

Special Article - Tools for Experiment and Theory

CUPID-0: the first array of enriched scintillating bolometers for $0\nu\beta\beta$ decay investigations

O. Azzolini¹, M. T. Barrera¹, J. W. Beeman², F. Bellini^{3,4}, M. Beretta^{5,6}, M. Biassoni⁶, C. Brofferio^{5,6}, C. Bucci⁷, L. Canonica^{7,8,16}, S. Capelli^{5,6}, L. Cardani⁴, P. Carniti^{5,6}, N. Casali⁴, L. Cassina^{5,6}, M. Clemenza^{5,6},

O. Cremonesi⁶, A. Cruciani^{3,4}, A. D'Addabbo⁷, I. Dafinei⁴, S. Di Domizio^{9,10}, F. Ferroni^{3,4}, L. Gironi^{5,6},

A. Giuliani^{11,12}, P. Gorla⁷, C. Gotti^{5,6}, G. Keppel¹, M. Martinez^{3,4}, S. Morganti⁴, S. Nagorny^{7,13}, M. Nastasi^{5,6},

S. Nisi⁷, C. Nones¹⁴, D. Orlandi⁷, L. Pagnanini^{7,13}, M. Pallavicini^{9,10}, V. Palmieri¹, L. Pattavina^{7,13,a}

M. Pavan^{5,6}, G. Pessina⁶, V. Pettinacci^{3,4}, S. Pirro⁷, S. Pozzi^{5,6}, E. Previtali⁶, A. Puiu^{5,6}, F. Reindl^{4,17,18},

C. Rusconi^{7,15}, K. Schäffner^{7,13}, C. Tomei⁴, M. Vignati⁴, A. Zolotarova¹⁴

² Materials Science Division, Lawrence Berkeley National Laboratory, Berkeley, CA 94720, USA

³ Dipartimento di Fisica, Sapienza Università di Roma, 00185 Rome, Italy

- ⁴ INFN, Sezione di Roma, 00185 Rome, Italy
- ⁵ Dipartimento di Fisica, Università di Milano-Bicocca, 20126 Milan, Italy
- ⁶ INFN, Sezione di Milano Bicocca, 20126 Milan, Italy
- ⁷ INFN, Laboratori Nazionali del Gran Sasso, Assergi, 67010 L'Aquila, Italy
- ⁸ Massachusetts Institute of Technology, Cambridge, MA 02139, USA
- ⁹ Dipartimento di Fisica, Università di Genova, 16146 Genoa, Italy
- ¹⁰ INFN, Sezione di Genova, 16146 Genoa, Italy
- ¹¹ CSNSM, Univ. Paris-Sud, CNRS/IN2P3, Université Paris-Saclay, 91405 Orsay, France
- ¹² DiSAT, Università dell'Insubria, 22100 Como, Italy
- ¹³ Gran Sasso Science Institute, 67100 L'Aquila, Italy
- ¹⁴ CEA-Saclay, DSM/IRFU, 91191 Gif-sur-Yvette Cedex, France
- ¹⁵ Department of Physics and Astronomy, University of South Carolina, Columbia, SC 29208, USA
- ¹⁶ Present address: Max-Planck-Institut fur Physik, 80805 Munich, Germany
- ¹⁷ Present address: Institut fur Hochenergiephysik der AW, 1050 Wien, Austria
- ¹⁸ Present address: Atominstitut, Technical University Vienna, 1020 Wien, Austria

Received: 1 March 2018 / Accepted: 14 May 2018 / Published online: 29 May 2018 © The Author(s) 2018

Abstract The CUPID-0 detector hosted at the Laboratori Nazionali del Gran Sasso, Italy, is the first large array of enriched scintillating cryogenic detectors for the investigation of ⁸²Se neutrinoless double-beta decay ($0\nu\beta\beta$). CUPID-0 aims at measuring a background index in the region of interest (RoI) for $0\nu\beta\beta$ at the level of 10^{-3} counts/(keV kg years), the lowest value ever measured using cryogenic detectors. CUPID-0 operates an array of Zn⁸²Se scintillating bolometers coupled with bolometric light detectors, with a state of the art technology for background suppression and thorough protocols and procedures for the detector preparation and construction. In this paper, the different phases of the detector design and construction will be presented, from the material selection (for the absorber production) to the new and innovative detector structure. The successful construct-

tion of the detector lead to promising preliminary detector performance which is discussed here.

1 Introduction

Scintillating cryogenic detectors are excellent devices for rare events investigations. Their use was first proposed in 1989 for the detection of solar neutrinos [1]. However, large target masses were needed, and the technology was yet not enough mature. Bolometers are nowadays extensively used both for applied physics [2] and fundamental physics [3].

One of the main challenge for next generation bolometric experiments is to increase the experimental sensitivity using larger mass detectors with lower background level in the region of interest (RoI). This is the case of CUORE [4], the first-ever ton-scale bolometric experiment searching for $0\nu\beta\beta$. CUORE, after 5 years of data taking, will reach a sen-

¹ INFN, Laboratori Nazionali di Legnaro, 35020 Legnaro , Padua, Italy

^ae-mail: luca.pattavina@gssi.it

sitivity of 9×10^{25} years [5] to the observation of 130 Te $0\nu\beta\beta$. Its limitation is given by the expected background index in the RoI which will be 0.01 counts/(keV kg years) [6], mainly ascribed to α -particle interactions on the detector surfaces.

CUPID [7,8] (CUORE Upgrade with Particle IDentification) aims at developing the technology of scintillating bolometers for the realization of a next generation $0\nu\beta\beta$ experiment with sensitivity of 10^{27} – 10^{28} years, depending on the isotope of interest. This goal establishes some technical challenges, the most relevant one is the operation of a ton of isotope with close-to-zero background level for a ton×year exposure [9] in the RoI around the $\beta\beta$ transition energy.

CUPID-0 (formerly LUCIFER [10]) is the first demonstrator of such technology, operating an array of 26 scintillating bolometers of Zn⁸²Se, 24 enriched at 95% and 2 naturals. One of the milestones of CUPID-0 is to demonstrate the feasibility of a close-to-zero background experiment, about 10^{-3} counts/(keV kg years), one order of magnitude better than CUORE. CUPID-0 is pursuing this goal using scintillating bolometers which allow the identification of the type of interacting particle, thus the rejection of the α -background [10–12], at a 12 σ level [13].

The detector is installed in the Hall A of the Gran Sasso Underground Laboratory (LNGS) of INFN, sited in Italy. This unique location ensures an effective shielding against high energy cosmic rays of about 3600 m.w.e..

In this work we describe in detail all the procedures for the realization and operation of the CUPID-0 detector, from the production of the fundamental units, the scintillating bolometers, to the processing of the thermal sensors for the signal read-out, and also the surface treatment for the reduction of surface contaminations. A review of the detector operations and performance will also be discussed.

2 Operation of scintillating bolometers

A scintillating bolometer is a scintillating crystal absorber which is operated as highly sensitive calorimeter at low temperature. The absorber is kept at cryogenic temperature, few tens of mK, in order to minimize its heat capacity¹. In these conditions an energy deposit induces a sizeable temperature variation measured by means of a Ge Neutron Transmutation Doped (NTD) thermistor [14]. This induces a perturbation of the crystal lattice which is mediated by phonons, these have energies of the order of few μ eV. Given that the RoI is at few MeV, the statistical fluctuation of the mediators is extremely small, thus allowing for an excellent energy resolution over a

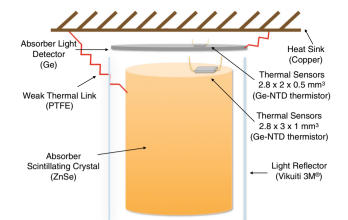


Fig. 1 Schematic view of a single module scintillating bolometer

wide energy range, at the level of several parts per thousand over a few MeV.

When the absorber is also a good scintillator at low temperature, a fraction of the deposited energy in the absorber is converted into a light signal. This can be read out by a suitable light detector (LD) facing the crystal. By means of the read-out of heat and light signal the identification of the type of interacting particle is feasible, thus allowing for the rejection of α particle interactions. Currently, the best choice for light detection in such ultra low background and low temperature environment is the operation of an auxiliary bolometer. CUPID-0 uses Ge absorbers equipped with a thermal sensor similar to the one used in the main absorber.

In Fig. 1 a schematic view of a CUPID-0 single module is shown. The ZnSe crystals and the Ge LDs are thermally coupled to the heat bath by means of a copper structure. The two absorbers are held in position by means of PTFE clamps. These act also as weak thermal link to the heat sink and at the same time compensate for the different thermal contractions of the absorbers and of the copper. The difference in mass between the ZnSe, about 450 g, and the LD, about 1 g, requires the design of different thermal sensors which must take into account the different heat capacities. The entire setup is enclosed in a light reflector which helps in maximizing the light collection efficiency and thus to make the particle identification more efficient.

3 Design of the CUPID-0 detector array

The CUPID-0 detector is completely different from any other previously designed bolometric experiment due to its high degree of complexity: large number of channels, extremely compact structure and different absorber dimensions². This

¹ The heat-capacity of a dielectric and diamagnetic crystals scales with the third power of the ratio between its temperature operation and its Debye temperature.

 $^{^2\,}$ ZnSe crystals and LDs have different dimensions, but also the height of each ZnSe crystal varies.

