental data deviate appreciably from the ex-
 151 pected behavior. In search of the source of the devia-
 152 tion, one may consider in particular the antiferromag-
 153 netic $Gd^{3+}O^{2-}Gd^{3+}$ super-exchange interaction oppos-
 154 ing the parallel alignment of the gadolinium moments
 155 at 4 K, as shown by the Curie-Weiss behavior of the
 156 susceptibility with negative θ , which likely constitutes

157 the major source of deviation from the independent ion
 158 approximation. On the contrary, since $L = 0$ for Gd^{3+}
 159 (S state), incomplete orbital quenching or the effect of
 160 crystalline field splitting cannot be invoked. Regardless,
 161 the results show that almost 80 % of the saturation mag-
 162 netization can be induced in GSO by $H = 5$ T at $T = 4$
 163 K.

2.2. Optical spectral response and timing

164 The optical properties of GSO:Ce have been mea-
 165 sured both under magnetic field and at cryogenic tem-
 166 peratures, using a GSO crystal of $(2 \times 2 \times 250)$ mm³ size.
 167 The crystal was inserted inside a small permanent mag-
 168 net giving a magnetic flux of 0.9 T and next placed in a
 169 cryostat into a liquid He bath. The cryostat had an opti-
 170 cal window allowing the crystal to be irradiated with a
 171 laser beam (YAG 355 nm, 200 Hz, 25 nJ/pulse).

172 Fig. 3 shows the timing and the emission spectra of
 173 the photoluminescence signal at room temperature and
 174 $B = 0.9$ T. The signal decay characteristic time is mea-
 175 sured to be $\tau = (25 \pm 0.8)$ ns. This value is in good
 176 agreement with what reported in [15].

177 Fig. 4 and 5 respectively show the lifetime and the
 178 photoluminescence emission spectra at different tem-
 179 peratures, ranging from 2 K to 280 K. No important
 180 variations are observed for the signal lifetime with re-
 181 spect to temperature. Minor systematic variations in
 182 proximity of the emission peak can be noticed but the
 183 emission bandwidth is not changed.

184 In summary, the measurements show that the photo-
 185 luminescence properties of the GSO crystal are not sig-
 186 nificantly altered by temperature from 300 K down to
 187

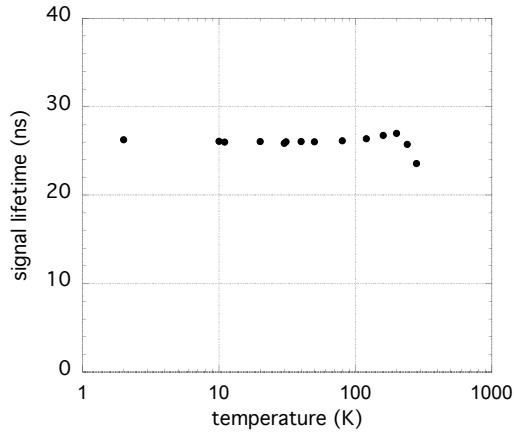


Figure 4: Signal lifetime τ at different temperatures

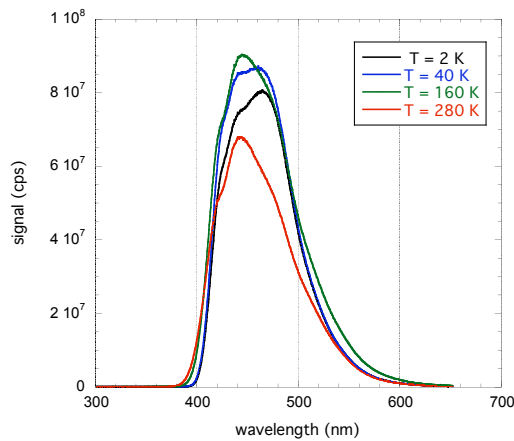


Figure 5: Emission spectra at different temperatures

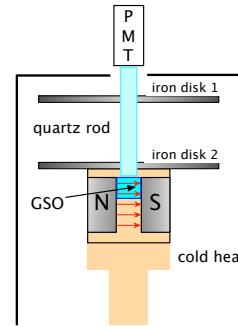


Figure 6: Scheme of the experimental setup used to study the GSO scintillation signal.

188 2 K nor are they altered by the presence of a magnetic
189 field.

190 3. Scintillation properties of the polarized material 191 under gamma irradiation

192 In the measurements presented in this section the
193 GSO crystal is kept at low temperature and is under the
194 influence of the magnetic field. As already stated in the
195 introduction, this is one of the necessary tests to check
196 the feasibility of a scintillating neutrino detector based
197 on a scintillating polarized material.

198 The GSO:Ce scintillation under irradiation of 511 keV
199 gamma source was studied both using a PMT and an
200 APD. Note that, since the aim of these measurements
201 is to check if the scintillating properties change at low
202 temperature and under the influence of the magnetic
203 field, particular care in reducing noise contributions has
204 not been taken. The main noise contribution in the ob-
205 served spectra is the microphonic noise originating from
206 the cryocooler.

