Combining in-beam $\gamma$-ray and conversion electron spectroscopy

The SAGE spectrometer

Thesis submitted in accordance with the requirements of the University of Liverpool for the degree of Doctor in Philosophy

by

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To my family and friends;
to all those who stood by me in this endeavour.

And to Prof. Spyros Dedousis
who left us forever on Easter day this year.

Αιώνια η μνήμη.
Σα βγεις στον πηγαμό για την Ιθάκη,
να σύχεσαι νάναι μακρώς ο δρόμος,
γεμάτος περιπέτειες, γεμάτος γνώσεις.

'Ιθάκη'
Κωνσταντίνος Καβάφης
Greek poet 1863-1933

A physicist is just an atom’s way of looking at itself

*Niels Bohr*
Danish physicist 1885-1962
Abstract

In-beam $\gamma$-ray and electron spectrometers are some of the most valuable tools in the study of excited states of nuclei; the latter particularly in the case of very heavy nuclei. However, if used separately they can provide only partial information of the nuclear deexcitation processes and consequently of nuclear structure. This becomes increasingly problematic in heavy nuclei, especially at low transition energies and high multipolarities, where internal conversion competes strongly with $\gamma$-ray emission.

To provide the means for a more complete spectroscopy in the superheavy nuclei region the Silicon And GErmanium (SAGE) spectrometer was designed and commissioned by a collaboration from the Universities of Liverpool in the UK and Jyväskylä in Finland and the STFC Daresbury Laboratory.

SAGE combines the JUROGAM II germanium-detector array with a highly-segmented silicon detector and allows efficient cross-coincidence measurements between $\gamma$ rays and conversion electrons. It uses digital front-end electronics and is coupled with the RITU gas-filled recoil separator and the GREAT focal-plane spectrometer for recoil-decay tagging studies.

Simulations and tests were performed to optimise the spectral response and electron transmission efficiency of SAGE. The spectrometer employs magnetic fields, to transport electrons to the silicon detector, electric fields to reduce the low energy background and a silicon detector coupled with high-gain preamplifiers to measure the electron energies. The electron part of the spectrometer is designed so it does not affect the performance of JUROGAM II.

In this thesis the individual parts of the spectrometer are presented and analysed. The different design criteria and limitations are described in such a way that the reasoning behind every component of SAGE is made clear.
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Abbreviations

AHO ........... Anisotropic harmonic oscillator
BGO ........... Bismuth Germanium Oxide
CFD ............ Constant Fraction Discriminator
CMOS ........... Complementary Metal Oxide Semiconductor
CPCI CPU ...... Compact Peripheral Component Interconnect Central Processing Unit
DSSD ........... Double-sided Silicon Strip detector
EM .............. Electromagnetic
FET ............. Field-effect transistor
FPGA ........... Field-programmable gate array
FWHM ........... Full Width at Half Maximum
GO-box .......... Gain and Offset box
GREAT ........... Gamma Recoil Electron Alpha Tagging focal plane spectrometer
GRT4 ........... Gamma-Ray Tracking 4 channel card
ICC ............. Internal Conversion Coefficient
ICE ............. Internal Conversion Electron
LDM ............ Liquid Drop Model
LGO-card ....... Liverpool Gain and Offset card
MCA ............. Multi Channel Analyser
MHO ............. Modified harmonic oscillator
mic-mac ........ Microscopic-macroscopic method
MWPC ........... Multiwire Proportional Counter
<table>
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<th>Abbreviation</th>
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<tr>
<td>PCB</td>
<td>Printed Circuit Board</td>
</tr>
<tr>
<td>PIN</td>
<td>P-type - Intrinsic - N-type semiconductor</td>
</tr>
<tr>
<td>RITU</td>
<td>Recoil Ion Transport Unit</td>
</tr>
<tr>
<td>SAGE</td>
<td>Silicon And GERmanium array</td>
</tr>
<tr>
<td>SGO-card</td>
<td>SmartPET Gain and Offset card</td>
</tr>
<tr>
<td>spec-amp</td>
<td>Spectroscopy amplifier</td>
</tr>
<tr>
<td>TDR</td>
<td>Total Data Readout</td>
</tr>
<tr>
<td>TDRi card</td>
<td>Total Data Readout interface card</td>
</tr>
<tr>
<td>TNT2</td>
<td>Tracking Nuclear Treatment card</td>
</tr>
<tr>
<td>VHS-ADC</td>
<td>Very High Speed Analog-to-Digital converter</td>
</tr>
<tr>
<td>Vpp</td>
<td>Peak-to-peak voltage</td>
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Chapter 1

Introduction

The main motivation of constructing the SAGE spectrometer is the investigation of nuclear properties at the extremes of atomic mass and atomic number, where internal conversion becomes as important as γ-ray emission. SAGE will also allow studies of E0 transitions which proceed via internal conversion only [Pa09a].

One very interesting, yet unanswered question regarding the superheavy nuclei region is the location of the next spherical proton shell closure above Z=82. Some models, like the liquid drop model (see Subsection 2.1.1), predict that nuclei with Z>104 cannot exist as they should fission instantly [Kr00]. According to theoretical predictions where shell structure is considered, a superheavy doubly magic nucleus will have enhanced stability [Ni68, Ni69, Ćw05]. This should lead to the formation of the so-called “island of stability”, the precise limits of which are yet unknown [Og00, St06].

The different theoretical approaches give varying estimates on the magic proton and neutron numbers in the superheavy nuclei region. Microscopic-macroscopic models (see Subsection 2.1.3), using various parameterisations of the potential (Yukawa plus-exponential with the Strutinsky shell correction, Woods-Saxon with the Strutinsky shell correction etc), predict the next doubly magic nucleus to have Z=114 and N=184 [Me67, Ni68, Ni69, Mo69, Pa89, Pa91, Mő92, Mő94, Ćw96, Sm97, Ch97, Pa04, Pa05].
Calculations using self-consistent nuclear structure models mainly disagree with the microscopic-macroscopic predictions. Only one potential parameterisation (SkI4) gives agreeable results [Be99] between the two, however, it does so at the cost of an unphysically large spin-orbit interaction. Comparisons of the mean-field calculations [Kr00, Be01] indicate that most of the non-relativistic mean-field models [Cw96, Ru97, Be99] give Z=124, 126 and N=184 whereas relativistic ones favour Z=120 and N=172 [La96, Ri96, Ru97, Be99, Be03, Af03]. For reviews on theoretical results see e.g. [Be03, Af03, Bu04, De06, Re06, Se06, So07] and references therein. A review of experimental studies relevant to these calculations is made in [He08].

The heaviest nucleus produced experimentally so far is 294118 in JINR [Og06]. Only three decay chains of this isotope were observed in the experiment over a period of 45 days. The extremely low production cross-sections of 1 pb or less for superheavy nuclei (such as element 118) [Ar00] make their in-beam spectroscopic study impossible with present day setups.