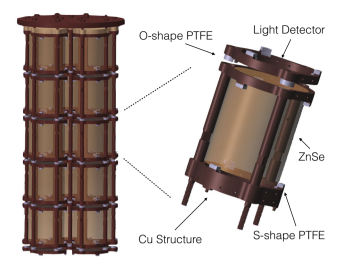


Fig. 2 Rendering of CUPID-0 detector and its single module

forced the collaboration to the design of a detector structure which had to be reliable, flexible and light at the same time.

CUPID-0 is an array of 26 ZnSe crystals, 24 highly enriched in ⁸²Se at a level of 95% and 2 naturals³. All the crystals are arranged in a tower like structure and in total there are 5 towers, four containing 5 crystals and one with 6 crystals. The position of the crystals inside the different towers is done in a way such that each tower has about the same weight (height) of about 2 kg (30 cm). The overall number of $\beta\beta$ nuclei included in the CUPID-0 detector is 3.8×10^{25} (natural + enriched crystals).

Each tower is about 30 cm tall, the ZnSe crystals are interleaved with Ge-LDs. As it is shown in Fig. 2, each ZnSe is faced to two LDs, one on the top and one on the bottom, so that there is redundancy in the light signal read-out in case of any issue related to their performance.

In order to minimize the amount of passive material next to the detector, which may induce a β/γ background in the RoI due to its internal radioactive contaminations, the only components selected for the detector construction are: copper, PTFE and light reflectors (VIKUITI-3M). All of the Cu pieces were machined from large Cu chunks from NOSV Cu of Norddeutsche Affinerie AG⁴.

The structure is composed by copper frames and columns. The innovative idea for making the simplest possible structure was to use a single copper frame both as ZnSe and LD holder, as it is shown in Fig. 2. In fact the top part of the circular copper component on its upper part holds the crystal, while the bottom part acts on the LD. Progressively on the top of the crystal there is a second frame (like the previous one, but inverted) where on its top part a new LD is set in place. Using this new type of design for each ZnSe and its adjacent LD only two copper frames are needed. The frames are kept together by means of 3 copper columns, whose lengths is specifically defined by each crystal height. We were able to push down the overall amount of copper in the detector structure and ancillary parts to 22% of the overall detector mass, an unprecedented value compared to any previous bolometric experiment.

The second material employed for the detector construction is PTFE, which was used to secure in place the two absorbers. This was machined in two sets of pieces: one with a S-shape for the ZnSe and one with a O-shape for the LD. The ZnSe crystals are supported with three S-shaped PTFE holders on the bottom and three on the top, while the LDs are clamped by three O-shape holders by means of a narrow slit made on the perpendicular axis of the holder. The overall fraction of PTFE in the structure amounts to 0.1%.

The last component used for the realization of the CUPID-0 detector is the light reflector VIKUITI from 3M. This plastic foil is shaped in cylinders and placed around each crystal, to maximize the light collection efficiency on the two LDs. The reflector completely surrounds the crystals avoiding any line of sight between the absorbers and the closest cryostat radiation shield at 50 mK. This allowed us to prevent the installation of a massive copper shield around the detector at ~10 mK, reducing by a large fraction the copper mass next to the detector.

4 Detector components

Each component employed for the detector construction was specifically selected for its intrinsic radiopurity. After material screening for each item a dedicated cleaning and purification technique was adopted, the final goal was to mitigate and prevent any recontamination. This section contains a detailed description of how each detector component was selected and handled before its final installation in CUPID-0.

4.1 Zn⁸²Se crystal absorbers

For the first time ever large mass Zn⁸²Se crystals enriched in ⁸²Se were grown. The scintillating elements were produced at the Institute for Scintillating Material at Kharkov (Ukraine).

The enriched Zn⁸²Se crystals were produced starting from highly pure raw materials, namely metal ⁸²Se and ^{nat}Zn. The radiopurity of these two metals was investigated at LNGS by means of γ -spectroscopy on a p-type HP-Ge detector [16]. This detector is characterized by an extremely low intrinsic background that allows the investigation of radioactive contaminations with extremely high sensitivity. In Table 1 we show the results of the γ -spectroscopic analysis of the two metals used for the crystal production.

³ The natural isotopic abundance of 82 Se is 8.82% [15].

⁴ Now Aurubis AG.

Table 1 Internal radioactive contamination for 2.5 kg of 96.3% enriched ⁸²Se metal beads and for 2.5 kg of ^{*nat*}Zn. Limits are computed at 90% C.L.. The measurements were carried out on October 2014

Chain	Nuclide	⁸² Se activity [μBq/kg]	^{nat} Zn activity [μBq/kg]
²³² Th	²²⁸ Ra	< 61	< 95
	²²⁸ Th	< 110	< 36
²³⁸ U	²²⁶ Ra	< 110	< 66
	²³⁴ Th	< 6200	< 6200
	^{234m} Pa	< 3400	< 4700
²³⁵ U	²³⁵ U	< 74	< 91
	⁴⁰ K	< 990	< 380
	⁶⁰ Co	< 65	< 36
	⁵⁶ Co	-	80 ± 20
	⁶⁵ Zn	-	5200 ± 600

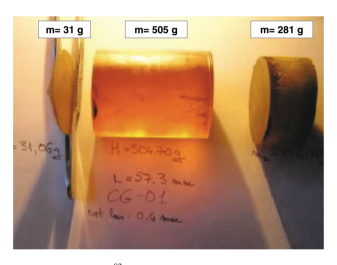


Fig. 3 Picture of a $Zn^{82}Se$ ingot as grown. The two edge of the boule are removed, while the central part is processed for the realization of the final crystal for the CUPID-0 detector

As described in [18], the synthesis of $Zn^{82}Se$ is made in vapour phase by evaporating Zn and enriched ⁸²Se in an Ar atmosphere at 950 °C. The synthesised powder then, after a two stage purification procedure, the first in Ar and the second in H₂ atmosphere, is charged inside a high density graphite crucible. All these procedures are performed in Ar flushed disposable glove-boxes, in such a way that the material is never exposed to air so to reduced any possible recontamination.

The 1 kg charge is sufficient for the production of a single crystal of 500 g, the rest is all recoverable material which is not included in the final crystal production given its poor crystalline quality, see Fig. 3. The crystal is grown using the Bridgman technique at 1500 $^{\circ}$ C at about 15 MPa of Ar pressure, with a growing speed of about 1.5 mm/h.

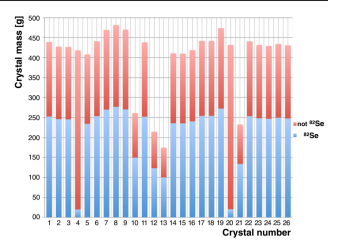


Fig. 4 Crystal masses. The ⁸²Se content for each crystal is shown. Crystal number 4 and 20 are crystal with natural Se isotopic abundance

The final crystal is then shaped and optically polished with specific materials and procedures, for further details on the crystal polishing and lapping see [18]. All these delicate procedures were carried out inside a clean-room where a Radon-abatement system was installed in order to reduce any possible recontamination of the crystals after the polishing. The ²²²Rn concentration inside the clean room is $<20 \text{ mBq/m}^3$ [17].

In Fig. 4, the masses of all CUPID-0 crystals and the fraction of the $0\nu\beta\beta$ source are shown. They range from about 150 g to about 500 g, due to the difficult growth conditions. From each ingot, the crystal cuts were performed so that the crystal mass and its quality were maximized. The two crystals with the lowest content of Se are the natural ones.

4.2 Ge light detectors

Operating reliable and robust LDs is of paramount importance, since the particle identification relies on the performance of these devices.

The Ge substrate/absorber, purchased at UMICORE Electro-Optic Material (Geel, Belgium), is a double side polished wafer of diameter 44.5 mm and an average thickness of $170 \,\mu$ m, with an impurity concentration $< 2 \times 10^{10}$ atoms/cm³.

All the long-standing experience in the development of cryogenic LDs [19] helped in defining the critical issues for an efficient particle identification and rejection, above all: energy resolution and signal amplitude. The former is strongly dependent on the operating conditions of the thermal sensor, as it is thoroughly discussed in [20]. The latter mostly depends on the detector design and on the ability to maximize the light detection efficiency. For this reason a dedicated procedure for the enhancement of the light collection efficiency using an anti-reflective coating was developed at

CSNSM (Orsay, France). Furthermore, a reflecting foil was employed for focusing on the LD the scintillation light coming from the ZnSe crystal.

A way to significantly improve the LD performance is to increase the light collection by minimizing its reflectivity. Such improvement can be achieved by means of a special anti-reflective coating on the side of the detector which is facing the scintillating bolometer.

One of the simplest methods to reduce the reflection is the so called *refractive index matching*. In the approximation of normally incident light from a transparent to an absorbing medium, the absorbed fraction can be calculated by the following formula:

$$R = \frac{(n_0 - n_1)^2 + k^2}{(n_0 + n_1)^2 + k^2} \tag{1}$$

where n_0 is the real refractive index of the transparent medium ($n_0 = 1$) and $n_1(k)$ is the real (imaginary) part of the complex refraction index of the absorbing medium, for germanium $n_1 = 5.48$ (k = 0.82) at 632 nm at room temperaure. If we consider a simple vacuum-germanium interface, the fraction of absorbed light is only 48% at 632 nm [21]. On the contrary, if a thin coating layer with refractive index n_i , with value between n_0 and n_1 is placed between the two media, we should evaluate first the *R* value for the vacuumcoating interface, then for the coating-germanium transition. The optimum value for anti-reflective material to be placed on the Ge detector is a material with $n_i \sim 2.4$ which leads to a gain on the light absorption of about 35% with respect to bare Ge, depending on the wavelength.