207 The GSO:Ce crystal ($1 \times 1 \times 1$) cm³, 0.5 % Ce doped,
208 was provided by the Hitachi Chemical Company. Its sur-
209 face is optically polished and it was laterally wrapped with
210 Teflon to prevent escaping of light.

211 3.1. Scintillation response of the polarized GSO read at 212 a PMT

213 To study the polarized GSO scintillation signal under
214 gamma rays irradiation, the experimental setup shown
215 in Fig. 6 has been arranged.

216 The crystal was enclosed into a double-C-shaped
217 copper mounting fixed on a cryocooler head. Inside
218 the copper mounting it is possible to accommodate also
219 two permanent magnets, generating a measured mag-
220 netic field of 0.98 T in a 1 cm gap (measurement done at
4

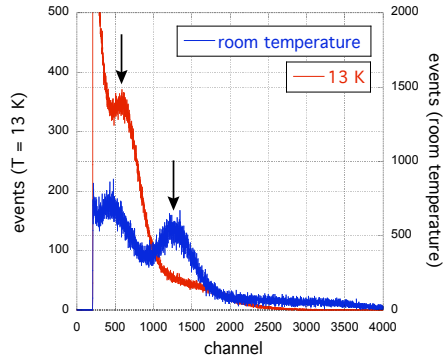


Figure 7: The 511 keV gamma energy deposition events at room temperature and at 13.5 K. The arrows indicate the full-energy peaks corresponding to absorption of the 511 keV photons.

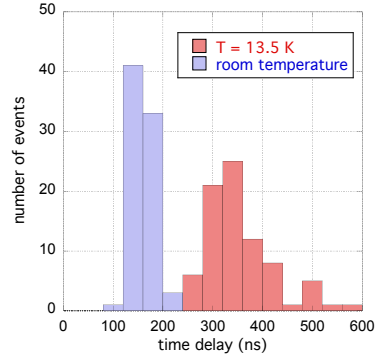


Figure 8: Distributions of measured time decays at room temperature and $T = 13.5$ K.

221 77 K). The photomultiplier tube (Hamamatsu R2027))
 222 is operated at room temperature, protected against mag-
 223 netic field disturbances by two soft iron plates; a 10 cm
 224 long quartz waveguide couples the light signal from the
 225 irradiated crystal to the PMT.

226 The stability of the PMT during measurements has been
 227 monitored sending 10 ns duration light pulses through
 228 an optical fiber.

229 The scintillation signals have been processed through a
 230 shaping amplifier (Silena mod. 7612), before entering
 231 an MCA (Ortec mod. 926).

232 The spectra from a Na^{22} radioactive source, placed un-
 233 der the GSO crystal, are shown in Fig. 7.

234 The results demonstrate that the light yield is not in-
 235 fluenced by a magnetic field of 1 T. The same result has
 236 been obtained with the GSO crystal kept at 13.5 K. As
 237 can be seen in Fig. 7, we observed a significant decrease
 238 (of the order of 40%) of the light signal at low temper-
 239 ature, that can be possibly attributed to the dependence
 240 of the light on temperature, as reported in [15] for GSO
 241 crystals under X ray irradiation.

242 3.2. Scintillation time response of the GSO crystal

243 The same setup shown in fig. 6 with the addition of
 244 a layer of plastic scintillator positioned above the cry-
 245 ocooler, was used to check the possible dependence
 246 of the scintillation time upon temperature and applied
 247 magnetic field.

248 Signals from the scintillation of the GSO produced
 249 when the crystal is traversed by cosmic rays, were ac-
 250 quired in coincidence with signals in the plastic scin-
 251 tillator. Typically, more than one hundred PMT wave-
 252 forms from the GSO were recorded on a 1 GHz band-
 253 width digital oscilloscope, both at room temperature

254 and at $T = 13.5$ K. The observed signals decay expo-
 255 nentially with a decay time τ_m which depends on the
 256 temperature, as shown in Fig. 8, where the distributions
 257 of the measured signal decay time at the two tempera-
 258 tures are plotted. The time decays (τ_m) of 163 ± 10 ns
 259 and of 330 ± 20 ns were obtained at $T = 300$ K and
 260 $T = 13.5$ K respectively, averaging over a sample of 90
 261 events at each value of temperature. The observed de-
 262 cay time τ_m is related both to the response time of the
 263 apparatus τ_a , defined as the delay of the light signal due
 264 to the quartz waveguide and the PMT, and the scintilla-
 265 tion time of the GSO. Since the measured scintillation
 266 time of the GSO crystal is $\tau_s = 60$ ns [15], τ_a is ob-
 267 tained through expression $\tau_a = \sqrt{\tau_m^2 - \tau_s^2} = 152 \pm 20$
 268 ns at room temperature. Also it is reasonable to expect
 269 that this value is temperature independent and therefore,
 270 assuming the same response time for the two tempera-
 271 tures, a scintillation time of (293 ± 20) ns is inferred at
 272 $T = 13.5$ K.