However the single-particle orbitals determining the stability of these elements can be probed through in-beam spectroscopic studies of lighter nuclei. Theoretical predictions indicate that the single-particle Nilsson orbitals near the Fermi surface in nuclei in the vicinity of the deformed nobelium, originate from spherical single-particle levels above the possible shell gap at Z=114 [He06]. A Nilsson diagram showing how the same orbitals are involved in the creation of the Z=114 and Z=102 shell gaps is presented in Figure 1.1.

254No and its neighbouring nuclei are found to be ideal for spectroscopic studies as they have reasonable cross-sections of the order of 2 µb [He06]. As well as the information obtained from in-beam and focal plane experiments in this region, the masses of three nobelium isotopes were established from direct mass measurements [Bl10] using the Penning-trap mass spectrometer SHIPTRAP [Bl07]. These measurements allow calculation of the binding energy.

In-beam studies, where γ rays and electrons were studied independently, have led to important results on the rotational structure, deformation and stability against
fission of the even-even nuclei \(^{254}\)No \([\text{He02, He06}]\) and \(^{250}\)Fm \([\text{Ba06, Gr08, Ro09}]\). \(\gamma\)-ray experiments have also been performed on \(^{252}\)No \([\text{He01}]\) \(^{246}\)Fm and \(^{248}\)Fm.

Odd-mass nuclei provide a more sensitive probe into the single-particle structure of superheavy nuclei. These nuclei pose a great experimental challenge when studied in independent conversion electron or \(\gamma\)-ray experiments. Such experiments were performed in the past for \(^{253}\)No (\(\gamma\)-ray spectroscopy \([\text{Re04, Re05, Ee06, Pa03a, Mo07, He09}]\) and conversion electron spectroscopy \([\text{He02, Pa03a, He09}]\), for \(^{255}\)Lr \([\text{Ke09}]\) and for \(^{251}\)Md \([\text{Ch07}]\).

In the spectroscopic studies of \(^{253}\)No a level scheme of two sequences of intra-band E2 transitions with M1 inter-band transitions was established. The highly converted M1 transitions were not observed in the first \(\gamma\)-ray experiments \([\text{Re04, Re05}]\) but were observed in a conversion electron study \([\text{He02, He09}]\). In the higher statistics \(\gamma\)-ray experiments that followed, M1 transitions with higher transition energy were observed but the low energy transitions still remained undetected. An in-beam experiment that would simultaneously study both conversion electrons and \(\gamma\) rays would yield a more complete level scheme.

In the case of \(^{251}\)Md the odd proton ground state is predicted to occupy the \(\pi[521]1/2^-\) orbital stemming directly from the spherical \(2f_{5/2}\) subshell, with excited states derived from the spherical \(1i_{13/2}, 1h_{9/2}\) and \(2f_{7/2}\) orbitals. The spin-orbit splitting between the \(2f_{5/2}\) and \(2f_{7/2}\) orbitals is critical for the creation of a possible spherical shell closure at \(Z=114\) (see Figure 1.1).

Establishing the relative position of the \(2f_{5/2}\) and \(2f_{7/2}\) orbitals provides information as to whether the theoretical prediction of a closed spherical shell at \(Z=114\) is true and helps to locate the “island of stability”. This can be done either through studies of single-particle excitations in odd-mass nuclei described above or multiparticle-multihole excitations in even-even nuclei, where the latter are considered to populate states involving the \(2f_{5/2}\) orbital.

The configurations of the experimentally observed bands can be deduced from the \(B(M1)/B(E2)\) reduced transition probabilities ratios. These are sensitive to the
g-factors of the orbitals involved and according to the geometric model are [Bo75]:

\[
\frac{B(M1)}{B(E2)} \propto \frac{K^2(g_K - g_R)^2}{Q_o^2},
\]

where \(g_R\) and \(g_K\) are the gyromagnetic ratios of the rotating core and the single-particle respectively (see Section 2.3), \(K\) is the projection of the total angular momentum on the nuclear symmetry axis (see Section 2.2) and \(Q_o\) is the intrinsic electrical quadrupole moment. \(B(M1)/B(E2)\) are easily measured if the electrons and \(\gamma\) rays are observed in the same experiment to provide accurate branching ratios.

Another region of interest for the SAGE spectrometer can be found in the light lead and mercury region. Here the deformed multiparticle-multihole structures intrude down in energy close to the spherical ground state, when approaching the neutron mid-shell at \(N=104\). A triplet of low-lying \(0^+\) states associated with different macroscopic shapes is observed. The properties of these low-lying \(0^+\) states and the inter-band transitions between the same spin-parity states of rotational oblate and prolate bands can be investigated through simultaneous conversion electron and \(\gamma\)-ray spectroscopy [An00, Pa07].

Nuclei in this region have been studied in the past in \(\alpha\)-decay fine structure measurements, in-beam \(\gamma\)-ray studies and in some cases internal conversion electron (ICE) measurements. Simultaneous \(\gamma\)-ray and ICE experiments performed with SAGE will measure the conversion electron strengths of the inter-band transitions (oblate to prolate) and the location and feeding of the low-lying \(0^+\) states. The measurements of the \(E0\) matrix elements in neutron mid-shell isotopes of lead and mercury will provide direct information on configuration mixing and shape changes in these nuclei.
Figure 1.1: Nilsson level diagram for protons with $Z \geq 82$ ($\epsilon_4 = \epsilon_2^2/6$), taken from [Fi96]. Calculations are made within the cranked Nilsson-Strutinsky framework as described in [Be85].
Chapter 2

Theoretical framework

2.1 Nuclear models

2.1.1 The liquid drop model (LDM)

One of the first models to successfully describe nuclear properties was the LDM, introduced in [Bo37]. Observations that the nuclear forces saturate and the nucleus has low compressibility and a well defined surface, led to the treatment of the nucleus as an incompressible nuclear fluid with the nucleons held together by the strong interaction [He94].

The LDM parametrises the energy using a volume term, a surface term and a Coulomb interaction term, based on the concept of a charged nuclear fluid. Including an isospin and a pairing term derived from the shell model, this leads to the Bethe-Weizsäcker mass equation [Kr88]:

$$BE(A, Z) = \alpha_v A - \alpha_s A^{2/3} - \alpha_c Z(Z - 1)A^{-1/3} - \alpha_i (A - 2Z)^2 A^{-1} + \delta.$$  \hspace{1cm} (2.1)

The coefficients $\alpha_v$, $\alpha_s$, $\alpha_c$ and $\alpha_i$ refer to the volume, surface, Coulomb and isospin terms respectively. $\delta$ is the pairing energy term and it is $+\alpha_p A^{-3/4}$ for even-even nuclei, $-\alpha_p A^{-3/4}$ for odd-odd nuclei and zero for odd $A$ nuclei. The LDM does not make any predictions on the values of the coefficients included in the mass formula but they are obtained by fitting Equation 2.1 to experimental data.
The LDM successfully predicts fission (not for superheavy nuclei), fusion and $\alpha$ decay. Because in the LDM the coefficients are fitted phenomenologically, the model describes well the average trends of nuclear binding energies. For the same reason though, the predictions made using the LDM have large uncertainties when extrapolating to exotic nuclei [Be03]. Additionally, because the model does not take into account shell structure, it fails to replicate many observables like the magic numbers, the discontinuities in neutron binding energies etc.

