GARFIELD + RCo digital upgrade: A modern set-up for mass and charge identification of heavy-ion reaction products

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Abstract. An upgraded GARFIELD + Ring Counter (RCo) apparatus is presented with improved performances as far as electronics and detectors are concerned. On the one hand fast sampling digital read out has been extended to all detectors, allowing for an important simplification of the signal processing chain together with an enriched extracted information. On the other hand a relevant improvement has been made in the forward part of the set-up (RCo): an increased granularity of the CsI(Tl) crystals and a higher homogeneity in the silicon detector resistivity. The renewed performances of the GARFIELD + RCo array make it suitable for nuclear reaction measurements both with stable and with Radioactive Ion Beams (RIB), like the ones planned for the SPES facility, where the physics of isospin can be studied.

1 Introduction

The international nuclear physics community is more and more interested in the development of new facilities based on accelerated radioactive ion beams [1]. For many years several laboratories worldwide have been active in this field. As far as the European ones are concerned, one can mention the following: the FAIR project [2] at GSI in Germany, Spiral II [3] at GANIL in France, HIE-ISOLDE [4] at CERN, FRIBS [5] at INFN Laboratori del Sud (LNS) in Catania and SPES [7] at INFN Laboratori Nazionali di Legnaro (LNL) in Italy. Together with the construction of these facilities, and in view of the forthcoming EURISOL project [8], new instrumentation has to be developed and the existing detectors have to be upgraded. In this framework, focusing on the charged particle detectors and their use at moderate energy in RIB facilities, it is important to achieve good isotopic separation in a wide range of fragment masses together with low-energy identification thresholds.

Several apparatuses exist in different laboratories with this aim. For example, CHIMERA [9–11] at LNS is a multi-detector based on 1192 silicon - CsI(Tl) telescopes exploiting Time of Flight (ToF), ΔE-E and scintillator fast-slow technique to detect and identify charged particles. At GANIL, the INDRA apparatus [12] features a Ionization Chamber (IC) stage in front of silicon detector and CsI(Tl) scintillators to reduce the energy threshold for charge identification. It detects the products emitted in nuclear reactions both with stable and unstable (Spiral) beams. Both these apparatuses cover the whole solid angle and presently rely mainly on analog electronics. They obtain mass and charge identification for light...
fragments (Z up to 3–4) through fast-slow Pulse Shape Analysis (PSA) in CsI(Tl) while silicon PSA has been obtained, in CHIMERA, via specific modules allowing the double CFD method. Apparatuses equipped with digital electronics exist in the U.S., like NIMROD [13] in Texas A&M University Cyclotron and HIRA [14] at the NSCL of MSU, which exploit several available particle identification techniques including PSA in silicon detectors [15–18]. Finally one has to mention the FAZIA demonstrator [19]; this array of 192 three-stage Si-Si-CsI modules is currently under construction. It presents the best performances reachable with this kind of detectors, thanks to specific detector solutions and a new designed digital electronics. FAZIA is a versatile system able to be coupled with other large arrays.

In order to meet the more demanding performances in experiments with the future RIB facility at LNL, the GARFIELD + Ring Counter (RCo) [20–22] charged particle and fragment detection array has been upgraded in order to obtain both light charged particle (LCP, \( Z \leq 2 \)) isotopic separation in all covered range (5°–150°) and a better position determination (granularity) at the forward angles where the kinematics concentrates most of the reaction cross section. Moreover, particular care has been devoted to increase the fragment (Z ≥ 3) identification capabilities. This upgrade allows for a more complete event reconstruction and therefore a better characterization of the different competing reaction mechanisms.

The GARFIELD array consists of two independent gas drift chambers designed especially to identify intermediate mass fragments and light charged particles. They exploit different techniques based on micro-strip gas chambers [23] and CsI(Tl) scintillators [24,25]. The forward angular range is covered by the RCo, a detector based on a three-layer telescope configuration: Ionization Chamber (IC), Silicon Strip Detectors and CsI(Tl) scintillators. In fig. 1 a sketch of the complete GARFIELD + RCo apparatus is shown. In this configuration a geometrical efficiency of ∼ 80% is achieved.

According to the needs of different experiments, different combinations of GARFIELD, RCo and ancillary detectors have been used. In particular a time-of-flight system, composed by large-area position-sensitive parallel plate avalanche counters (PSPPAC), has been used instead of the RCo to detect evaporation residues and/or fission fragments in fusion/fission nuclear reactions. This allowed to perform experiments aimed at a detailed study of light particle pre-equilibrium emission [26]. Further experiments have been performed by using a set of phoswich detectors made of three different scintillation layers (two plastics and one CsI(Tl) crystal) read by a unique photomultiplier, formerly used in the FIASCO experiment [27]. For this kind of experiments a system of BaF₂ scintillators (HECTOR apparatus [28,29]) was used in place of the backward drift chamber: this coupling was considered in order to detect high-energy gammas in coincidence with evaporation residues and with light charged particles. With this set-up we have studied the Giant Dipole Resonance [30], the isospin mixing [31], the Jacobi shape transition [32] and the dynamical dipole emission [33]. Recently the first version of the GARFIELD + RCo set up has been used to study odd-even effects in the isotopic fragment distribution (staggering) up to the multi-fragmentation threshold [34,35]. The upgraded set-up has been recently exploited in low-energy light nuclei reactions to get information on the statistical and non-statistical behavior of the light particle emission mechanism [36]. In the next future, the FAZIA [19] telescopes could be coupled to GARFIELD, to perform nuclear dynamics and thermodynamics studies in the perspective of the RIB of SPES.