The best thickness for the layer can be determined by fulfilling the conditions for an optimal anti-reflective coating in the approximation of single-layer interference. The coating thickness should be $d = \lambda/4$, where λ is the wavelength of the incident light. This method works well for monochromatic light sources. In our case, we can take $\lambda = 645$ nm, which corresponds to the maximum of the intensity emission for ZnSe scintillation, even though the wavelength distribution is rather broad [22].

Several cryogenic tests on anti-reflective coatings were performed at LNGS and at CSNSM during the past years [23]. The best results were achieved with a SiO coating, which has a refractive index of ~ 2.5 . The optimal thickness is approximately 65 nm for a SiO coating on a Ge absorber. The increase of the absorbed light fraction was 34%, in agreement with the expectations. For these reasons a SiO coating was deposited on CUPID-0 LDs.

Forty Ge absorbers were prepared at CSNSM for CUPID-0. Before the SiO deposition each Ge wafer was previously etched with a mixture of nitric (HNO₃ 70%), acetic (CH₃COOH 100%) and hydrofluoride (HF 40%) acids in proportion 5:3:3. This mixture is aggressive enough to react

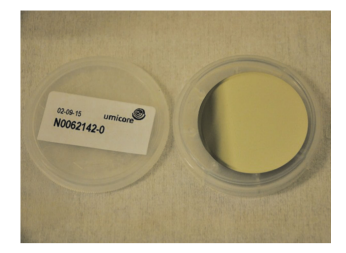


Fig. 5 A high purity Ge slab before applying any procedure

with the Ge surface. The thickness of the removed surface layer was $\approx 10 \ \mu m$ with an etching time of about 1 min.

After the chemical etching the surface was also treated with an Ar ion bombardment. The gas was ionised with an electron gun, the Ar pressure during the bombardment was 3×10^{-3} mbar. These procedures remove any possible residual oxides and improve the surface quality for the coating process. The deposition is performed using a tantalum box, where the SiO is heated up to T ~ 1000 °C. The deposition is performed under vacuum: the pressure in the evaporation chamber is P< 10^{-7} mbar (see Fig. 5).

The evaporation rate was tuned to be in the range of 0.5-1 nm/s. The deposition thickness is controlled with a high precision (< 0.1 nm) piezoelectric quartz crystal; its resonance frequency depends on the deposited mass. The average final thickness of the SiO coatings was between 70 and 80 nm. This is a good approximation of the value required by the single-layer interference anti-reflective coating, as previously discussed (see Fig. 6).

4.3 VIKUITI light reflector

The light reflector foil installed around the crystal is a VIKUITI multi-layer specular reflector produced by 3M. The foil has been characterized at different temperatures and it ensures a reflectivity greater than 98% for wavelengths between 400 and 800 nm [24] over a wide temperature range, from 300 to 20 K.

The emission spectrum of ZnSe at 10 K has different components, the most intense is at 645 nm [22], thus ensuring an excellent light collection efficiency.

The reflector radiopurity was investigated with ICP-MS at the LNGS employing an innovative mineralization procedure for the sample preparation [25]. The measured concentration

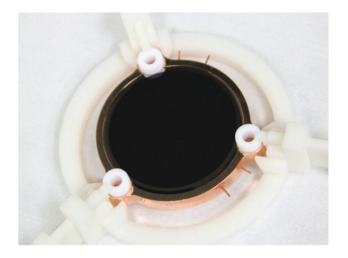


Fig. 6 Light detector before its assembly. The side with the antireflective coating is visible. The coating results in a dark internal circle, while 2 mm on the edge are uncoated

Table 2 Bulk contaminations of the VIKUITI-3M light reflector. The measurement was carried out at the LNGS by means of a mineralization procedure and an ICPMS analysis. The overall mass of VIKUITI-3M used in CUPID-0 is 17 g. The total activity is $0.8 \pm 0.2 \mu$ Bq and $2.9 \pm 0.7 \mu$ Bq for ²³²Th and ²³⁸U, respectively

Chain	Nuclide	VIKUITI-3M activity [µBq/kg]
²³² Th	²³² Th	49 ± 12
²³⁸ U	²³⁸ U	170 ± 50

of the elemental Th and U were 12 ± 3 and 14 ± 4 ppt, respectively, see Table 2.

4.4 Ge-NTD thermal sensor

The CUPID-0 thermal sensor production started in March 2012. Nine wafers of high-purity Ge, $\emptyset = 65.5$ mm and thickness 3.25 mm, were irradiated at the MIT Nuclear Reactor Laboratory, Boston (MA, USA), see Fig. 7. The exposure to high neutron fluxes is needed in order to uniformly dope the wafers. The required dopant concentration to enable the operation of the Ge as thermal sensor is at level of 10^{16} atom/cm³. Such high and uniform doping level will take the semiconductor close to the metal-to-insulator region. The electrical conductivity of these heavily doped semiconductors has an exponential dependence on the temperature, making these sensors the most suitable technology for our purposes [14]:

$$\rho(T) = \rho_0 e^{(T_0/T)^{0.5}},\tag{2}$$

where T_0 depends on the Ge-NTD doping level and ρ_0 on the doping level and on the sensor geometry. As a consequence, a fluctuation of the doping level will strongly affect the sen-

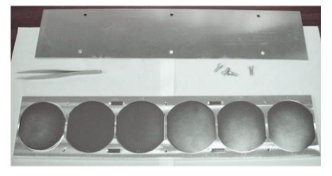


Fig. 7 Six out of 9 Ge wafers, diameter of 65.5 and thickness 3.25 mm, installed in an Al holder before neutron irradiation at the MIT Nuclear Reactor Laboratory (MA, USA)

sitivity of the thermometer. The target values are $T_0 = 4.2$ K and $\rho_0 = 1.5 \Omega$ for a sensor sensitivity of about 1 M Ω/μ K.

An accurate and precise measurement of the flux and neutron dose to which the Ge wafers are exposed is of paramount importance for the success of the experiment. These values will depend the detector response and resolution. The entire neutron dose should not exceed 2% of the nominal value, which is estimated to be at the level of 4×10^{18} n/cm³.

To achieve such high accuracy and precision nominal neutron dose, we decided to irradiate: 3 wafers at $\pm 7\%$ of the target dose, 2 wafers at $\pm 3\%$, 1 wafer at $\pm 2\%$ and 3 wafers at the nominal value. This choice was driven by the fact that the Nuclear Reactor facility could ensure dose within 5% of the target value. While irradiating the Ge wafers some witness samples were also exposed to the same neutron flux. These were used for an accurate off-line evaluation of the total exposure, by means of Neutron Activation Analysis [27]. At last, a fine tuning of the neutron dose was carried out at the LENA Nuclear Research Reactor Laboratory (Pavia, Italy). The wafers that received a neutron dose closest, but lower, than the nominal value underwent a neutron exposure at the LENA reactor. The LENA most intense neutron flux is about 3 orders of magnitude lower than the MIT reactor flux, thus allowing a more accurate neutron irradiation. A detailed mapping of the LENA reactor neutron fluxes was performed [26] before irradiating the sensors. This allowed us to select the irradiation channel more suitable for our purposes, thus with a ratio thermal/fast neutron flux of about 20.

On the two large area of the Ge wafer a 4000 Å gold layer is deposited, which serves as sensor Ohmic contacts, see Fig. 8. After the deposition the wafer is diced in sensors of the desired size. The final sensor dimensions for the ZnSe and LDs are $3 \times 2.8 \times 1 \text{ mm}^3$ and $2 \times 2.8 \times 0.5 \text{ mm}^3$, respectively, see Fig. 9, where 2.8 mm is the distance between the two golden pads.

The sensors, before their coupling to the absorbers, are equipped with gold wires. A dedicated ball-bonding machine



Fig. 8 Doped Ge wafer with a 4000 Å gold deposition. The Au on the two large surface of the wafers serve as Ohmic contacts for reading-out the sensor

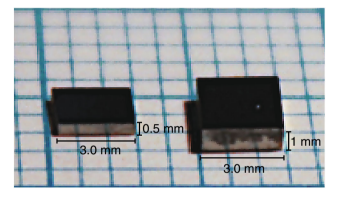


Fig. 9 Ge Neutron Transmutation Doped sensors for ZnSe (right) and light detector (left). The dimensions for the two sensors are $3 \times 2.8 \times 1 \text{ mm}^3$ and $2 \times 2.8 \times 0.5 \text{ mm}^3$, respectively

was used for the bonding of gold wires of 25 μm of diameter on the golden Ohmic contacts.

4.4.1 Sensors gluing

The mechanical and thermal coupling of the sensors to the crystal absorbers (referred to as gluing) is a well-known concern in the construction of bolometers because it influences the quality of the detector performance. In particular, the R&D towards the Cuoricino experiment [28] established stringent requirements to this delicate process, involving the geometry of the glue interface between sensor and absorber, the selection of glue, and the environmental conditions (temperature and humidity) in which the operation has to be performed.

According to these constraints, it is preferable to deposit the glue in a matrix of dots, in order to compensate for the different thermal contractions at low temperatures. The most suitable number of dots for a $2.8 \times 3 \text{ mm}^2$ surface Ge-NTD is 9. These dots must have a diameter of 0.7 mm, while their height is determined by imposing a 0.05 mm gap between the crystal and the absorber, see Fig. 10.

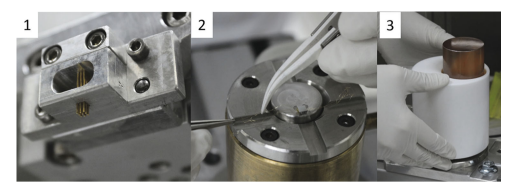
Moreover, the glue is deposited on the sensor surface instead of on the crystal, because the former is easier to reprocess in case of gluing failure, while the cleaning of the crystal surface may require an entire crystal surface polishing treatment.