One of the major predicaments of the LDM is its inability to calculate fission barriers for the heaviest elements. According to this model elements with $Z>104$ should fission instantly due to their large electric charge [Kr00]. In these nuclei stability arises solely due to shell effects. In order to do any realistic calculations in the superheavy nuclei region, shell structure must be taken into account.

### 2.1.2 The spherical shell model

The shell model [Ma49] considers the nucleons to move in almost unperturbed single-particle orbits, due to the Pauli exclusion principle, within the uniform average potential created by the nucleons themselves. Different approaches can be taken when describing the nuclear potential such as the square well, Gaussian well, exponential well, Yukawa well, harmonic oscillator and Wood-Saxon (see Figure 2.1). In addition to the nuclear potential a spin-orbit term is introduced to the Hamiltonian in order to express the interaction between the intrinsic spin and the orbital momentum and also an $\hat{l}^2$ term, where $l$ is the orbital angular momentum, in order to flatten the effective radial shape of the potential.

The shell model reproduces the experimentally observed shell gabs, the so called magic numbers, at proton and neutron numbers 8, 20, 28, 50 and 82. For neutrons, 126 is a magic number as well. It also explains spin and magnetic moments of nuclei close to the magic numbers. On the other hand it does not explain some of the bulk properties that are successfully described by the LDM.
2.1.3 The Strutinsky shell correction

By combining the macroscopic LDM with microscopic shell corrections the much more precise microscopic-macroscopic (mic-mac) method is produced. Considering that a nuclear property, such as the total ground-state energy, consists of a smooth and an oscillatory part as shown in Figure 2.2, then in order to calculate it correctly both these parts need to be considered. In the case of the mic-mac method the Strutinsky shell correction [St67, St68] is used to extract the oscillatory part of a nuclear property and then combine it with the smooth part derived using a macroscopic model (e.g. LDM).

Using the Strutinsky procedure the energy is written as:

\[ E = E_{LDM} + E_{SHELL}, \quad (2.2) \]

where

\[ E_{SHELL} = \sum_{i=1}^{A} \epsilon_1(\delta) - \tilde{E}_{SHELL}, \quad (2.3) \]

with \( \tilde{E}_{SHELL} \) used to subtract the part of the energy that was already included in \( E_{LDM} \). \( \epsilon_1(\delta) \) denotes the eigenvalues of the Nilsson potential and \( \delta \) is a deformation.
parameter [He94].

Figure 2.2: Plot of a nuclear property showing the smooth and oscillatory parts. Figure adapted from [Pa10].

One important outcome of the mic-mac method, relevant to this work, is the prediction of enhanced stability against fission of superheavy elements. This is shown in Figure 2.3 where fission barriers for uranium, californium and hassium were calculated with and without taking shell structure into account. According to the LDM calculations hassium should be completely unbound and have a half-life of about $10^{-19}$ s. On the other hand when shell-model corrections are applied the fission barrier increases dramatically and the calculated half-life of the element increases by several orders of magnitude, even up to $10^{15}$ s for $^{292}$Hs.

2.1.4 The deformed shell model

Nuclei in the vicinity of closed shells are near spherical and can be described very well by a spherical potential. However when moving away from close shells sphericity is no longer valid and strong evidence, such as the existence of rotational bands and large intrinsic quadrupole moments, show that nuclei are deformed in the regions between
acting nuclei decreases the Coulomb
pulsion between them at the moment
they touch, and thus increases the prob-
Despite these obvious advantages, all
tries to synthesize new elements
using calcium-48 ions between 1977
and 1985 failed. However, improved ex-
perimental techniques and the availab-
ility of intense beams of calcium ions
have increased the sensitivity of these
xperiments by at least three orders
of magnitude. This has allowed us, to-
gether with colleagues at the Lawrence
Livermore National Laboratory in the
US, to probe deep into the superheavy-
If the theory is right, elements on
the islands of stability should not decay
via spontaneous fission. Instead they
should undergo alpha decay. As a result,

\[ H_{\text{def}} = -\frac{\hbar^2}{2m} \nabla^2 + \frac{1}{2} m \left( \omega_x^2 x^2 + \omega_y^2 y^2 + \omega_z^2 z^2 \right), \tag{2.4} \]
where \( \omega_i \) defines the frequency of the potential at the \( i^{th} \) direction. For axially sym-
metric nuclei \( \omega_x = \omega_y = \omega_\perp \), with \( \omega_\perp \) being the frequency perpendicular to the
symmetry axis \( z \) [He94].

The eigenstates of the AHO are conventionally labelled using the asymptotic Nils-
son quantum numbers:

\[ [N n_z \Lambda] \Omega^\pi, \]

where \( N \) is the total number of oscillator quanta, \( n_z \) the number of oscillator quanta along the symmetry axis \( z \), \( \Lambda \) is the projection of the orbital angular momentum \( l \) onto the symmetry axis and takes values \( \Lambda = N, N - 2, \ldots, 1 \) or 0, \( \Sigma \) is the projection of spin \( s = \pm \frac{1}{2} \) on the symmetry axis, \( \Omega = \Lambda \pm \Sigma \) is the projection of the total angular momentum \( j \) onto the symmetry axis (see Figure 2.4) and \( \pi \) is the parity of the level.

Figure 2.4: Schematic representation of the coupling of orbital angular momentum and spin in a deformed nucleus.

Nilsson [Ni55] modified the AHO potential to include a spin-orbit term to allow derivation of a more realistic single-particle spectrum. He also added an \( \hat{l}^2 \) term to flatten the radial shape of the potential. This potential is called Nilsson potential or modified harmonic oscillator (MHO) potential. The Nilsson Hamiltonian is:

\[
H_{\text{Nilsson}} = \hbar \omega_0(\delta) \left[ -\frac{1}{2} \Delta' + \frac{r^2}{2} - \beta r^2 Y_{20}(\hat{r}') \right] - k \hbar \omega_0 \left[ 2 \hat{l} \cdot \hat{s} + \mu \left( \hat{l}^2 - \langle \hat{l}^2 \rangle_N \right) \right],
\]

where \( \omega_0(\delta) \) arises when one considers a deformation dependent oscillator length with
the deformation parameter $\delta$, $\omega_0(\delta) = \omega_0 \left(1 + \frac{2}{3} \delta^2\right)$, $\beta = \frac{1}{3}\sqrt{16\pi/5\delta}$, $2k$ is the spin-orbit strength and $k\mu$ the $\hat{l}^2$ orbit energy shift [He94].