In sects. 2 and 3 the main features of the set-up will be recalled and the major upgrades, which are hereafter listed, will be described in detail:

1) design and implementation of new digitizing Front End Electronics (FEE) for all detector channels [37], along the same lines as of what has been done for new generation apparatuses, such as, for instance, HIRA [14] for charged particles or AGATA for γ-rays [38];
Fig. 2. (Color online) Picture of the micro-strip pads mounted in one of the GARFIELD chambers. Cabling and pre-amplifiers are visible in the upper part of the figure.

2) replacement of the old RCo Silicon detectors with neutron-transmutation-doped ones (uTD) [39];
3) increase of the granularity of the RCo CsI(Tl) scintillators.

2 GARFIELD

The GARFIELD apparatus consists of two large-volume gas detectors employing micro-strip gas chambers (MSGC) as $\Delta E$ stage, complemented by CsI(Tl) scintillators as residual energy detectors. This allows to achieve low identification thresholds (0.8–1 MeV/\(u\)) using the $\Delta E$-E technique. The two drift chambers have cylindrical symmetry and are placed back to back with respect to the target. The first chamber (C1 in fig. 1) covers the angular region from $\theta = 95^\circ$ to $\theta = 150^\circ$ and has a side gap of $\Delta \phi \approx 45^\circ$ which allows the placement of different ancillary detectors. The second one covers the angular region from $\theta = 30^\circ$ to $\theta = 85^\circ$ and the complete azimuthal angle. The two gas volumes are, respectively, divided in 21 and 24 azimuthal sectors and are filled with CF$_4$ at a typical pressure of 50 mbar. Each sector is equipped with one trapezoidal micro-strip pad [23] divided in 4 different collecting zones providing the $\Delta E$ signals (see sect. 2.1). The ensemble of all 21 (24) trapezoidal pads forms a “ring” around the target (see fig. 2). The pads are positioned almost perpendicularly ($82^\circ$) to the beam axis (see fig. 3). A total of 84 (96) micro-strip signals are read out in the first and the second chamber, respectively.

As shown in fig. 3, each sector is completed by four CsI(Tl) crystals with photodiode readout which provide the residual energy signals ($E$). These detectors also define the boundaries of the sensitive gas volume and are positioned in a radial configuration with respect to the interaction point (see sect. 2.2).

In each sector one motherboard carrying 8 pre-amplifiers is mounted inside the gas volume, close to the detectors in order to minimize their input capacitance. The pre-amplifiers have been designed to have a large dynamic range (up to 10 V) and low noise, less than 130 ENC (Equivalent Noise Charge), together with low power consumption (about 200 mW each) [40]. Their nominal gain is 45 mV/MeV silicon equivalent. A water recirculating cooling system is used to dissipate the pre-amplifiers generated heat.

The original version of the array was equipped with an analog electronic chain mainly composed by the CAEN N568B shaping amplifiers, SILENA FAIR 9418/6-V32 peak sensing ADCs, SILENA FAIR 9418/6-T32 TDCs and C208 CAEN Camac CFDs. ADCs and TDCs were part of a VME acquisition system, based on FAIR protocol [41]. Also VME delays based on the FAIR protocol were part of the electronic chain. The complete analog chain has been replaced by new digitizing Front End Electronics (FEE) developed ad hoc by the collaboration, also compatible with the FAIR framework.

The core of the new FEE is a single channel board based on a fast sampling ADC followed by a Digital Signal Processor (DSP). The main features of the 125 MHz - 12 bit digitizing cards are described in [37]. The DSP is used to perform on-line pulse shape analysis [42] and extract the relevant parameters from the sampled signal: energy, timing and pulse shape information are obtained using the algorithms described in [37, 43, 44] and later summarized (see sect. 2.3).

2.1 Gas micro-strip detectors

Each GARFIELD micro-strip pad has a keystone geometry and it is divided in four collecting zones (1A, 1B, 2A, 2B, as sketched in fig. 4). The pads are composed of very small anode/cathode alternated metallic electrodes (dis-
tance = 50 μm) that are deposited through photo-lithography technique on glass. The anodes are 10 μm large, electrically connected into the four collecting areas and biased at 430 V. The cathodes have a trapezoidal shape (width from 85 μm to 140 μm in the 1A and 1B zones and from 85 μm to 190 μm in the 2A and 2B ones) and are all grounded.