The glue used for the process must have a density high enough to avoid the merging of the dots after their deposition. Besides that, it has to work at cryogenic temperatures through several thermal cycles, and it must fulfil the radiopurity constraints required by the experiment. The selected glue is the bi-component epoxy Araldite Rapid by Huntsman Advanced Materials [29]. It has a viscosity of 30 Pa s, a very quick pot life of about 3 min, but also a very short curing time of about 1 h and low radioactivity, less than 8.9 $\times 10^{-4}$ Bq/kg for ²³²Th and 1.0×10^{-2} Bq/kg for ²³⁸U [30].

To improve the reproducibility of the detector performance, a R&D was carried on to develop a semi-automated system for the sensor-to-absorber coupling of the CUPID-0 bolometers, similar to the one developed for the CUORE experiment [31]. The CUPID-0 gluing system was developed starting from the previously acquired knowledge [32]. The result was a semi-automated system in which the dots are placed through a matrix of spring-loaded tips applied to a x-z Cartesian robot, see Fig. 10-1. The introduction of automated elements reduces the variability induced by manual work and gives the advantage of precise timing, which is useful considering the short epoxy life-time.

The gluing procedure is divided in three steps: the tool preparation, the glue dispensing and the crystal deposition. Among the three, only the second one is automated. Firstly, the Cartesian robot is equipped with the correct spring-loaded tip matrix according to the kind of Ge-NTD to be glued (a nine tip matrix for the $2.8 \times 3 \times 1 \text{ mm}^3$ Ge-NTDs for ZnSe crystals and a six tip matrix for the $2.8 \times 2 \times 0.5 \text{ mm}^3$ Ge-NTDs for LDs). Then the sensor is placed on a positioning device on the Cartesian robot, where it is held in place by vacuum, see Fig. 10-1 and -2. In parallel, the crystal is prepared in a dedicated PTFE holder that will be inserted onto the sensor positioning device after the glue handling phase. This begins with the mixing of the two epoxy components through a dispensing gun provided with a disposable static mixer; the mixed glue is poured and then levelled in a small PTFE container placed on the Cartesian robot. The x-z arm dips the tip matrix in the glue container and then presses it onto the sensor surface to deploy the glue dots. The correct size of the single glue dot is determined by choosing a proper diameter of the tip (0.53 mm) in combination with the depth of the glue container (0.70 mm). The fact that the tips are spring-loaded ensures a uniform collection/deposition of the glue by each tip.

Fig. 10 Sensor gluing: (1) Matrix of spring-loaded tips used for the $3 \times 2.8 \times 1 \text{ mm}^3$ NTDs. (2) Placing NTD on the positioning device and held by vacuum. The Spring-loaded tip matrix is lowered on the NTD after dipping in the glue. After that, (3) the ZnSe crystal is placed on the top of the NTD and it is kept in place by a PTFE holder



Finally, the crystal is lowered on the Ge-NTD thanks to the PTFE crystal holder in which it was previously hosted, see Fig. 10-3, that ensures a gap of 0.05 mm between the crystal and Ge-NTD surfaces. This, together with the fact that the sensor is held by vacuum for all the glue curing time prevents the dots to merge in a layer, preserving the shape of the dot matrix.

All the absorbers are also equipped with a Si resistor which are used for injecting fixed amount of energy in the crystals [33]. These produce signals similar to the one induced by particle interaction, and they are used for the correction of the detector gain drift caused by the continuous cooling down of the experimental apparatus.

The gluing activity of CUPID-0 was performed inside a Radon-free cleanroom to ensure low radioactive conditions ($< 20 \text{ mBq/m}^3$) and a very stable environment in terms of temperature and humidity, mandatory conditions since these parameters influence the glue's intrinsic properties (especially viscosity).

4.5 Ultra-clean copper

The CUPID-0 detector structure, is mainly composed by copper, it makes about 22% of the overall detector mass. Minimizing the concentration of radioactive impurities, especially from the surface of the detector structure is important for suppressing possible high-energy β/γ background sources. For this reason a dedicated cleaning procedure was developed for the abatement of surface contamination in copper, similar to the one described in [34].

The frames that hold the crystals are made of electrolytic tough pitch copper, also known as NOSV copper. Some impurities, including decay products of ²³²Th and ²³⁸U decay chain can accumulate on the material surface as a consequence of an exposure to an uncontrolled atmosphere. Additional impurities are also deposited during several mechanical machining steps needed to produce the copper holders. The concentration of contaminant in the material is usually modelled as a gradient from the first external layer to the inner bulk, caused by diffusion. The radioactive contaminants of ²³²Th and ²³⁸U are usually present on copper surfaces to a depth of about 20 μ m [35].

All the NOSV Cu components were cleaned within a period of 6 months. The cleaning protocol consists of five general macro steps divided in sub-steps for a total of 61 individual processes. The time required for cleaning one set was about ten days. The total number of pieces cleaned was 268: 78 columns (with 26 different lengths), 70 frames, one large copper plate for the tower installation, and several spare parts.

4.5.1 Cleaning process protocol to reduce the radioactive contamination levels in Cu components

The cleaning procedure, developed at the Legnaro National Laboratories (LNL) of INFN consists of a sequence of successive treatments: Tumbling, Electropolishing, Chemical etching and Magnetron plasma etching (T+E+C+M). The storage of the Cu parts after each cleaning step was performed in a clean-room to avoid possible re-contamination of the surface.

- Pre-cleaning process: the pre-cleaning is performed for removing any lubricant residues deposited on the copper surfaces, and it directly affects the efficiency of the electropolishing process. It is performed wiping the Cu surface using specific wipes and a sequence of three different solvents: tetrachlorethylene to disolve organic materials, acetone to degrease and remove tetrachloroethylene, and ethyl alcohol to dissolve the acetone from the Cu surface. Later, the copper pieces are cleaned in ultrasound baths (33 kHz) at 40 °C for 10 min with deionized water and NGL 17.40 P.SP powder soap. Right after the bath, the copper pieces are dried with alcohol and nitrogen taking precautions to avoid re-contamination.
- Chemical etching pre-electrochemical process: the Cu surfaces must be prepared for the electropolishing process. Tumbling was used for CUPID-0 pieces, except for some delicate parts that cannot undergo the tumbling process, due to the high precision machining and small holes (less than 1 mm). In this case the tumbling step cleaning protocol is substituted by an ammonium persulfate chemical treatment with a concentration of 20 g/L for 2 h.

- *Electrochemical process*: the electropolishing can remove surface layer up to 50 μ m for the frames and 100 μ m for the other parts. To avoid the removal of Cu in specific area of the detector components such as the threads, PTFE protections were used. The Cu pieces are placed inside an electrochemical solution of 40% of butanol – 60% of phosphoric acid, using a specifically designed anode for each type of Cu component. During the process, the amount of Cu etched from the surface is monitored through a tuning of the current flowing inside the electropolishing solution and through the time spent inside the acid bath. After the electrochemical treatment, the residual electropolishing solution is removed with ultrasonic cleaning.
- Chemical etching process and Passivation: the chemical etching is applied to reduce the radioactive contaminants from the areas screened with the PTFE protections. The erosion rate is about 2 μ m/min. This chemical etching is performed using deionized water heated at 72 ± 4 °C with a recipe of sulfamic acid, ammonium citrate in powder form adding hydrogen peroxide and butanol in liquid form, named SUBU. The copper pieces are fixed to a sample holder through a copper wire and drawn in the SUBU solution for 5 min, the pieces rotate to enhance the reaction rate. After the SUBU process, the copper pieces are passivated in sulfamic acid solution at a concentration of 20 g/L for 5 min and cleaned in ultrasonic bath to finally be dried and packed.
- Plasma cleaning: the plasma cleaning constitutes an important step in the cleaning protocol. It is carried out in a vacuum system and it is the last phase before the assembly of the detector. The erosion rate during plasma cleaning is about 1µm/h. The plasma cleaning is a process based on DC magnetron sputtering technique which consists in the erosion of a target (copper pieces) through the impact ions of Ar gas (plasma). The copper pieces are fixed on a different sample holder for each kind of component, inside of a class 100 cleanroom in order to handle the copper pieces in a controlled environment. The holder is placed in the vacuum chamber and before the plasma cleaning, 12 h of baking at 100 °C is performed in order to degas adsorbed chemical substances and humidity. After the baking, a uniform magnetic field of 1.5×10^{-2} T is generated and a 30 W power plasma is supplied for 5 min.

5 Detector assembly

All activities for the construction of CUPID-0 detector were carried out in an underground Rn-suppressed clean room, with a Rn contamination of less than 20 mBq/m³ located in the Hall C of the LNGS. The cleanroom contained two



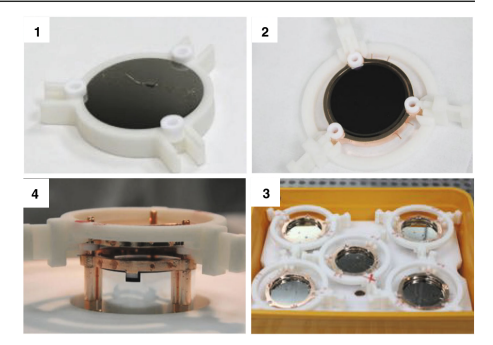
Fig. 11 Photos of the two working areas inside the cleanroom. On the left is shown the station were the Ge-NTD thermal sensors are glued to the absorbers, and on the right is shown the area where the tower are assembled

separate workstations: one for gluing the sensors to the ZnSe crystals and to the Ge wafers (see Fig. 11, left), and one for building and instrumenting the towers (see Fig. 11, right). A nitrogen-flushed storage container was also installed in the clean room for hosting the detector after its assembly.