Other potential parametrisations, such as the deformed Woods-Saxon potential can be used instead of the MHO to calculate properties of deformed nuclei. Different potentials might work better in different areas of the nuclear chart but in general the potential to be used is a matter of personal preference.

As mentioned above for the case of superheavy elements shell effects lead to increased stability. Especially in the case of the superheavy nuclei that are far from the spherical closed shells stability arises solely from deformation. An example of this would be the deformed shell gap at the region of nobelium ($Z=102$) [He06].

To illustrate the effect of deformation on the magic numbers a single-particle level spectrum plotted as a function of quadrupole deformation, $\epsilon_2$, is shown in Figure 2.5. To keep the figure as simple as possible the axially deformed harmonic oscillator potential without the use of a spin-orbit and an $\hat{l}^2$ term was used in the calculations. Because these terms were omitted, the model does not reproduce the correct magic numbers but is still good enough in order to illustrate the effect of deformation on them. The quadrupole deformation is defined as $\epsilon_2 = (\omega_\perp - \omega_z)/\omega_0$, with $\omega_0 = \frac{1}{3}(2\omega_\perp + \omega_z)$.

From Figure 2.5 it becomes obvious that with increasing deformation single-particle levels shift forming shells gaps at different proton or neutron numbers than in the case of the spherical nucleus. This explains the presence of the deformed proton shell at $Z=102$ which should not exist if the nucleus was spherical. In more realistic calculations the produced single-particle spectra are more complicated but can reproduce the observed shell gaps accurately. An example of this is the calculations made within the cranked Nilsson-Strutinsky framework presented in Figure 1.1. In this figure the deformed shell gap in the vicinity of nobelium can be clearly seen.
Figure 2.5: Single-particle level spectrum calculated using the axially deformed harmonic oscillator potential without the use of a spin-orbit and an $\hat{l}^2$ term. The arrows indicate deformations corresponding to the ratios of $\omega_\perp : \omega_z = 1 : 2, 2 : 3, 1 : 1, 3 : 2, 2 : 1$ and $3 : 1$. The figure is adapted from [Wo92].
2.2 Nuclear rotation

In the case of spherical nuclei it is impossible to distinguish their orientation, due to axial symmetry, so rotation is forbidden. Deformation though introduces an anisotropy allowing the definition of orientation and making nuclear rotation possible.

The nucleus can be described as a core surrounded by the valence nucleons with the axis of rotation perpendicular to the symmetry axis. The total angular momentum, $I$, of the rotating nucleus is defined as the sum of the collective angular momentum, $R$, generated by the collective rotation of the nucleus and the intrinsic angular momentum, $J$, created by the motion of valence nucleons,

$$I = R + J.$$ (2.7)

Figure 2.6 shows how the collective angular momentum couples to the total intrinsic angular momentum produced by two valence single particles with intrinsic angular momenta $j_1$ and $j_2$.

![Figure 2.6: Coupling of the collective and single-particle angular momenta. $\omega$ is the rotational frequency around the rotation axis $x$ and $z$ is the symmetry axis.](image)

The signature quantum number, $r$, is defined in the case of nuclear rotation.
Signature relates with the $\mathcal{R}$ invariance of the system which arises from the invariance of the intrinsic Hamiltonian to rotations of 180° about any axis perpendicular to the symmetry axis [Bo75].

Rotational bands can be separated into three categories:

- **$K = 0$ bands**

  In this case the allowed values of signature are:
  \[
  r = (-1)^I, \tag{2.8}
  \]

  which gives the selection rules:
  \[
  I = 0, 2, 4, \ldots \text{ for } K = 0 \text{ and } r = +1, \tag{2.9}
  \]
  \[
  I = 1, 3, 5, \ldots \text{ for } K = 0 \text{ and } r = -1. \tag{2.10}
  \]

  This leads to two rotational bands with opposite values of signature, called signature partner bands. The levels within each band are separated by $2\hbar$ and their energy is given by:
  \[
  E_{K=0} = \frac{\hbar^2}{2\mathcal{J}^{(0)}} I(I + 1), \tag{2.11}
  \]
  where $\mathcal{J}^{(0)}$ is the rigid body static moment of inertia.

- **$K \neq 0$ bands**

  In the case of $K \neq 0$ signature is defined as:
  \[
  r = (-1)^{I+K}, \tag{2.12}
  \]

  and changes sign within a band for alternate spin values.

  For $K \neq 0$ bands particle-rotor coupling occurs which is described by the rotation of the nuclear core (rotor) coupled to the single particles in a way analogous to the classical Coriolis force.
In the case of large deformation or strong coupling, Deformation Aligned (DAL) scheme, \( K \) is a good quantum number and there is little signature splitting. In this case the rotational band is defined by:

\[
I = K, K + 1, K + 2, \ldots
\]

with energies:

\[
E_{\text{DAL}} = \frac{\hbar^2}{2\mathcal{J}(0)} \left[ I(I + 1) - K^2 \right].
\]

When the nucleus is weakly deformed or the particle rotation is fast, the coupling is weak and \( K \) is no longer a good quantum number. This case is called Rotation Aligned (RAL) and has the nucleonic angular momentum \( j \) as a good quantum number. The band members have spins defined by:

\[
I = j, j + 2, j + 4, \ldots
\]

and energies:

\[
E_{\text{RAL}} = \frac{\hbar^2}{2\mathcal{J}(0)} (I - j_x)(I - j_x + 1).
\]

- \( K = \frac{1}{2} \) bands

In this case the Coriolis force has only a diagonal contribution to the energy. The energy is defined as:

\[
E = \frac{\hbar^2}{2\mathcal{J}(0)} \left[ I(I + 1) + \alpha(-1)^{I+\frac{1}{2}}(I + \frac{1}{2}) \right],
\]

with \( \alpha \) the decoupling parameter. Signature is now:

\[
r = (-1)^{I+\frac{1}{2}}
\]

and the spins of the band states are:

\[
I = \frac{1}{2}, \frac{3}{2}, \frac{5}{2}, \ldots
\]

Because the energy is in this case proportional to \( \left( I + \frac{(1+r\alpha)}{2} \right)^2 \) the spins of the signature partner bands are displaced in different directions along the momentum axis. Thus if the decoupling parameter is numerically larger than unity this displacement leads to inversion of the normal spin sequence [Bo75, Pa10].
CHAPTER 2. THEORETICAL FRAMEWORK

2.3 Magnetic dipole moment and gyromagnetic ratios

Measurements of the magnetic dipole moments and the gyromagnetic ratios (g-factors) of a nucleus and their comparison with theoretical predictions yield useful information on nuclear structure.