The MSGC operation is based on the avalanche multiplication process in gas, which is used to amplify the signals due to primary ionization electrons. A proper electric field forces the free electrons formed along the track of the impinging particle to drift toward the anode. The anodic plane is screened towards the drift region by a Frisch grid positioned 3 mm apart: in this way the primary signals due to primary ionization electrons. A proper multiplication process in gas, which is used to amplify the electrons start feeling the very strong MSGC electric field (∼ 10^7 V/m) only after crossing the grid, thus producing the electron avalanche. Using CF4 gas at 50 mbar, typical gain factors of 30–50 are achieved; this is one of the main advantages of using MSGC with respect to normal ionization chambers. CF4 gas is used due to its high density (0.19 mg/cm³ at a pressure of 50 mbar and at a temperature 20°C) and high electron drift velocity (10 cm/μs at 1 V/cm/mbar). An automatic recirculating and filtering system makes the gas continuously flowing in the chamber to minimize the oxygen contamination.

The obtained signals carry the information on the energy loss (∆E) and on the polar angle (θ) of the impinging ion [23,45,46]. Every MSGC collecting zone allows for an angular resolution of ∆φ = 7.5° and ∆θ = 1°–3° [20, 21], the latter exploiting the measured drift time of the electrons. The energy resolution of the chambers is about 6% [20,21] for elastically scattered 32S at 200 MeV leaving about 10 MeV in the gas.

2.2 CsI(Tl) scintillators

As shown in fig. 3 each GARFIELD sector contains 4 CsI(Tl) scintillators, the shape of which has been designed in order to optimize the geometrical efficiency. A total of 96 (84) crystals are present in the forward (backward) chamber. Each detector covers ∆φ = 15° and ∆θ ≃ 15° (θ_foward) = 35°, 47°, 60°, 75°; (θ_back) = 105°, 120°, 133°, 145°. The crystal thickness is 4 cm in order to stop the charged products expected in the energy range of interest. The thallium doping is about 1200 ppm and contributes, together with the chosen diffusive wrapping, to maximize the light collection [20,21,47]. The CsI(Tl) crystals are optically coupled to Hamamatsu S2042-05 photodiodes closely connected to the pre-amplifiers. The energy resolution is about 3% for 5 MeV α particles and 2–3% for 8 MeV/u Li and C elastically scattered beams on Au target.

2.3 Digital FEE algorithms

As already stated, the major upgrade of the GARFIELD array is related to the digital readout electronic chain [37]. Each pre-amplifier output signal is now directly fed into a fast sampling ADC and online processed by a DSP. The algorithm implemented on the signal processor extracts time, energy and particle identification information from the sampled signal. According to the different detector type specific calculations are performed.

The energy information is derived from the maximum amplitude of a semi-Gaussian shaper with a selectable value of peaking time (usually 4–5 μs). This shaper has been designed to faithfully reproduce the response of the analog shaping amplifier previously employed. The shaper is an Infinite Impulse Response filter, implemented using a recursive algorithm. In order to reduce truncation error propagation on our 16 bit DSP, the algorithm exploits double precision calculations (32 bit) and to reduce calculation time the input signal is decimated 16 fold before filtering, thus increasing its sampling period from 8 to 128 ns. Since the ballistic deficit of the digital version, as a function of the signal rise time, is the same of the analog shaper, the calibration points obtained in the past using various ion energy combinations can still be used.

Since all the digitizing channels are triggered by the global trigger of the experiment, a zero suppression step is needed to reduce the data flow to the acquisition. This is done channel by channel inside the DSPs, where the data output is validated only if the shaper maximum amplitude is above a selected threshold. The global trigger is composed by the sum of simple (singles) or composite (coincidences) trigger signals: generally a downscaled single logic signal (i.e. the reduced logic OR of all the GARFIELD CsI(Tl) detectors and/or the logic OR of all the RCo Si detectors) plus more structured ones are used. The more complex triggers are related to all possible coincidences among different parts of the apparatus, which can be from time to time needed depending on the specific experiment (i.e. OR Garfield and OR RCo).

Timing information from the sampled signals are obtained using a digital CFD algorithm based on cubic interpolation [48]. This procedure calculates a digital CFD (dCFD) time mark corresponding to a specific percentage (typically 25%) of the signal maximum amplitude. This amplitude is evaluated by the DSP as the average of the absolute maximum and the following 15 samples of the digitized signal in order to reduce the influence of electronic noise fluctuations. The 8 ns digitizer sampling period is known within 1/10⁸ and it is stable within a few units in 10⁵ so that the conversion from samples
Fig. 5. (Color online) GARFIELD microstrip signals and associated (coincident) CsI(Tl) signal. Bottom: CsI 5–8 signals acquired in coincidence with microstrip signals. Top: Signals of a GARFIELD microstrip pad obtained in coincidence with CsI 5–8, respectively (from left to right). As expected the delay of the microstrip signal with respect to the one of the associated CsI(Tl) increases as a function of the polar angle with respect to the beam (larger drift time).

to nanoseconds is straightforward and, therefore, no time calibration is needed. Since the on-board DSP features fixed-point arithmetics and a relatively time-consuming division algorithm, the cubic interpolation, needed to extract the time mark, is best performed off-line. The DSP just calculates the dCFD threshold and it identifies the four samples needed for interpolation (two before and two after the threshold crossing) which are then sent to the data acquisition. This allows to reduce the processing dead time and to achieve a better precision in the time mark computation. The time mark is in generally used to avoid contamination of different beam bunches. In the following, the specific algorithms used for the different detectors will be described.