The first step in the CUPID-0 tower construction was the pre-assembling of the LDs: each Ge wafer equipped with Ge-NTD was held in its copper frame using three PTFE Oshape holders. Since the Ge wafers are $170 \,\mu m$ thick specific assembly tools were developed in order to avoid accidental damage and recontamination of the copper frames during the assembly procedures. These tools were made with a ENVI-SIONTEC ULTRA 3SP 3D printer using a highly radiopure plastic resin (see Table 3). These tools consisted of a mounting template and of a handling system (see Fig. 12-1 and -4 respectively). The assembly procedure for the 31 LDs of the CUPID-0 detector consisted in the following steps: positioning of the Ge wafer with the PTFE O-shape holders on the mounting 3D-printed template, installation of the Cu frame on the LD and connection of the Ge-NTD gold wires to the Cu frame for the sensor read-out, fitting and tightening of the

Table 3 Radioactivecontamination of thepolymer-resin used for the	Chain	Nuclide	Activity [mBq/kg]
3D-printing of the detector	²³² Th	²²⁸ Ra	< 9.3
assembling tools (3SP WHITE D7)		²²⁸ Th	< 10.3
	²³⁸ U	²²⁶ Ra	< 3.8
		²³⁴ Th	< 73
		^{234m} Pa	< 0.25
	²³⁵ U	²³⁵ U	< 5.3
		⁴⁰ K	81±35

Fig. 12 Light detectors assembly: (1) Positioning of the Ge wafer with the PTFE O-shape holders on the mounting template. (2) Positioning of the copper holder and connection of the Ge-NTD gold wires for the detector read-out. (3) Storage of the assembled light detectors in a vacuum box. (4) Fit and tighten of the handling tool and removal from the mounting template



handling tool and removal from the mounting template and finally their storage of the assembled LDs in a vacuum box, see Fig. 12.

The second step in assembling the CUPID-0 detector was to physically assembly the towers using Cu, PTFE, 26 crystals and 31 pre-assembled LDs. The towers were manually built, one floor at the time ⁵, starting from the lower one. In order to maintain a suitable operational working height, the towers were assembled in an automatically adjustable table, named garage. After the assembly of the first floor, the tower is lowered by the height of this floor, in this configuration the operator always works at the same level.

The main steps for the assembly of a single tower are shown in Fig. 13. The first operation is the positioning of the first pre-assembled LD on the bottom copper holder of the tower and the installation of the first three columns that will host the ZnSe crystal, then the positioning of the ZnSe crystal on the bottom copper frame equipped with three Sshaped PTFE clamps. The third step is the installation of the VIKUITI-3M reflector and the top copper holder equipped with three S-shaped PTFE clamps. At this point the electrical connections between the gold-wire and the mechanical structure are finalized. The wires are crimped into the insulated copper tubes glued into the frames. Finally the second LD is installed on the top frame of the previously installed ZnSe crystals. This procedure is repeated until a tower of 5 (or in one case 6) floors.

After the completion of the assembly of the five towers, they are installed on a copper plate which is directly coupled to the cryogenic system, see Fig. 14.

6 Cryostat

The CUPID-0 cryostat is the same cryogenic infrastructure that hosted the CUORICINO [28] and the CUORE-0 [36] detectors. This system was upgraded in order to meet our stringent requirements in terms of low vibrational environment and increased number of read-out channels. The cryostat was commissioned at the LNGS in 1988 and it is a Oxford TL1000 dilution refrigerator with a copper He dewar. The ³He/⁴He dilution unit has a cooling power at 100 mK of about 1 mW. This ensures the possibility to install in the system a large number of read-out channels without spoiling the cryostat performance in terms of cooling power.

In Fig. 15, the cryogenic system hosting the CUPID-0 detector is shown. The detector is installed right below a Roman Pb shield by means of a spring and it is thermally coupled to the Mixing Chamber stage (MC) by means of a high-purity (99.999%) copper foil of 50 μ m thickness. The MC, which is the coldest point of the system, ensures the detector cooling down to the designed base temperature of 7.5 mK.

Two major upgrades were implemented in the system compared to the previous configuration: increasing the number of read-out channels and installing a mechanical decoupling system as anti-vibrational damping system. Upgrading the number of read-out channels was a mandatory step given the fact that for each ZnSe crystal there is also a LD; this doubles the number of wires from room temperature down to the detectors. When hosting the CUORE-0 experiment, the cryostat was able to read out up to 52 bolometers; now the system can handle up to 136 detectors, irrespectively if they are ZnSe crystals or LDs. For CUPID-0 67 channels are used: 26 for ZnSe crystals, 31 for LDs and 10 thermometers

 $^{^5\,}$ A floor is defined as a single module, this is composed by a $Zn^{82}Se$ and its most adjacent LD.

Fig. 13 Single tower assembly: a positioning of the first pre-assembled LD on the bottom copper holder of the tower; installation of the first three columns that will host the ZnSe crystal. b Positioning of the ZnSe crystal on the bottom copper frame equipped with three S-shaped PTFE clamps. c Positioning of the VIKUITI-3M reflective foil. d Positioning of the top copper holder equipped with three S-shaped PTFE clamps; connection of the Ge-NTD gold wires in the inner copper pins. e Positioning of the top pre-assembled LD. f Coupling of the pre-assembled LD with the top ZnSe copper

holder



Fig. 14 CUPID-0 detector installed on the cryogenic system, just below the 10 cm thick Roman Pb shielding

for monitoring the stability of the detectors and of the system. The remaining available channels might be employed for a future upgrade of the detector.

The signals are extracted from the detectors using NbTi-NOMEX[®] ribbon cables. The NbTi-NOMEX cables run from room temperature down to the MC, see Fig. 16, while from the MC to the detectors there are twisted 60 μ m constantan wires. All the cables are thermalized at the different temperature stages of the cryostat, and on the MC they are plugged into custom-made junction boards through Zero Insertion Force (ZIF) connectors, which connect the ribbon to the constantan wires, as shown in Figs. 16 and 17.

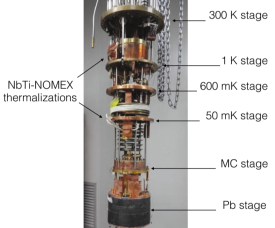


Fig. 15 Dilution unit photo. The different thermalization stages are identified by the arrows. A Roman Pb shield is hung on the MC stages by means of a vibrational damping system. The junction board is installed on the Pb stage, this is shown in Fig. 17

The NbTi-NOMEX ribbon cables are made of 13 twisted pairs of 100 μ m NbTi twisted wires. Their low radioactivity and electrical properties [37] make them the best choice for the detector read-out. They are characterized by low thermal conductivity, becoming superconducting below 10 K, low parasitic capacitance (100 pF/m), negligible cross-talk, and 500 twisting/m.

The second major upgrade of the cryogenic system consisted in a mechanical double stage anti-vibrational decoupling system, similar to the one developed in [38]. The main purpose for the development and installation of such system was driven by the fact that any microphonic noise source has to be minimized, in order to prevent any spoiling of the LD

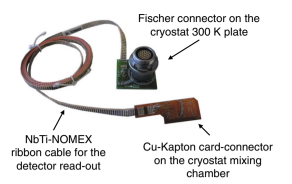


Fig. 16 NbTi-NOMEX cable from 300 K to the mixing chamber. On the room temperature side they are soldered to Fischer 27-pin connectors on the other end they are soldered to customized Cu-Kapton Zero Insertion Force connectors

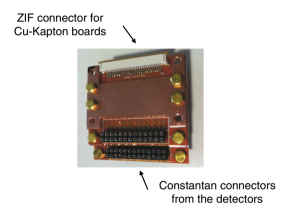


Fig. 17 Junction boards on the MC for connecting the NbTi-NOMEX cables to the constantan wires from the detectors

bolometric performance. In fact the LDs, having a higher sensitivity compared to the ZnSe crystals, requires a much lower vibrational environment compared to a system where only massive crystals are operated [39].

Figure 18 shows a scheme of the designed mechanical decoupler which is directly hanging from the MC using the top brass ring. The circular brass piece holds in place the 10 cm thick Roman Pb shield by means of three customdesigned wires made of harmonic steel (red color of Fig. 18, where just one wire connector is shown). The Roman Pb shield is connected to the harmonic steel wires by means of three harmonic steel wings mechanical anchored on the Pb (purple color of Fig. 18). The characteristic longitudinal resonance frequency of this first decoupling stage is about 12 Hz. On the top of the Pb shield is encapsulated a steel spring which is mechanically connected to the detectors by means of Cu cylinder housed inside the Pb shield. The Cu connector is mechanical decoupled from the Pb acting as second mechanical decoupling system, and it is characterized by longitudinal resonance frequency of about 5 Hz. The Cu connector acts as shielding from the radioactivity of the

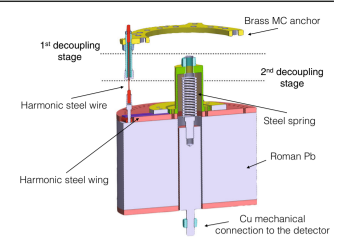


Fig. 18 Rendering of the double stage mechanical decoupling system. This is installed directly on the cryostat mixing chamber with the top brass anchor and on the bottom part there is a Cu mechanical connection for the detector installation

spring, which due to mechanical reasons is not made from high purity materials.