The magnetic moment of a single nucleon consists of an orbital and a spin part [Bo69]:

\[ \vec{\mu} = g_l \vec{l} + g_s \vec{s}, \]  \hspace{1cm} (2.20)

where \( g_l \) is the orbital g-factor and \( g_s \) the spin g-factor in units of nuclear magnetons \((\mu_N = e\hbar/(2m_p) = 5.0508 \times 10^{-27} \text{JT}^{-1})\). The bare orbital g-factors are \( g_{l}^p = 1\mu_N \) for protons and \( g_{l}^n = 0\mu_N \) for neutrons. For a free nucleon the spin g-factors are \( g_{s}^p = 5.58\mu_N \) and \( g_{s}^n = -3.82\mu_N \) for the proton and neutron respectively, whereas \( s = 1/2 \) for fermions. The spin g-factor for a bound nucleon can vary significantly from the values above, but generally \( g_s^p \) is positive and \( g_s^n \) negative.

The magnetic dipole moment of a nucleus is defined as [Ej89]:

\[ \mu(I) = \left[ g_R I + (g_K - g_R) \frac{K^2}{I+1} \right] \mu_N, \]  \hspace{1cm} (2.21)

where \( g_K \) arises from the single particle contribution in the nuclear magnetic dipole moment due to electric currents generated by the nuclear motion and \( g_R \) refers to the collective rotation of the nucleus. \( g_R \) is usually approximated by [Ej89]:

\[ g_R \simeq k \frac{Z}{A}, \]  \hspace{1cm} (2.22)

where the quenching factor \( k \) is roughly 1.

In an odd nucleus (e.g. \(^{251}\text{Md}\) described earlier) the B(M1)/B(E2) ratios can be used to determine the \( g_K \) factor (Equation 1.1). By comparing this factor with theoretical calculations within a certain nuclear model the single-particle configuration can be assigned.

As an example of this process the assignment of the two-quasiparticle configuration of the \( K^\pi = 8^- \) isomer in the even-even nucleus \(^{250}\text{Fm}\) taken from [Ro09]
is briefly explained. $B(M1)/B(E2)$ ratios are produced for all the observed transitions from the measured intensities. These values are also theoretically calculated from Equations 2.28 and 2.29 by hand or using the program $BM1BM2$ [Pa09b] that is based on [Dö83, Dö87]. In this approximation an intrinsic representation of the radiative multipole operators based on the rotor-plus-particle coupling concept is used to calculate electromagnetic transition amplitudes with cranking states.

The intrinsic electric quadrupole moment used in the calculations is taken as $Q_0=12.6$ eb and the quenching factor in Formula 2.22 as $k=1$ or $k=0.8$. $k=0.8$ was used because rotational nuclei with $140 \leq A \leq 200$ are found to have $g_R$ values consistently lower than $g_R = Z/A$ [Bo69].

The $g_K$ values of the single-particle states involved are calculated with the $swbeta$ program [Čw87]. This program uses a Hamiltonian including an axially deformed Woods-Saxon potential, the spin-orbit interaction and the Coulomb potential for protons to calculate single-particle energies, wave functions, LDM constants, effective barriers for the unbound states, single-particle quadrupole moments and $g$-factors. The single-particle $g_K$ values computed with the program are combined using the generalised Landé formula (Equation 2.23) to give the $g_K$ value of the two-quasiparticle excitations, labeled as $g_J$:

$$g_J = \frac{1}{2}(g_1 + g_2) + \frac{1}{2}(g_1 - g_2) \left[ \frac{j_1(j_1 + 1) - j_2(j_2 + 1)}{J(J+1)} \right], \quad (2.23)$$

where $g_1$ and $g_2$ are the $g_K$ values of the two quasi-particle states and $j_1$ and $j_2$ their spins [Kr88].

For $^{250}$Fm the comparison of the experimental with theoretical results is made in Figure 2.7. The intensity ratios could be experimentally determined for the $16^-$, $15^-$ and $14^-$ states. The figure shows the comparison of theory and experiment for all three of these states. Within uncertainties a two neutron, $\frac{7}{2}^+[624]_\nu \otimes \frac{9}{2}^-[734]_\nu$, and two proton, $\frac{7}{2}^-[514]_\pi \otimes \frac{9}{2}^+[624]_\pi$, configurations can be assigned to this band. The two neutron configuration is favoured by both the quenched and unquenched results, so it can be concluded that the $K^\pi = 8^-$ band is formed by the two neutron configuration.
Figure 2.7: Relation between the intensity ratio, $B(M1)/B(E2)$, and the $g_K$ factor for the $16^-$, $15^-$ and $14^-$ states of the $K^\pi = 8^-$ isomer of $^{250}$Fm. The experimental results (blue lines and dotted lines their errors) are compared with the two neutron $^7_2[624]_\nu \otimes ^9_2[734]_\nu$ configuration (red circles) and the two proton $^7_2[514]_\pi \otimes ^9_2[624]_\pi$ configuration (green squares). The unquenched and quenched values are as explained in the text. Figure adapted from [Ro09].
2.4 Electromagnetic transitions

SAGE will be used in experiments employing fusion evaporation reactions. In a fusion reaction the projectile and target nuclei interact strongly forming a compound nucleus in an exited state with a lifetime between $10^{-19}$ and $10^{-16}$ s. The compound nucleus releases its excess energy and angular momentum by evaporation of particles and emission of $\gamma$ rays. Depending on its mass the decay can progress through different channels as described in Figure 2.8.

In lighter systems the compound nucleus decays to a particle-stable evaporation residue through light-particle emission, namely neutron, proton and $\alpha$-particle emission with fission increasingly competing at heavier systems [Fr96].

After the internal energy becomes less than the particle emission threshold the dominant mode of deexcitation is via electromagnetic transitions. The nucleus loses energy rapidly mainly by the emission of E1 $\gamma$ rays (about $10^{-15}$ s after particle evaporation). After roughly $10^{-12}$ s it reaches the yrast line [Gr67] and starts decaying towards the ground state which it reaches after about $10^{-9}$ s. In some cases the nucleus might populate an isomeric state\(^1\) and thus reach the ground state after a longer time period.

Electromagnetic transitions can proceed via several mechanisms: (1) $\gamma$-ray emission, (2) Internal conversion and (3) Internal pair formation (alternatively called electron-positron pair emission and only possible if the excitation energy is above 1.02 MeV). The first two will be described, as they are the processes studied with SAGE. Internal pair formation is not discussed here, but can be found in [Ro49, Wi66].

2.4.1 $\gamma$-ray emission

In the process of $\gamma$-ray emission a transition of multipolarity $L$ is allowed between two states of angular momenta ($I_i, I_f$) and parities ($\pi_i, \pi_f$), where $i$ denotes the initial and $f$ the final states, if it obeys a set of selection rules [Kr88, Mo66].