273 Time measurements, at both temperatures, were re-
 274 peated without magnetic field and no observable vari-
 275 ations were observed at the 10% level. It should be
 276 noted that the longer decay time at lower temperatures
 277 may be attributed to a temperature effect on the physical
 278 mechanism of emission of scintillation light, as already
 279 reported by Mori *et al* [15].

280 3.3. Measurements using APD

281 The signal from GSO crystals in a magnetic field
 282 could be detected using APDs. It is not obvious how-
 283 ever that these devices work properly at cryogenic tem-
 284 peratures, since it is believed that their functioning at
 285 temperatures below 77 K is limited by the so-called
 286 freeze-out effect [18], especially if the material (n or p -
 287 type) is heavily doped.

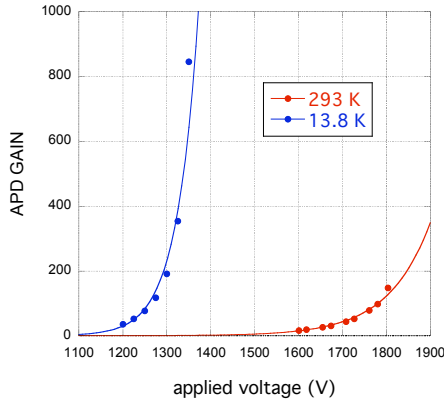


Figure 9: The measurement of the APD gain versus applied voltage at room temperature and at 13.5 K.

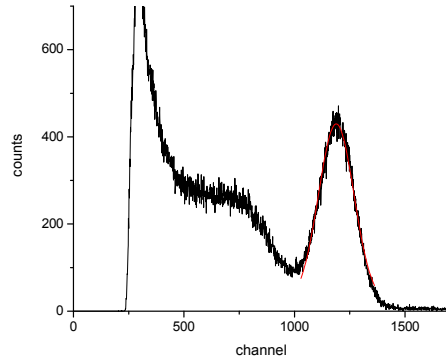


Figure 10: Cs^{137} source spectrum at room temperature. The voltage applied to the APD is 1800 V.

288 On the other hand, improvements are expected when the
 289 APD is made of high purity silicon, as suggested by re-
 290 cent results that were obtained with semiconductor de-
 291 tectors [21].

292 Therefore we carried out preliminary measurements on
 293 bare beveled silicon APDs from Advance Photonics, 16
 294 mm diameter, UV sensitive.

295 The gain of the APD was measured at $T = 300$ K and
 296 $T = 13.5$ K, using a 5.9 keV X-ray source (Fe), as func-
 297 tion of the applied voltage.

298 The results are shown in Fig. 9, in which the temper-
 299 ature dependence of the energy required to produce an
 300 electron-hole pair has also been taken into account to
 301 calculate the APD gain.

302 At the lower temperature, a limit of approximately 10^3
 303 in the gain is set by discharges, which appear when the
 304 applied voltage exceeds ~ 1400 V.

305 Nevertheless, the values of the gain allowed the acquisi-
 306 tion of the spectrum of a Cs^{137} γ -ray source, position-
 307 ed just above the GSO crystal, directly coupled to the APD
 308 which collected the light produced in the scintillation
 309 process. No optical grease was used to minimize re-
 310 flections at interfaces, therefore the amount of light col-
 311 lected would be smaller than expected if that precaution
 312 were adopted.

313 The spectra taken at two different temperatures ($T =$
 314 300 and 13 K) are shown in Fig. 10 and 11. As it was
 315 the case with PMT readout, in this experimental config-
 316 uration as well, the light yield at $T = 13.5$ K is measured
 317 to be a factor of two lower than at room temperature,
 318 thus confirming previously obtained results [15].

319 These measurements show that indeed it is possible to
 320 collect the scintillation light from GSO crystals using

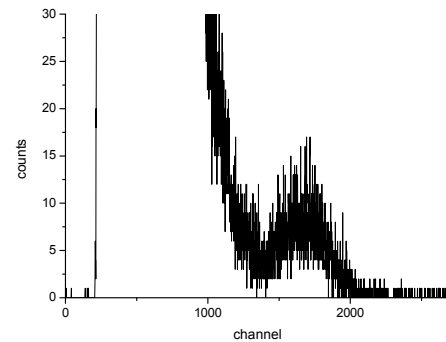


Figure 11: Cs^{137} source spectrum at 13 K. The voltage applied to the APD is 1325 V.

321 APDs kept at cryogenic temperatures.

322 4. Conclusions

323 In this work the feasibility of a new generation de-
324 tector for neutrino physics has been investigated. It has
325 been demonstrated that a GSO crystal kept at low tem-
326 peratures and in a high magnetic field is a suitable ac-
327 tive material. This was achieved through the systematic
328 study of its magnetic properties and optical response at
329 cryogenic temperature and in the presence of a magnetic
330 field up to 5 T. The scintillation properties of the polar-
331 ized material under gamma irradiation have been stud-
332 ied and the possibility to collect the light signal with a
333 photomultiplier tube or an avalanche photodiode kept at
334 low temperature, has been successfully tested.

335 Cosmic ray events have also been used to measure the
336 time response of the GSO crystal at low temperatures,
337 which is about a factor of five longer than the value at
338 room temperature [15].

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