The thermal connection is ensured by several 100 μ m thick (99.999 % purity) Cu strips between the MC and the first damping stage and by 2 -softer- 50 μ m thick 6 cm long, 2 cm wide copper stripes between the first damping stage and the Cu detector top plate.

7 Detector readout

The readout system of CUPID-0 shares the same general structure as that of the CUORE experiment [40]. Many of the operating parameters were optimized for CUPID-0. Figure 19 shows the block diagram of the readout chain for a single detector, valid for both crystals and LDs. The thermistor R_B is biased with a DC current through a pair of load resistors $R_L = 30 \ G\Omega$ (10 GΩ). The bias generator V_L can be set between -25 and $25 \ V$ with 16-bit resolution. The voltage across the thermistor is amplified by the two-stage amplifier A_1 and A_2 . The total gain can be set between 27 and 10000 V/V with 12-bit resolution. The input stage of A_1 is based on a JFET differential pair.

We designed two different options: (a) low parallel noise, less than 100 f A below 50 °C, and about $3.5 nV/\sqrt{Hz}$ series white noise (twice this value at 1 Hz); and (b) low series noise, $1.2 nV/\sqrt{Hz}$ (twice this value at 1 Hz), with larger parallel noise. In principle, the choice depends on the value of NTD impedance. In practice we observed that in both cases this stage was not limiting the resolution, and option a) was found adequate for all detectors. The thermal parallel noise of the load resistors R_L , whose maximum value and temperature of operation are constrained by its extremely large values, tens of G Ω .

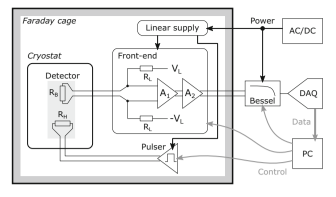


Fig. 19 Block diagram of a readout channel, from the detector to the DAQ

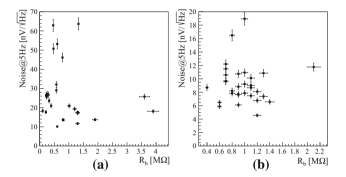


Fig. 20 Detectors RMS noise at 5 Hz as a function of the Ge-NTD dynamic impedance for ZnSe (a) and LD (b)

In Fig. 20a, b, the RMS noise at 5 Hz is shown for the ZnSe crystals and the LD, respectively. The measured noise in the present setup is larger than expected considering the value of the detector impedances, the contributions from the front-end amplifier and the load resistors. Future optimization will be focused on further reducing the observed noise, minimizing mechanical vibrations and mitigating the pre-amplifier noise, attempting to reach this limit.

The amplified signals are routed out of the Faraday cage to the antialiasing filters (Bessel–Thomson, 6 poles) and the data acquisition system (DAQ) [41]. The cutoff frequencies, settable in 4 steps, are 15, 35, 100, 120 Hz for the crystals and 15, 100, 140, 220 Hz for the LDs. The final values are set at 35 and 100 Hz for the ZnSe crystals and LDs, respectively. The DAQ hardware consisted of six National Instruments (NI) PXI-6284 digitizer boards in a NI PXI-1036 chassis. Each board digitized 16 differential channels with 18 bit resolution over a range of ± 10.5 V. The ZnSe crystals and the LDs are acquired with a sampling frequency of 1 and 2 kHz, respectively. The recorded waveform have a length of 2 and 0.5 s for the two types of bolometer.

The Pulser board is used to generate voltage pulses, which are injected onto the detector by resistor R_H [42]. The pulses are triggered and tagged by the DAQ, and used for relative calibration during data taking. Their noise is negligible, typi-

cally at the order of 10 ppm RMS, and their thermal stability is better than 1 ppm/°C, reducing the need for calibration runs with radioactive sources. Similar boards are also used to stabilize the temperature of the mixing chamber and of the detector holder through PI (proportional-integral) control loops. The power supply is provided by a two stage system: a commercial floating AC/DC generator with a custom filtering solution [43], followed by two custom linear power supplies with low noise (1.6 μV peak to peak between 0.1 and 100 Hz) and high stability (about 1 ppm/°C), which serve also as reference voltages for the front-end amplifier and the bias generator [44]. In this way the entire system is able to maintain a stability better than 10 ppm/°C. The front-end and the Pulsers are housed in 19" 6U and 19" 3U standard racks respectively. A total of 66 channels are available, which are also used to read out diagnostic thermometers. Two Pulser boards (4 channels each) are used for stabilization, and one (two channels) is used for PI control. The entire system is remotely controlled by a PC through an optically coupled CAN bus.

8 Detector configuration

CUPID-0 detector was cooled down in February 2017 and the first operation to be performed was the evaluation of the detector temperatures. This consists in measuring the resistance of the Ge-NTD sensors on all the ZnSe crystals and LDs. According to the sensor design we are expecting a base resistance (R_{base}) of hundreds of M Ω on the ZnSe and one at least order of magnitude larger values for the LD, given the reduced sensor mass. The spread in the distribution of the base temperature provides information on the uniformity of the absorber properties, namely the heat capacity of the system: ZnSe + Ge-NTD. This discrepancy is highlighted if we look at the spread of the distribution of the working resistance (R_{work) of the detectors. The R_{work} is the Ge-NTD resistance once the operational parameters of the detector are set, e.g. biassing current.

In Fig. 21a, b, the distributions for the R_{work} for the ZnSe crystals and LDs are shown. The distribution of the LD resistances tells us that the production of such devices is highly reproducible and there is a robust control of the critical aspects for the detector production. For the ZnSe, on the other hand the spread of the distribution is large and this is due to two different aspects: the first is because the crystals have different masses, hence different heat capacities, and the second because there is limited control of the ferromagnetic impurities inside the absorber. We performed a bias scan in order to evaluate the best operating condition of each ZnSe crystal and LD. This was done varying the bias current of each detector and evaluating the best signal-tonoise ratio for each configuration. In Figs. 22, 23, we show

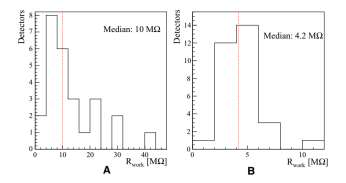


Fig. 21 Distribution of the Ge-NTD sensor resistances for ZnSe (a) and LD (b) at the operating conditions

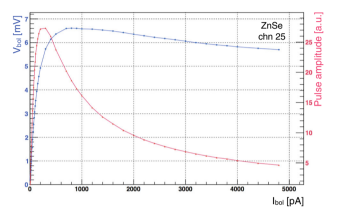


Fig. 22 Characteristic load curve of a CUPID-0 ZnSe crystal operated with a Ge-NTD thermal sensor. The figure shows how varying the detector biasing voltage acts on the signal amplitude (red) and on the voltage drop across the sensor resistance (blue)

how the signal amplitude varies as a function of the detector biasing voltage for a ZnSe detector and how the sensor is able to stand sizeable power dissipation without affecting its working resistance. In the detectors the reference signals are generated by a Si resistor coupled to the crystal operated as Joule heater.

In order to better estimate the best operating conditions of the detector, the signal-to-noise ratio is the key parameter to be optimized for each ZnSe crystal and LD. In fact, also the noise amplitude has to be taken into account, especially the parallel Johnson noise that develops across the resistors of the biasing circuit, which becomes more relevant at high values of the Ge-NTD resistances, hence lower temperatures. A compromise between low noise condition – higher temperature - and large signal amplitudes - lower temperature must be established. A reference pulse is generated on each absorber dissipating the same amount of energy through the Si resistors. While varying the biasing voltage we monitor how the amplitude of the reference signal varies and how the detector noise changes. In Fig. 24 we show the signalto-noise ratio for a set of measurements at different detector operating bias for ZnSe and LD's. The reported values are

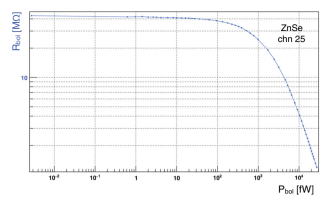


Fig. 23 Characteristic load curve of a CUPID-0 ZnSe crystal operated with a Ge-NTD thermal sensor. The figure shows how a Ge-NTD stands high power dissipation without affecting the sensor operational conditions

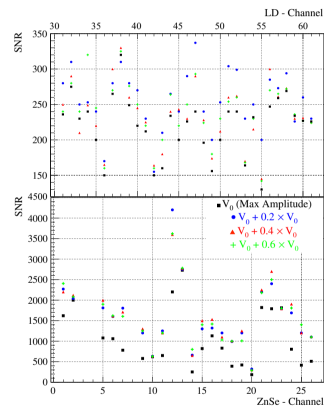


Fig. 24 Signal-to-noise ratio (SNR) scan for LDs (top) and ZnSe crystals (bottom) varying the voltage bias. It is calculated as the ratio of the filtered pulser amplitude to the $\sigma_{baseline}$. V₀(Max Amplitude) represents the biasing voltage which gives the maximal pulse amplitude. The scan is performed at bias higher than V₀(Max Amplitude) because we expect a stronger reduction of the noise compared to the signal amplitude

estimated filtering the acquired pulses by means of the Optimum Filter technique [45].