\(^1\)Isomeric is an excited state whose half-life is long compared to most other excited states [Wa99].
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Figure 2.8: Production and decay mechanisms of a compound nucleus [Pa10].

- Conservation of angular momentum:

  \[ \vec{I}_i = \vec{I}_f + \vec{L} \]  \hspace{1cm} (2.24)

- Triangle rule:

  \[ |I_i - I_f| \leq L \leq I_i + I_f \]  \hspace{1cm} (2.25)

- Parity rule:

  \[
  \Delta \pi(EL) \equiv \pi_i \cdot \pi_f = (-1)^L \\
  \Delta \pi(ML) \equiv \pi_i \cdot \pi_f = (-1)^{L+1}
  \]  \hspace{1cm} (2.26)

The above apply for all but monopole transitions. M0 transitions do not exist [Ku93] and E0 transitions can only proceed via internal conversion (see Subsection 2.4.2).
Another important observable is the transition probability. This is especially useful when angular correlations or linear polarisation measurements cannot be used for the determination of the type or multipole order of a transition (see for example [Mo76] for a detailed description of these methods). The total transition probability, $T(\sigma L)$, is a function of the reduced transition probability, $B(\sigma L)$, and for the first five values of $L$ they are related as shown in Table 2.1 [Bo69, Mo76].

Table 2.1: Relation between $T(\sigma L)$ and $B(\sigma L)$. $E$ is in MeV, $T(\sigma L)$ in $s^{-1}$, $B(EL)$ in $e^2\text{fm}^{2L}$ and $B(ML)$ is in units of $\mu_N^2\text{fm}^{2L-2}$.

<table>
<thead>
<tr>
<th>$T(E1)$</th>
<th>$1.59 \times 10^{15}$</th>
<th>$E_1^3$</th>
<th>$B(E1)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T(E2)$</td>
<td>$1.22 \times 10^9$</td>
<td>$E_2^5$</td>
<td>$B(E2)$</td>
</tr>
<tr>
<td>$T(E3)$</td>
<td>$5.67 \times 10^2$</td>
<td>$E_3^7$</td>
<td>$B(E3)$</td>
</tr>
<tr>
<td>$T(E4)$</td>
<td>$1.69 \times 10^{-4}$</td>
<td>$E_4^9$</td>
<td>$B(E4)$</td>
</tr>
<tr>
<td>$T(E5)$</td>
<td>$3.43 \times 10^{-9}$</td>
<td>$E_5^{11}$</td>
<td>$B(E5)$</td>
</tr>
<tr>
<td>$T(M1)$</td>
<td>$1.76 \times 10^{13}$</td>
<td>$E_1^3$</td>
<td>$B(M1)$</td>
</tr>
<tr>
<td>$T(M2)$</td>
<td>$1.35 \times 10^7$</td>
<td>$E_2^5$</td>
<td>$B(M2)$</td>
</tr>
<tr>
<td>$T(M3)$</td>
<td>$6.28 \times 10^0$</td>
<td>$E_3^7$</td>
<td>$B(M3)$</td>
</tr>
<tr>
<td>$T(M4)$</td>
<td>$1.87 \times 10^{-6}$</td>
<td>$E_4^9$</td>
<td>$B(M4)$</td>
</tr>
<tr>
<td>$T(M5)$</td>
<td>$3.79 \times 10^{-13}$</td>
<td>$E_5^{11}$</td>
<td>$B(M5)$</td>
</tr>
</tbody>
</table>

The reduced transition probability and decay half-life ($\tau$) are related as [Bo69]:

$$B(\sigma L) \propto (E^{2L+1} \cdot \tau)^{-1}. \quad (2.27)$$

Rough estimates of transition probabilities (called Weisskopf estimates) are derived in [We51]. In [Mo76] these are repeated together with estimates for the reduced transition probabilities and tabulated values for the first five electric and magnetic transitions as shown in Table 2.2.
Table 2.2: The Weisskopf estimates of $T(\sigma L)$ and $B(\sigma L)$. $E$ is in MeV, $T(\sigma L)$ in $s^{-1}$, $B(EL)$ in $e^2 fm^{2L}$ and $B(ML)$ is in units of $\mu^2 N fm^{2L-2}$.

<table>
<thead>
<tr>
<th>$\sigma L$</th>
<th>$B^W(\sigma L)$</th>
<th>$T^W(\sigma L)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>E1</td>
<td>$6.45 \times 10^{-2} A^{2/3}$</td>
<td>$1.02 \times 10^{14} A^{2/3} E^3$</td>
</tr>
<tr>
<td>E2</td>
<td>$5.94 \times 10^{-2} A^{4/3}$</td>
<td>$7.23 \times 10^{7} A^{4/3} E^5$</td>
</tr>
<tr>
<td>E3</td>
<td>$5.94 \times 10^{-2} A^2$</td>
<td>$3.37 \times 10^{1} A^2 E^7$</td>
</tr>
<tr>
<td>E4</td>
<td>$6.29 \times 10^{-2} A^{8/3}$</td>
<td>$1.06 \times 10^{-5} A^{8/3} E^9$</td>
</tr>
<tr>
<td>E5</td>
<td>$6.93 \times 10^{-2} A^{10/3}$</td>
<td>$2.38 \times 10^{-10} A^{10/3} E^{11}$</td>
</tr>
<tr>
<td>M1</td>
<td>$1.79$</td>
<td>$3.12 \times 10^{13} E^3$</td>
</tr>
<tr>
<td>M2</td>
<td>$1.65 A^{2/3}$</td>
<td>$2.21 \times 10^{7} A^{2/3} E^5$</td>
</tr>
<tr>
<td>M3</td>
<td>$1.65 A^{4/3}$</td>
<td>$1.03 \times 10^{1} A^{4/3} E^7$</td>
</tr>
<tr>
<td>M4</td>
<td>$1.75 A^2$</td>
<td>$3.25 \times 10^{-6} A^2 E^9$</td>
</tr>
<tr>
<td>M5</td>
<td>$1.92 A^{8/3}$</td>
<td>$7.29 \times 10^{-13} A^{8/3} E^{11}$</td>
</tr>
</tbody>
</table>

Equation 1.1 from Chapter 1 comes from the reduced transition probabilities as calculated within the geometric model [Bo75]. These are expressed as:

$$B(M1; KI_i \rightarrow KI_f) = \frac{3}{4\pi} \left( \frac{e\hbar}{2Mc} \right)^2 (g_K - g_R)^2 K^2 \langle I_i K 10 \mid I_f K \rangle^2,$$  \hspace{1cm} (2.28)

for M1 transitions and

$$B(E2; KI_i \rightarrow KI_f) = \frac{5}{16\pi} e^2 Q_0^2 \langle I_i K 20 \mid I_f K \rangle^2$$  \hspace{1cm} (2.29)

for E2 transitions.