2.3.1 Drift chamber signal analysis

For the MSGC detectors both energy and drift time information are needed: the energy is evaluated as previously described, while the electron drift time in the gas, which corresponds to the polar angle information, is obtained as the time difference between two dCFD time marks. They are calculated, respectively, on the CsI(Tl) and on the microstrip signals.

Figure 5 shows signal shapes from ions hitting a given sector. The tracks produce CsI(Tl)-microstrip coincidences, represented with four colors, one for each of the four polar regions corresponding to the CsI(Tl) crystals (from 5 to 8). Each event is triggered by one of the CsI(Tl), thus the CsI(Tl) waveforms (bottom panel) start at about the same time mark (there is an almost fixed delay between the global trigger and the CsI(Tl) trigger).

However, signal pairs have been realigned in time in such a way that the 12.5% threshold of the CsI(Tl) falls exactly at 3500 ns. Since CsI(Tl) scintillators are placed at different polar angles, the electron drift times associated to the different tracks (microstrip signal) change and in particular grow from CsI5 to CsI8 (and from CsI4 to CsI1, respectively, for the C1 chamber), as one can observe in the top panel in fig. 5.

Also the signal rise time is sensitive to the track polar angle, as shown in fig. 6, where the signals recorded from the four collecting zones have been time aligned for a better comparison. The rise time is evaluated as the difference between two dCFD time marks calculated on the same signal, one at 12.5% and the other at 62.5% of the maximum amplitude. The dependence of the rise time on the polar angle is somewhat milder than the one of the drift time as it will be shown in sect. 2.4.2.

2.3.2 Algorithms for the CsI(Tl)

Time, energy and particle identification information are all needed by the CsI(Tl) detectors. The time and energy information are obtained as described in the general section. Further information is related to the particle identification procedure: to identify particles stopped in the CsI(Tl) crystal, the well-known technique of shaping the charge signal with two different time constants is used [25]. Pulse shape identification of light charged particles is obtained correlating the amplitudes of two semi-Gaussian shapers with a peaking time of 2.5 $\mu$s and 5 $\mu$s, respectively (see sect. 2.4.3).

Before the upgrade, pulse shape identification in CsI(Tl) detectors was not available, since it would have required twice the number of shaper amplifiers and ADC’s. Timing information was available via standard CFD and TCD modules, which were removed after the upgrade. The use of digital electronics has thus extended the capabilities of the GARFIELD apparatus also allowing to simplify the front end electronics.
2.4 Upgraded GARFIELD performances

2.4.1 $\Delta E$-$E$ identification

The microstrip electrodes and the CsI(Tl) scintillators give a total number of 360 three stage telescopes. The $\Delta E$-$E$ correlation, where $\Delta E$ is the energy lost in one or both sections of the drift chamber and $E$ is the residual energy released by the incoming particle in the CsI(Tl), is shown as an example in fig. 7 for the reaction $^{16}$O+$^{65}$Cu at 16 MeV/u beam energy. LCP and fragments shown in the figure are produced with charge from $Z = 1$ to $Z \approx 8$. For $Z = 2$ the maximum energy is around 110 MeV, whereas for $Z = 6$ is around 140 MeV. Using a linearization algorithm a Particle IDentification (PID) parameter can be obtained.

The charge (mass) ion identification procedure adopted so far [25,49–52] is a two-step process:

1) For each employed detector, in a bi-dimensional scatter plot ($\Delta E$-$E$ or fast-slow) several points are sampled on the ridges of well-defined isotopes.

2) The set of points for a given ridge are fitted one by one using an analytical function (if it exists) [25,49,50] or via polynomial functions (spline) [51,52].

Event by event, isotopes are identified in charge (mass), by minimizing the distance of the measured signals with respect to the identification function. This distance provides information on the dispersion around the most probable value, which corresponds to the charge (mass) value.

As an example, the PID histogram obtained from the correlation of a MSGC with the CsI7 in sector 13 is presented in fig. 7 (inset). Fragment charges are well identified when the FoM is of the order of 0.4–0.5 with neighboring charges. The charge (or the mass) can be still expected to well-separated Gaussians with a peak-to-valley ratio $\approx 2$ and less than $\approx 5\%$ of contamination by other isotopes.

The charge identification thresholds are around 0.8–1 $MeV/e$, as can be seen in fig. 7 (inset). Fragment charges are well identified when the FoM is of the order of 0.4–0.5 with neighboring charges. The charge (or the mass) can be still expected to well-separated Gaussians with a peak-to-valley ratio $\approx 2$ and less than $\approx 5\%$ of contamination by other isotopes.

As a set of fig. 7, one gets values of the FoM ranging from $\approx 1$ to $\approx 2$.

2.4.2 Drift and rise time of drift chamber signals

As previously explained, the polar angle information can be extracted using the electron drift time. Light charged particles give small signals, due to the low energy loss in the gas. If the signals are comparable to the noise of the electronic chain, it is more difficult to obtain a definite angular information. The fast amplification stadium of the N568 amplifiers, together with the C208 CFD threshold procedure determination, made it possible to obtain sufficiently precise drift times only for fragments with $Z \geq 3$ in the analog case. The same result was easily obtained in the digital electronics case, as shown in fig. 8, where the data for the drift time of fragments with $Z \geq 3$ emitted in the reaction $^{40}$Ca+$^{16}$O at 16 MeV/u beam energy are displayed: the events are sorted as a function of each CsI(Tl) scintillator in the sector hit by the impinging particles and the well-separated contributions in the spectrum are represented with different colors. The shortest times correspond to CsI5 in fig. 3, whereas the largest to CsI8. Moreover, with digital electronics, a clean drift time information can also be extracted for $\alpha$-particles, due to a better signal-to-noise ratio of the digitizing electronics.