Summarizing, for each absorber, load curve measurements are performed by evaluating the configuration that maximizes the signal amplitude, see Fig. 22. Then, a nar-

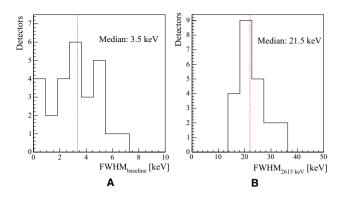


Fig. 25 Distribution of the ZnSe energy resolutions. The FHWM resolution at 0 keV, which is defined as the detector baseline noise, in shown in (a). While the distribution for the 2615 keV γ -line energy is shown in (b)

rower scan in proximity of region of operation is carried out, aiming at defining the best signal-to-noise ratio, see Fig. 24.

9 Detector performance

The overall detector performance is benchmarked by means of a ²³²Th calibration source deployed next to the detector, but outside of the cryostat. This is used to calibrate the energy response of the detector at the RoI and of the baseline energy resolution⁶. Unfortunately, we are only able to calibrate the ZnSe crystals and not the LDs, due to their low mass. The best method to calibrate such small devices would be to place a permanent X-ray source on the detector, as it was already done in [13]. In CUPID-0 we decided not to install any sort of permanent source on the LDs for obvious reason related to the ultra-low background conditions in which the measurement is carried out.

In Fig. 25a, b we show the distribution of the detector FHWM resolutions for the ZnSe detectors at 0 keV (FWHM_{baseline}) and at 2.6 MeV (FWHM₂₆₁₅), the high energy and high intensity γ -line produced by the ²³²Th source. The median value of the FWHM_{baseline} reveals to us that the cryogenic system and the electronics are performing at the cutting edge, given that the baseline noise in first approximation is independent of the absorber properties.

The average detector energy resolution is computed at 2.6 MeV, the most intense high energy gamma line next to the region of interest. The exposure-weighted harmonic mean FWHM energy resolution results to be 23.0 ± 0.6 keV. The spread in the energy resolution is driven by the limited crystal quality, in fact while an ideal bolometer is supposed to be a crystal with a single-crystalline structure, our ZnSe cystals

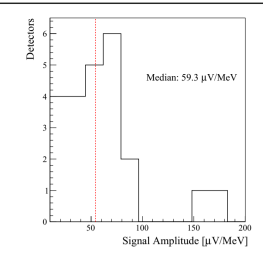


Fig. 26 Distribution of the ZnSe signal amplitudes for each crystal

have polycrystalline structures. This characteristic strongly affects the thermalization of phonons inside the crystal, hence the energy resolution. In fact, in this case the amplitude of the thermal signals will depend on the point of interaction inside the crystal, thus it will spoil the detector energy resolution, expecially for multi-Compton events occuring in one single crystal. We would like to underline the fact that there are still effective methods for improving the energy resolutions, and the most important one consists in taking advantage of the heat-light correlation in ZnSe crystals. In our group, while operating ZnSe bolometers, we were able to improve the detector energy resolution by 25% [46] by means of the heat-light de-correlation.

In Fig. 26, we show the distribution of the signal amplitudes for the Zn⁸²Se crystals. The median value of the distribution is 59.3 μ V/MeV, which is a value comparable with other large mass bolometers like ^{*nat*} TeO₂ [30].

This is the first time that a large number of bolometric LDs is operated: 31 channels. Their performance had not yet been investigated on such a large scale. Due to the lack of a calibration source for the LD, we investigate the LD performance by means of the pulses generated by Si resistor coupled to the absorber. In Fig. 27, the distribution of the signal-to-noise ratio of the heater pulses recorded on each LD is shown. The LD performance is extremely reproducible showing no major outlier. This proves the control and robustness of this technology for the scintillation light read-out.

Finally in Table 4, we show the overall operating and performance parameters of all ZnSe crystals and LDs. The risetime and the decay time are also shown. These are computed as the time interval between the 10 and 90% of the leading edge of the pulse amplitude and as the 90 and 30% of the trailing part of the pulse amplitude, respectively. Furthermore we also report the noise amplitude evaluated at 5 Hz which is within the signal bandwidth. This shows that the detector res-

⁶ The energy resolution at 0 keV is evaluated as the detector baseline resolution and it is evaluated on acquired baseline where no pulses are recorded.

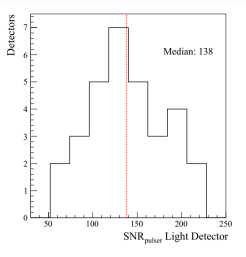


Fig. 27 Distribution of the LD SNRs. The signal amplitude is evaluated on test pulses generated by a current pulse through a Si resistor coupled to each LD. The amount of dissipated energy is the same for each detector

olution is limited not by the electronics noise (see Sect. 7), but by the detectors themselves.

We would like to underline the fact that the energy calibration of the LD it is not needed in order to perform the particle identification and rejection, because this is carried out on the relative signal amplitude. Moreover, in Table 4 given the reproducibility of the LD performance we only show the median value for the different parameters.

10 Detector radiopurity

In order to achieve the extremely low-background index in the RoI for a sensitive investigation of $0\nu\beta\beta$ decay, the detector radiopurity is fundamental. All the materials used for the detector realization were chosen for their ultra-low concentration of impurities. Nevertheless, the final detector radiopurity can be spoiled if dedicated procedures are not adopted, while producing or handling detector components. In order to validate and to prove the firm control of all the procedures adopted for the detector production, it worths to analyzing the internal contamination of all the crystals used for CUPID-0. At the same time, the study of the internal contaminations of the detector is fundamental for the development of a reliable and robust background model for the study of the possible background sources in the RoI.

Thanks to the excellent particle discrimination, we can select with high efficiency only events induced by α particle interactions, by applying a cut on the shape of the scintillation light signal, this class of data selection is thoroughly discussed in [13,46]. In Fig. 28 we show the energy spectrum of the CUPID-0 detector for α interacting particles. For details on the data selection see [48]

In the energy spectrum, the peaks between 4 and 7 MeV are induced by natural radioactive decays occurring in the crystal bulk, while the excess of events at higher energies are induced by pile-up events, usually defined as Bi-Po cascade. In the ²³⁸U and ²³²Th decay chains there are two nuclides which undergo a β decay and after a few msec the daughter nucleus undergoes an α decay. The poor time resolution of the bolometers does not allow disentanglement of these two events which then produce a single events with a total energy equal to the Q-value of the two transitions minus the energy carried away by the neutrino. These cascades are ²¹⁴Bi \rightarrow ²¹⁴Po for the ²³⁸U chain and ²¹²Bi \rightarrow ²¹²Po for the ²³²Th one.

All the other events in the energy spectrum are mostly ascribed to surface α contaminations [35].

In Table 5 we report the measured internal radioactive contaminations for the CUPID-0 enriched crystals. We also show the highest and lowest concentrations observed in each nuclide. The overall detector radiopurity complies with the needs for a extremely low background index in the RoI of ⁸²Se, which is expected to be at the level of few 10^{-3} counts/(keV kg years). The average detector purity is not better than other bolometric detectors for $0\nu\beta\beta$ [47], nevertheless if we look at the best values for each nuclide, the achieved radiopurity is competitive with the previously mentioned results.

An important observation that can be inferred from the table is the wide spread in the radiopurity level. In fact there is about one order of magnitude difference between the lowest and highest nuclide activity. All of the procedures established for the crystal production were not sufficient for keeping under complete control the impurity concentration in the crystals. Nevertheless, from preliminary studies we can state that the crystal purity has strongly improved during the crystal production; the crystals which show the highest purity are the one produced at end of the production campaign. A speculative explanation could be given by the fact that the crucibles employed for the crystal growth have not undergone a full purification process, which actually occurred during the crystal production.

11 Conclusion

CUPID-0 is the first large array of scintillating bolometers for $0\nu\beta\beta$ decay investigations. The detector is made of 26 crystals, 24 of them are enriched in ⁸²Se at the level of 95% and 2 have natural isotopic abundance. The large $0\nu\beta\beta$ source mass and the particle discrimination capability may allow CUPID-0 to achieve background levels in the ROI unprecedented for bolometers. This is feasible thanks to the excellent performance and reproducibility of the LDs which are operated together with the Zn⁸²Se bolometers.