For a rotational band with a quantum number $K \neq 1/2$ the spectroscopic electric quadrupole moment can be described as [Ej89]:

$$Q(I) = \frac{3K^2 - I(I + 1)}{(I + 1)(2I + 3)} eQ_o(K)$$  \hspace{1cm} (2.30)

For the same band the magnetic dipole moment is defined as in Equation 2.21.
2.4.2 E0 transitions

The E0 transitions are the only allowed transitions between two $0^+$ states. They do not change the angular momentum of the nucleus but alter the nuclear surface [Mo76]. E0 components are also observed in transitions between states of the same spin and parity in competition with M1 and E2 components.

E0 transitions mainly proceed through internal conversion. The photon has unit intrinsic spin; so a transition between two states with zero spins and even parities through the emission of a single $\gamma$ ray is strictly forbidden. Additionally it can proceed through the emission of two $\gamma$ rays, double internal conversion or simultaneous $\gamma$-ray and electron emission. However, all these processes are orders of magnitude less likely to happen than internal conversion. E0 can also take place through internal pair formation if that is energetically allowed [Go66].

The absolute transition probability, $W(E0)$, depends on the electronic factor and the monopole transition strength. The electronic factor, $\Omega$, is not completely independent of nuclear properties as it is described by the electron wave functions within the nucleus. This is the case because the monopole interaction exists only while the electron is within the nuclear charge distribution [Ch56]. The monopole transition strength $\rho$ depends on the nuclear radius ($R \simeq 1.2A^{1/3}$) and the position vector of each proton, $\rho \approx \sum_p \langle r^2_p/R \rangle$ [Go66].

$$W(E0) = \frac{1}{\tau(E0)} = \rho^2(E0) \times [\Omega_{ic}(E0) + \Omega_\pi(E0)],$$

(2.31)

where $\tau(E0)$ is the partial mean life of the initial state for E0 decay and $\Omega_{ic}E(0)$ and $\Omega_\pi E(0)$ the electronic factors for internal conversion emission and electron-positron pair emission, respectively [Ki05].

2.4.3 Internal conversion

Internal conversion is the processes in which the electromagnetic multipole field of the nucleus interacts with the atomic electrons, resulting in an electron emission [Kr88]. It was first discovered and named as “conversion of the $\gamma$-radiation” by
Meitner and Hahn \cite{Ha24, Me24}. A few years later Hulme gave the first correct theoretical interpretation of the effect \cite{Hu32}, and together with Taylor and Mott set the foundation for the development of a relativistic theory on internal conversion \cite{Hu36, Ta32, Ta33}.

The emitted electron is one of the atomic electrons and unlike $\beta$ decay it is not created during the decay process. This implies that internal conversion is a threshold process with the threshold depending on the binding energy, $B_i$, of the atomic shell involved. The kinetic energy of the conversion electron is the energy of the transition (equal to the energy of the corresponding $\gamma$ ray if the nuclear recoil is not considered) minus the binding energy of the electron \cite{Kr88, Ej89}.

$$T_e = \Delta E - B_i,$$

(2.32)

$B_i$ depends on the electronic shell from which the electron comes and so conversion electrons are labelled according to the shell they originate from, K, L, M and so on. This means that a conversion electron spectrum shows discrete peaks belonging to different shells for each transition (unlike the continuum from $\beta$ decays). With sufficient resolving power, peaks from the different subshells can be discerned. An example of a high resolution spectrum taken from \cite{Gr60} is shown in Figure 2.9.

The vacancy left in the atomic shell after a conversion electron is emitted is filled rapidly by electrons from higher orbitals. This leads to the emission of X-rays (characteristic K X-rays when a K electron is emitted and so on) that can be used to identify internal conversion when studying a $\gamma$-ray spectrum.

Internal conversion and $\gamma$-ray emission are competing modes of decay. To show which one is preferred in each case, the internal conversion coefficient (ICC) is defined as the ratio of the electron emission rate to the $\gamma$-emission rate,

$$\alpha = \frac{\lambda_e}{\lambda_\gamma},$$

(2.33)

$\alpha$ is the total ICC giving the probability of electron emission relative to $\gamma$-ray emission. Since different electronic shells correspond to different binding energies a partial ICC
can be defined for each shell as:

\[ \alpha_i = \frac{\lambda_{ei}}{\lambda_\gamma}, \]  

(2.34)

with \( i \) representing the shell/subshell. The total electromagnetic decay rate can be written either using the total \( \text{ICC} \) or partial \( \text{ICCs} \) as:

\[ \lambda_t = \lambda_\gamma (1 + \alpha) = \lambda_\gamma (1 + \alpha_K + \alpha_{L_1} + \alpha_{L_{II}} + \alpha_{L_{III}} + ...). \]  

(2.35)

The study of \( \text{ICCs} \) has many useful applications both in nuclear structure and
in other fields. By comparing experimental and calculated values it is possible to determine multipolarities and mixing ratios of transitions, which can lead to spin and parity assignments of excited nuclear states. Knowledge of accurate ICCs is needed in determining total transition rates, and are also used in nuclear reaction calculations, nuclear medicine, environmental control and the nuclear fuel cycle [Ba02, Ki08, Ge08].

A simple nonrelativistic way to define the ICCs in the case of a point nucleus for energies above thresholds gives:

\[
\alpha_{EL} \approx \frac{Z^3}{n^3} \left( \frac{L}{L+1} \right) \left( \frac{e^2}{4\pi\epsilon_0}\frac{\hbar c}{E} \right)^4 \left( \frac{2m_e c^2}{E} \right)^{L+5/2}, \tag{2.36}
\]

for the electric multipoles and

\[
\alpha_{ML} \approx \frac{Z^3}{n^3} \left( \frac{e^2}{4\pi\epsilon_0}\frac{\hbar c}{E} \right)^4 \left( \frac{2m_e c^2}{E} \right)^{L+3/2}, \tag{2.37}
\]

for magnetic multipoles [Kr88].

More realistic definitions of the ICC can be found in Rösel et al. [Ro78] using the relativistic Hartree-Fock-Slater method and in Band et al. [Ba02] using Dirac-Fock calculations.

From Equations 2.36 and 2.37 some general characteristics of the ICCs can be seen [Ro66]:

- They increase rapidly with increasing nuclear charge (atomic number \(Z\)).
- They increase with multipole order (\(L\)).
- They decrease with increasing transition energy (\(E\)).
- They decrease for higher atomic shells (\(n\)).

These general features are present in all ICC calculations. ICCs deviate from this pattern at threshold energies where the ejected electron has zero kinetic energy [Ro78].