In fig. 9 (left panel) $\alpha$-particle data are shown, which have been collected by one of the outer microstrips for one GARFIELD sector and selected through the $\Delta E$-$E$ identification technique. The energy of the $\alpha$-particles ranges from 8 to 130 MeV.

In the right panels of figs. 8 and 9 the correlation between rise time and drift time is presented. The two quantities are clearly correlated, even though the drift time shows up a major sensitivity to the polar angle, as expected. From the finite width of the borders of each CsI(Tl) drift time spectrum in figs. 8 and 9 we can estimate the angular resolution. It is found to be in the 1–3 degree range in agreement with previous tests [20,21].
of procedure described in ref. [25] are shown. The energy range
fig. 10, the mass distributions obtained following the pro-
and d-t, respectively. For

\[ Z \]

attributed to

performed for a precise energy calibration of the CsI(Tl)
calibration phase [25]. Dedicated measurements have been
studied to quickly identify the isotopes and to speed up the

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Fig. 9. (Color online) Left: Drift times in coincidence with CsI
5 to 8 of one sector. Data refer to fragments (Z \geq 3). Different colors in the time spectra correspond to the
four CsI(Tl) of fig. 3. Right: Rise time vs. drift time correlation.
Loci associated to different CsI(Tl) scintillators are well
separated.

Obtaining the angular information also for low ionizing
alpha particles is one of the advantages of using the
digital electronics in GARFIELD.

2.4.3 Pulse shape analysis of CsI(Tl) signals

It is well known that CsI(Tl) allows for particle identification
through pulse shape analysis. The fast-slow technique
was applied for GARFIELD according to the procedure described in sect. 2.3.2 [25]. In fig. 10 (top panel)
fast-slow bi-dimensional plots for one of the CsI(Tl) de-
tectors are shown: the ΔE-E correlation. Loci associated to different CsI(Tl) scintillators are well
separated. In the bottom panel of fig. 10, the mass distributions obtained following the procedure described in ref. [25] are shown. The energy range of α-particles ranges from 8 to 130 MeV.

The FoM obtained for Z = 1 are 0.98 and 0.96 for p-d
and d-t, respectively. For Z = 2 one gets FoM = 1.29 for
\[ ^3He - \alpha \] discrimination. An automatic procedure has been
studied to quickly identify the isotopes and to speed up the
calibration phase [25]. Dedicated measurements have been
performed for a precise energy calibration of the CsI(Tl)
crystals: in this way the light output response function with respect to the energy and charge of the reaction products has been determined.

3 Ring counter

The RCo is a three-stage annular detector array with a
truncated cone shape covering the \( \theta = 5^\circ - 17^\circ \) angular range. The first stage is an ionization chamber (IC), fol-
lowed by silicon strip detectors (Si) and CsI(Tl) scintil-
lators. The ion chamber is divided into eight azimuthal sectors (each one covering \( \Delta \theta = 45^\circ \)) sharing an unique

gas volume. Eight separate pie-shaped silicon detectors are
mounted on an aluminum holder behind every IC sector:
each of them is segmented into eight independent annular
strips. The third stage of the RCo has been renewed in order to increase the granularity: the two old CsI(Tl) crys-
tals positioned after each silicon pad have been replaced
by six smaller-size 4.5 cm thick CsI(Tl) crystals (for a total number of 48 devices read by photodiodes). The geomet-
rical shapes of the new CsI(Tl) crystals precisely fit the
angular cone subtended by the IC.

A remotely controlled collimator system has been im-
plemented in order to apply different screens in front of the
RCo under vacuum and therefore while running the ex-
periment. Different screens are used for focusing, shielding or
collimation purposes. The RCo is mounted on a sliding
plate and can be remotely moved back and forth in order
to replace the front collimator. The RCo operational
position is inside the forward cone of the GARFIELD drift
chamber at 177 mm from the target as shown in fig. 1. The pre-amplifiers are mounted on the same sliding plate of the RCo close to the detectors, in order to reduce their input capacitance. They are thermally connected to shielded metallic boxes to be refrigerated through a water cooling system.

3.1 The ionization chamber

The IC is 6 cm long with three thin electrodes (1.5 μm aluminum Mylar): an intermediate anode and two grounded entrance and exit cathodes. At present the detector is the same of the first version [22] but novel signal processing is performed using the same digital electronics described for GARFIELD. At the operating pressure of ~50 mbar, a resolution of about 6% is obtained for the energy lost by elastically scattered 17 MeV $^32$S ions (their average energy loss in IC being ≈6 MeV).