Channel ID	Name	Tower	Type	Mass [g]	R _{base} [MΩ]	\mathbb{R}^{work} [M Ω]	Noise@5Hz $[nV/\sqrt{Hz}]$	Signal Amplitude [µV/MeV]	Rise time [ms]	Decay time [ms]	FWHM ^{baseline} [keV]	FWHM ₂₆₁₅ [keV]
1	CG-01	1	Enriched	439.40	10	4.17	11.7	40	10.1	24.7	2.59	21
2	CG-13	1	Enriched	427.86	54	14.71	25.7	81	12.0	37.6	2.58	35
3	CG-28	1	Enriched	427.00	29	9.63	18.1		9.5	17.3	I	I
4	NAT-1	1	Natural	418.39	34	11.28	26.3		5.5	20.7	I	I
5	CG-26	1	Enriched	408.22	7	3.94	20.9	51	11.1	33.1	3.22	19
6	CG-02	2	Enriched	441.29	16	4.90	19.3	46	9.3	23.6	3.86	22
7	CG-15	2	Enriched	469.64	25	11.25	17.2	47	12.6	26.2	2.88	20
8	CG-29	2	Enriched	480.90	23	6.42	17.7		8.5	13.7	I	I
6	CG-14	2	Enriched	470.59	27	9.56	26.5	66	10.3	21.5	3.57	20
10	CG-16	2	Enriched	260.52	2	1.22	10.1	13	9.2	26.4	6.09	19
11	CG-03	3	Enriched	438.65	11	4.34	13.7	23	10.3	27.4	4.84	22
12	CG-20	3	Enriched	214.62	14	6.42	21.0	179	18.1	48.3	0.96	15
13	CG-23	3	Enriched	174.89	33	9.85	25.7	156	12.0	34.2	1.47	14
14	Xtra-4	Э	Enriched	409.88	63	12.69	23.6	26	11.5	26.7	6.85	25
15	CG-18	e,	Enriched	410.31	LL	18.13	29.0	57	14.5	33.3	3.17	24
16	CG-22	Э	Enriched	418.92	65	21.26	50.7	68	15.1	47.5	3.84	18
17	CG-04	4	Enriched	442.32	150	30.78	46.1	69	15.8	40.4	4.10	25
18	Xtra-2	4	Enriched	442.43	101	23.63	62.9	73	14.4	35.7	5.05	25
19	CG-17	4	Enriched	474.22	73	20.76	32.1	36	14.0	36.6	5.11	29
20	NAT-2	4	Natural	431.21	216	41.40	63.7	38	15.9	32.9	8.41	38
21	CG-21	4	Enriched	233.08	23	10.06	27.2	61	14.6	58.1	2.40	20
22	CG-10	5	Enriched	440.47	14	6.91	17.2	68	14.4	39.3	1.94	19
23	CG-08	5	Enriched	431.00	62	28.19	53.2	84	18.1	44.0	3.36	17
24	CG-24	5	Enriched	429.62	42	14.33	26.7	62	11.9	26.1	3.06	29
25	CG-25	5	Enriched	434.51	14	5.32	18.3	35	9.7	20.5	4.84	26
26	CG-27	5	Enriched	431.18	11	5.77	13.7	21	10.6	26.8	5.36	25
ZnSe Median					28	9.95	24.6	59.3	13.5	35.7	3.47	22
I D Median						116	27		1. (ī		

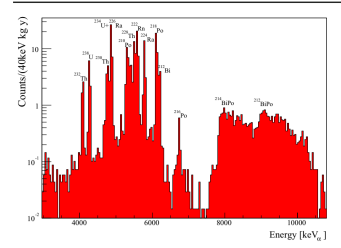


Fig. 28 Total alpha energy spectrum of all detectors

Table 5 Evaluated internal α radioactive contamination for the CUPID-0 detector. The values reported refer to the overall detector radiopurity (CUPID-0), the lowest (Best) and highest (Worst) measured contamination in a single crystal. ²¹⁰Po values refers to the sum of bulk and surface contaminations. Limits are evaluated at 90% C.L.

Chain	Nuclide	Activity [µBq/kg] CUPID-0	Activity [µBq/kg] Lowest	Activity [µBq/kg] Highest
²³² Th	²³² Th	2.5 ± 0.2	< 0.54	8.6 ± 1.2
	²²⁸ Th	13.6 ± 0.4	2.3 ± 0.8	26.9 ± 2.2
	²²⁴ Ra	10.9 ± 0.3	2.1 ± 0.6	23.1 ± 0.2
	²¹² Bi	12.2 ± 0.6	<3.7	24.2±3.5
²³⁸ U	²³⁸ U	5.1 ± 0.2	<1.2	12.7 ± 1.5
	²³⁴ U	5.3 ± 0.8	1.0 ± 2.0	14.7±4.3
	²³⁰ Th	5.3 ± 0.2	<2.4	16.4±1.7
	²²⁶ Ra	17.0 ± 0.4	3.8 ± 0.9	18.4 ± 1.8
	²¹⁸ Po	17.4 ± 0.4	3.4 ± 0.6	19.8 ± 1.9
	²¹⁰ Po	18.8 ± 0.6	9.1 ± 0.3	45.4 ± 2.6

The large number of detectors, 31 LDs and 26 Zn⁸²Se crystals, required an upgrade of the cryogenic infrastructure. The cryostat is now able to host up to 136 channels, thus also a possible second phase of the CUPID program. Furthermore an innovative and effective vibration damping system was installed which allowed CUPID-0 to achieve relevant results on the detector performance, in terms of baseline energy resolution.

The state of the art technologies in low background techniques and in cryogenic detector design have been implemented inside the CUPID-0 detector. The preliminary results obtained with this innovative detector are a significant step towards the demonstration of the feasibility of a next generation cryogenic bolometric experiment for the investigation of $0\nu\beta\beta$ down to the meV scale. Acknowledgements This work was partially supported by the European Research Council (FP7/2007-2013) under contract LUCIFER no. 247115. We are particularly grateful to M. Iannone for the help in all the stages of the detector construction, A. Pelosi for construction of the assembly line, M. Guetti for the assistance in the cryogenic operations, M. Lindozzi for the development of cryostat monitoring system and the mechanical workshop of LNGS (in particular E. Tatananni, A. Rotilio, A. Corsi, and B. Romualdi) for continuous and constructive help in the overall set-up design. We acknowledge the Dark Side collaboration for the use of the low-radon clean-room.

Open Access This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. Funded by SCOAP³.

References

- L. Gonzalez-Mestre, D. Perret-Gallix, Nucl. Instr. Meth. A 279, 382 (1989)
- J.N. Ullom, D.A. Bennett, Supercond. Sci. Technol. 28, 084003 (2015)
- S. Pirro, P. Mauskopf, Annu. Rev. Nucl. Part. Sci. 67, 161–181 (2017)
- C. Alduino et al., [CUORE Collaboration], Adv. High Energy Phys. 2015, 879871 (2015)
- C. Alduino et al., [CUORE Collaboration], Eur. Phys. J. C 77, 532 (2017)
- C. Alduino et al., [CUORE Collaboration], Eur. Phys. J. C 77, 543 (2017)
- 7. CUPID Interest Group, arXiv:1504.03599 (2015)
- 8. CUPID Interest Group, arXiv:1504.03612 (2015)
- 9. D.R. Artusa et al., Eur. Phys. J. C 74, 3096 (2014)
- 10. J.W. Beeman et al., Adv. High Energy Phys. 2013, 237973 (2013)
- 11. J.W. Beeman et al., Eur. Phys. J. C 72, 2142 (2012)
- 12. E. Armengaud et al., Eur. Phys. J. C 77, 785 (2017)
- 13. D.R. Artusa, Eur. Phys. J. C 76, 364 (2016)
- 14. E.E. Haller, *Neutron transmutation doping of semiconductor materials* (Springer, Boston, 1984)
- 15. M. Berglund, M.E. Wieser, Pure Appl. Chem. 83, 397-410 (2011)
- 16. J.W. Beeman et al., Eur. Phys. J. C 75, 591 (2015)
- 17. G. Zuzel et al., J. Phys. Conf. Ser. 798, 012109 (2017)
- 18. I. Dafinei et al., J. Cryst. Growth **475**, 158–170 (2017)
- 19. S. Pirro et al., Nucl. Instr. Meth. A **559**, 361–363 (2006)
- 20. J.W. Beeman et al., J. Instr. 8, P07021 (2013)
- 21. D.E. Aspnes et al., Phys. Rev. B 27, 985-1009 (1983)
- 22. I. Dafinei et al., IEEE Trans. Nucl. Sci. 57, 1470-1474 (2010)
- 23. M. Mancuso et al., EPJ Web Conf. 65, 04003 (2014)
- 24. A. Langenkämper et al., e-Print: arXiv:1703.07152
- 25. S. Nisi et al., Int. J. Mod. Phys. A 32, 1743003 (2017)
- 26. A. Borio di Tigliole et al., Progr. Nucl. Energy 52, 494-502 (2010)
- 27. A. Borio di Tigliole et al., Progr. Nucl. Energy 70, 249-255 (2014)
- 28. E. Andreotti et al., Astropart. Phys. 34, 822–831 (2011)
- 29. http://www.huntsman.com/advanced_materials/a/Home
- 30. CUORE-0 Coll., J. of Instrum. **11**, P07009 (2016)
- 31. D.R. Artusa et al., Adv. High Energy Phys. 2015, 879871 (2015)
- C. Rusconi, "Optimization of the bolometric performances of the CUORE-0/CUORE and LUCIFER detectors for the neutrinoless double beta decay search", Ph.D. Thesis, Università Degli Studi dell'Insubria, Italy (2011)
- 33. E. Andreotti et al., Nucl. Instr. Meth. A 664, 161 (2012)

- 34. F. Alessandria et al., Astropart. Phys. 45, 13-22 (2013)
- 35. M. Clemenza et al., Eur. Phys. J. C 71, 1805 (2011)
- 36. K. Alfonso et al., Phys. Rev. Lett. **115**, 102502 (2016)
- 37. E. Andreotti et al., J. Instr. **4**, P09003 (2009)
- 38. S. Pirro, Nucl. Instr. Meth. A5(59), 672–674 (2006)
- 39. C. Ligi et al., J. Low Temp. Phys. **184**, 590 (2016)
- 40. C. Arnaboldi et al., arXiv:1710.06365 (2017), submitted to JINST (to be updated)
- 41. C. Arnaboldi at al., Nucl. Instr. Meth. Phys. Res. A 617, 327 (2010)
- 42. P. Carniti et al., arXiv:1710.05565 (2017), submitted to JINST (to be updated)
- 43. C. Arnaboldi et al., Rev. Sci. Instr. 86, 124703 (2015)
- 44. P. Carniti et al., Rev. Sci. Instr. 87, 054706 (2016)
- 45. E. Gatti et al., Rivista del Nuovo Cimento 9, 1 (1986)
- 46. J.W. Beeman et al., J. Instrum. 8, P05021 (2013)
- 47. C. Arnaboldi et al., J. Cryst. Growth 312, 2999–3008 (2010)
- 48. CUPID0 Coll., O Azzolini, e-Print: arXiv:1802.07791