3.2 The silicon detectors

The performance of the silicon detectors has been substantially improved. On the one hand, the new digitizing electronics has added pulse shape identification capabilities to the apparatus (e.g., by measuring both the amplitude and the rise time of each pre-amplifier signal) [15–18]. On the other hand, the study performed within the FAZIA framework [19] demonstrated the better performances achievable using silicon detectors with good resistivity uniformity (less than few percent) [54]. For this reason new nTD silicon detectors [39] (which are known to feature a high controlled doping homogeneity) have been mounted. The major advantage of a better homogeneity is the reduction of the pulse shape dependence on the particle interaction zone in the detector [55]. Furthermore, at present the silicon detectors have been reverse mounted: the pulse shape discrimination capability is, in fact, enhanced by forcing the particle to enter through the low field side [52,56]. Each silicon pad covers one sector and the rear electrode (junction side) is segmented into eight strips. The polar angle interval covered by each strip is quoted in table 1. Values in the table refer to silicon detectors of the same of the first version [22] but novel signal processing is performed using the same digital electronics described for GARFIELD. At the operating pressure of ~50 mbar, a resolution of about 6% is obtained for the energy lost by elastically scattered 17 MeV $^32$S ions (their average energy loss in IC being ≈6 MeV).

### Table 1. Radii and polar strip angles.

| Strip | Int. radius | Ext. radius | Min. angle | Max angle |
|-------|-------------|-------------|------------|-----------|
| 1     | 77.9        | 85.0        | 15.6       | 17.0      |
| 2     | 70.8        | 77.8        | 14.2       | 15.6      |
| 3     | 63.7        | 70.7        | 12.9       | 14.2      |
| 4     | 56.6        | 63.6        | 11.5       | 12.8      |
| 5     | 49.4        | 56.4        | 10.1       | 11.4      |
| 6     | 42.3        | 49.3        | 8.6        | 10.0      |
| 7     | 35.2        | 42.2        | 7.2        | 8.6       |
| 8     | 26.2        | 35.1        | 5.4        | 7.2       |

Fig. 11. Left: scheme of the strips of the silicon detector. The dotted lines correspond approximately to the different CsI(Tl). Right: CsI(Tl) scintillator limits for one of the eight sectors of the RCo.

In this way charge split and cross-talk are drastically reduced. The bulk resistivity of the detector is of the order of 3400 Ω·cm, the full depletion voltage is 100 V with recommended bias of 120 V. The reverse current of each strip is about 30 nA. The silicon detectors feature relatively thin window thicknesses (50 nm junction side, 350 nm ohmic side) in order to get low thresholds even in the reverse mounting configuration. The thickness of the silicon detectors is around 300 μm, and therefore the punching through energies are about 6 MeV/u for protons and α-particles and 7–11 MeV/u for light fragments. A resolution of the order of 0.3% both for 5 MeV α particles and for elastically scattered 17 MeV/u $^32$S (about 500 MeV residual energy) has been obtained. The preamplifiers used for the silicon strips have a gain of 5 mV/MeV and are fed into the same digital read-out cards used for the other channels.

3.3 The CsI(Tl) scintillators

The granularity of RCo has been increased bringing the number of CsI(Tl) scintillators from 16 to 48. For each sector the 4 Si inner strip polar angular region is now azimuthally divided in two parts: in place of the previous single crystal ($\Delta \phi = 45^\circ$), two CsI(Tl) scintillators are located ($\Delta \phi = 22.5^\circ$ each). Moreover the 4 outer strip region is now covered by four CsI(Tl) crystals, each covering...
half of the polar and half of the azimuthal angular range defined by the strips. The smaller dimension of the CsI(Tl) allows for better doping (and therefore scintillation) uniformity and better light collection of the single detector (better photodiode area to crystal dimension matching). Having scintillators of similar size also favors response uniformity among the various detectors. The right panel of fig. 11 shows a front cross section of the position of the six CsI(Tl) of each sector. A picture of the upgraded RCo during its mounting is shown in fig. 12. In the region where the silicon detectors were not yet mounted the upgraded CsI(Tl) configuration can be seen.

As a consequence of studies on the resolution and Light Output performances of CsI(Tl) scintillators as a function of the doping content and homogeneity [24, 57, 58], the present CsI(Tl) crystals have been chosen with a Tl doping in the range of 1500–2000 ppm. The 48 crystals, 45 mm thick, have been wrapped with micro-porous polyvinylidene fluoride membrane [59] and Teflon tape. They have been optically coupled with Hamamatsu photodiodes (S2744-08, 10 × 20 mm²), 500 μm thick, dark current (≤ 15 nA).

The preamplifiers have the same characteristics as the ones described for GARFIELD.

The algorithms employed in the analysis of silicon detector signals are the same as those applied for GARFIELD micro-strips (see sect. 2.3.1). Energy, time of occurrence and rise time information are extracted. The main parameters, in this case, are the amplitude (i.e. energy) and rise time information: their correlation allows for fragment identification via pulse shape analysis.

The algorithms employed in the analysis of the CsI(Tl) scintillators are the same used for the GARFIELD scintillators (see sect. 2.3.2).

3.5 Performances of the detectors of the RCo

Two complementary methods can be used to identify the reaction products.

1) The ΔE-E technique. Considering the RCo three fold stage one can extract information on:
   - Particles stopped in the silicon detector: from the energy lost in the ionization chamber vs. the residual energy in a silicon strip.
   - Particles stopped in the CsI(Tl) scintillators: from the energy lost in a silicon strip vs. the residual energy in the scintillator.

2) The pulse shape analysis which gives information on:
   - Particles stopped in the silicon detector through energy-rise time correlations.
   - Particles stopped in CsI(Tl) through fast-slow correlations.

3.5.1 ΔE-E from IC-Si

A typical ΔE-E scatter plot for a IC sector signal (ΔE) and a silicon strip signal (E) correlation is shown in fig. 13 (left panel) for the fragments produced in the reaction $^{32}$S + $^{40}$Ca at 17 MeV/u. Charged particles and fragments are clearly visible, together with low-energy heavy residues formed through fusion reactions.

The different nuclear species can be identified in charge and their kinetic energy can be obtained, after proper calibration. Especially for the low energy region of the heavier fragment, the “in gas” energy loss corrections, estimated according to refs. [60, 61] have to be considered. The high quality of the detectors and electronics can be demonstrated by the wide range of charge identified ions: as shown in fig. 13, the ridges are rather separated up to scandium, corresponding to the evaporation residue region and down to 0.8–1.0 MeV/u in energy. In the right panel of fig. 13 the PID charge identification function is shown. The FoM for increasing charge of the identified fragments to the beam charge varies from 0.8 to 0.5, whereas in the residue region (Z around 20) the FoM becomes of the order of 0.4.
optimized for greater is slightly shifted from the right value since the procedure is optimized for greater $Z$ values.

Fig. 13. (Color online) Left: $\Delta E$-$E$ correlation “IC vs. silicon” for the reaction $^{32}$S + $^{48}$Ca at 17 MeV/u at $\theta \approx 8^\circ$. The line which extends to the largest silicon energy corresponds to $^{32}$S ions. Right: PID for the same data. The $Z = 2$ distribution is slightly shifted from the right value since the procedure is optimized for greater $Z$ values.

Fig. 14. (Color online) $\Delta E$-$E$ correlations obtained for a representative pair of a silicon strip and a CsI(Tl). The left panel shows the resolution for $Z = 1$–$3$, whereas the right panel focuses on ions $Z > 3$. The two well-separated isotopic lines of $^7$Be and $^8$Be are easily recognized in the right panel, starting at a $\Delta E$ energy of $\approx 50$ MeV. The maximum energy for $\alpha$-particles is around 140 MeV, for carbon 270 MeV and for neon 430 MeV.

3.5.2 $\Delta E$-$E$ plots from Si-CsI(Tl)

Thanks to the excellent energy resolution of the silicon detectors, the Si-CsI(Tl) $\Delta E$-$E$ correlation allows to identify the reaction products both in $Z$ and $A$. In particular it is evident from fig. 14 that the mass separation is certainly achievable up to $Z = 11$–$12$. Even though the available statistics in the studied reaction does not allow to determine the upper $Z$-value of mass separation, a quantitative evaluation of the quality of the detector resolution can be inferred from the FoM.

In fig. 15 the results of mass identification using the procedure described in ref. [49] is shown. The obtained FoM ranges from 1.1 at $Z = 1$ down to 0.77 at $Z = 10$, showing that $Z = 10$ is still not at the upper limit for good mass separation.

3.5.3 PSA in silicon detectors

As already mentioned, the reverse mounting of the silicon detectors allows for a better exploitation of PSA. Moreover, the use of the fast digitizing electronics allows for extracting several parameters from the signal waveform, a few of them well suited for PSA.

In fig. 16 the energy vs. rise time signals have been plotted [15–18,52]. Data refer to the results of a typical detector strip. A good charge separation has been achieved for all the products. The lower energy threshold for charge identification is $\approx 2.5$–5 MeV/u, which corresponds approximately to the energy needed by the fragment to travel along the first 30 $\mu$m of the detector (see refs. [55,56]).

Zooming the energy-rise time plot it appears that some marginal mass resolution is achievable in some regions of the correlation plot (see fig. 16, bottom panel). Efforts are in progress in order to improve the resolution in view of future experiments. In this respect the use of preamplifiers of higher sensitivity is promising [55]. In fact, the ion separation capability of silicon via PSA strongly depends on the rise time determination and this latter is influenced by the noise especially for small signal amplitudes. Increasing the preamplifier sensitivity gives better signal-to-noise ratio, taking into account the sampling ADC internally generated noise. As a consequence, a better precision in rise time measurements can be achieved. Figure 17 shows the quality of the separation for the same RCo detector when using a larger sensitivity (45 mV/MeV@Si): the improved behavior clearly appears, even if some saturation effect is present.

3.5.4 Pulse shape analysis in CsI(Tl) detectors

As for the GARFIELD scintillators the PSA method (see sect. 2.4.3) is applied to the RCo CsI(Tl) signals. Typical results are shown in fig. 18 (top panel). In the insets a zoom for $Z = 1$ and $Z = 3,4$ shows the achieved mass identification. The $\alpha$-particle energy ranges from 25 to 130 MeV.
In the lower panels of fig. 18 the mass identification obtained for $Z = 1$ up to $Z = 4$ in the whole energy range is shown (for $Z = 4$ only the isotope $^7\text{Be}$ is well separated and identified). The calculated FoM for masses of $Z = 1$ and $Z = 2$ particles is larger than 1.5 and for $Z = 3$ it is 0.82.
are higher for the PSA with respect to the $\Delta E-E$; on the contrary the FoM obtained by the PSA are better than those obtained by $\Delta E-E$ due to the resolution of the gas $\Delta E$. In addition, if two particles/fragments enter the same IC sector, but in different silicon strips, only the PSA can discriminate between them. The obtained results with PSA will make it possible to perform specific experiments at very low IC pressure, or even to remove it, depending on the charge-energy distributions of the ions to be detected.

iii) Mass identification of fragments (up to at least $Z = 11$) from the Si-CsI(Tl) $\Delta E-E$ correlation and pulse shape - rise time correlation.

iv) Mass identification of light isotopes (up to $Z = 3$) exploiting the pulse shape analysis of CsI(Tl) crystals, with thresholds of the order of $\geq 6 \text{ MeV}/u$, due to the preceding $300 \mu \text{m}$ silicon detector.

### 4.3 Results of the coupled apparatuses

In order to show which kind of physics the coupled apparatuses are able to perform, we point to the high quality of the charged particle identification and of the energy determination. Indeed if one calculates the correlation functions between couples of identified isotopes one can infer the contribution of excited levels decaying to a given pair of isotopes. In a recent paper, an experiment performed with GARFIELD and RCo apparatuses in their first version [34,35] has demonstrated that the position of the levels of excited parent nuclei can be well recognized by correlation functions. The further improved granularity and particle identification obtained with the present version of the apparatuses will allow to perform this analysis with a reduced distortion and smaller width of the correlation peaks.

The overall quality of the detector response (namely mass, energy and position determination) is tested by an example of $Q$-value estimation from a 3-body event reconstruction [61]. Indeed in a recent experiment [36] performed with GARFIELD + RCo, the $Q$-values of a specific reaction, i.e. $^{12}\text{C} + ^{12}\text{C}$ going to $^{16}\text{O} + \alpha + \alpha$, show values compatible with the calculated ones from the mass difference, as it is clearly shown in fig. 19, where one can see three different peaks. The rightmost peak corresponds to the reaction $^{12}\text{C}^{(12}\text{C},\alpha\alpha)^{16}\text{O}$, leaving the $^{16}\text{O}$ residue in its ground state ($Q = -0.11 \text{ MeV}$). The central peak corresponds to the reaction $^{12}\text{C}^{(12}\text{C},\alpha\alpha)^{16}\text{O}^*$, with the $^{16}\text{O}$ residue in one of its excited bound states at $E^* = 6.05, 6.13, 6.92$ and $7.12 \text{ MeV}$, which are so close in energy that cannot be resolved in the experiment and the measured average energy expense is $Q \approx -6.55 \text{ MeV}$. (We note that this estimation largely bases on data from the CsI detectors and therefore cannot be compared with specific high-resolution spectrometers). The leftmost arrow corresponds to the opening ($Q = -15.78$) of the reaction $^{12}\text{C}^{(12}\text{C},\alpha\alpha)^{15}\text{O}$, where neutrons, not detected in this experiment, have a broad energy distribution.

### 5 Conclusions

The improved GARFIELD + RCo apparatus has been described. The array, designed to study nuclear reactions in the $5-20 \text{ MeV}/u$ energy range at LNL, will be employed also in future experiments with the radioactive SPES beams. The main properties of the set-up are the large acceptance, the good energy resolution, the low energy thresholds and the high granularity especially in the forward part (RCo) which allow for correlation function studies. Moreover, the GARFIELD+RCo complex is the first European 4$\pi$ charged particle detector array fully provided with digital electronics. The new digitizing boards allowed for a sensible reduction of the needed front-end channels, keeping or even improving the performances obtained with the analog electronics.

Different techniques are used to identify particles, such as $\Delta E-E$ correlations and pulse shape analysis in scintillators and silicon detectors: these complementary techniques allow for extensive charge and, for the lightest products, mass identification in a large energy and angular range.

This improved apparatus well matches the needs of advanced studies on nuclear reactions, especially where the complete event reconstruction is required [34–36]. For example the studies of pre-equilibrium LCP emission [26] can now be extended to $Z = 1$ and $Z = 2$ isotopes, allowing deeper comparisons with models. A better reconstruction of warm fragments [35] via correlation functions will be possible with the extended mass identification and the increased granularity.

Fig. 19. (Color online) $Q$-values of reactions $^{12}\text{C}^{(12}\text{C},\alpha\alpha)^{16}\text{O}$. The arrows correspond to $\alpha$-decay chains, starting from the $^{24}\text{Mg}^*$ compound nucleus and leaving an $^{16}\text{O}$ residue either in its ground state or in one of its excited bound states at $E^* = 6.05, 6.13, 6.92$ and $7.12 \text{ MeV}$. The leftmost arrow corresponds to the upper limit ($Q = -15.78$) of the opening of the 4-body channel going to $^{12}\text{C}^{(12}\text{C},\alpha\alpha)^{15}\text{O}$. For more details see text.

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