The dependencies of the internal conversion coefficients on physical parameters are shown in Figure 2.10, where the ICCs for different elements, transition multipolarities and transition energies are compared. Figure 2.11 concentrates on just one superheavy
element, but similar behaviour is present in all the heavy nuclei. The high ICCs for these elements, especially at low transition energies, emphasise the importance of studying internal conversion in this region of the chart of nuclei.

The dependence of the energy of conversion electrons on the binding energy is clearly seen in Figures 2.10 and 2.11. The sudden increase on the ICCs seen in these figures takes place when the transition energy becomes equal to the binding energy of the next available atomic electron shell. These thresholds are obvious for fermium at the lower multipolarity transitions. As the multipolarity increases the ICCs of higher shells dominate over the ones from lower shells and a threshold is not visible in the plot. For example for an 145 keV E1 transition in fermium the $\alpha_K$ is one order of magnitude greater than $\alpha_L$ and $\alpha_M$, whereas for an E5 transition of the same energy $\alpha_K$ is seven orders of magnitude smaller than both of them.

When the nucleus is considered as having finite size there is a non-zero probability for electrons penetrating into the nucleus. The ICCs calculated using Equations 2.36 and 2.37 and in [Ro78] do not take into account the nucleus-electron interaction while the electron is inside the nucleus. The surface current model [Sl51] used in the calculations in [Ba02] incorporates this interaction approximately.

For highly hindered transitions this effect is significant and corrections are needed in the calculations. For unhindered transitions it is still quite important for M1 transitions and less for M2, M3 and higher multipolarities [Ba02]. To estimate the magnitude of this effect, calculations that use no-penetration models are compared with ones that use surface current models. The effect for an 100 keV M1 transition varies from $\sim 0.01\%$ for light elements to $\sim 16\%$ for superheavy elements [Ba02, Ki08].

The most widely used ICC tables are those of Hager and Seltzer [Ha68], Rösel et al. [Ro78] and Band and Trzhaskovskaya [Ba78]. All these calculations use the relativistic Hartree-Fock-Slater method. More recently Band et al. [Ba02] published tables that show great improvements in accuracy using the Dirac-Fock method. The most precise calculations so far are made with the BrICC conversion coefficient calculator [Ki08] using Dirac-Fock. This is shown in [Ge08] where a comparison of the above-mentioned
calculations with measured values is made.

Here a hypothetical example of multipole order and multipole mixing ratio assignment is shown. Let us consider that in an internal conversion experiment studying fermium a 200 keV transition was measured. From the absolute values of ICCs and their ratios $\alpha_i/\alpha_j$ (see Figure 2.12 top diagram) for this transition one can assign its multipolarity [Ej89]. This can also be made using the normalised values of ICCs (normalised so that for every transition $\sum \alpha_i = 1$), shown in the bottom diagram of Figure 2.12, and comparing the theoretical and experimental patterns for the L_I, L_{II} and L_{III} shells. For mixed multipole transitions the multipole mixing ratio, $\delta^2$, can also be assigned. This is possible because the conversion coefficients are sensitive to the multiplicities but are independent of the transition matrix elements.

It can be proved that the $M1, E2$ multiple mixing ratio can be calculated from the $\alpha_{L_I} : \alpha_{L_{II}}$ ratios using the relation:

$$\delta^2 = \frac{\alpha_{L_I}^c(M1) - \alpha_{L_{II}}^c(M1) \cdot \alpha_{L_{II}}^m/\alpha_{L_{II}}^m}{\alpha_{L_{II}}^c(E2) \cdot \alpha_{L_{II}}^m/\alpha_{L_{II}}^m - \alpha_{L_I}^c(E2)}$$  \hspace{1cm} (2.38)

where the ICCs marked with superscript $m$ refer to measured values and those marked with superscript $c$ refer to calculated values.
Figure 2.10: Internal conversion coefficients for electric (top) and magnetic (bottom) transitions with multipolarity $L \leq 5$ as a function of transition energy for elements with $Z=20$ and $Z=100$ [Ki08]. The positions of the bound electron shells are indicated.
Figure 2.11: Internal conversion coefficients for E2 transitions (top) and M1 transitions (bottom) in fermium (Z=100) with respect to transition energy.
Figure 2.12: Absolute (top) and normalised, so that for every transition $\sum \alpha_i = 1$, (bottom) internal conversion coefficients for different multiples for a 200 keV transition in fermium.
Chapter 3

Concept of an ice and $\gamma$-ray spectrometer

3.1 Historical review of $\gamma$-ray and electron spectrometers

The first great leap in the detection efficiency and resolution of charge particle and $\gamma$-ray detectors was made with the introduction of solid state detectors. For $\gamma$ rays, organic (plastic) and inorganic scintillators (NaI(Tl), BaF$_2$, BGO etc.) and semiconductor detectors (Ge(Li) and HPGe) were used [Ne66, Kn00]. For charged particles, semiconductors like silicon were found to be ideal in most cases [Gi66, Kn00].

A major advancement in the detection of $\gamma$ rays was the construction of large $\gamma$-ray detector arrays surrounding the target region [No94]. The first generation of such arrays consisted of crystal scintillators coupled to photomultiplier tubes (PMTs), for example NaI arrays such as the crystal balls at Argonne [Ja83] and Heidelberg [Si80] and BaF$_2$ arrays such as the Chateau de Cristal in Strasbourg [Be84].

The use of HPGe detectors with anti-Compton suppression shields (initially with NaI and later on using BGO crystals [No94]) led to the second generation arrays such as tessa [Tw84] and Hera [Di84] followed by the third generation arrays like
CHAPTER 3. CONCEPT OF AN ICE AND $\gamma$-RAY SPECTROMETER

Gammasphere [Le90] and Euroball [Eb92]. Fourth generation arrays that use \(\gamma\)-ray tracking (AGATA [Si05] and Greta [Le03]) are under development.

The story in electron spectroscopy has been somewhat different. The necessity of discriminating electrons from other charged particles and reducing the low energy electron background has led to the use of magnetic and electric fields in combination with a detector arrangement.

The first semicircular spectrometers [Da12, Da13] were built almost a century ago, but they lacked space-focusing. Later on, long and short lens spectrometers helped to solve this problem [Mi79]. Better focusing and dispersion was achieved with the double focusing \(\pi\sqrt{2}\) spectrometer [Si46] and the orange spectrometer [Vi56, Ko65].

Electron spectrometers can be separated into mainly two categories: devices that use longitudinal magnetic fields to transport the electrons to the detector and devices that use fields transverse to the electron trajectories [Si66].

Examples of spectrometers from both categories can be found in [Ej89] and [Mi79]. A good example of the first type is SACRED [Bu96, Ka04] operated in Jyväskylä. Orange spectrometers and the more compact mini-oranges [Ki72] are the best examples of the second type. Successful use of mini-orange arrays has also been reported [Ga01]. Spectrometers based on different magnet arrangements than in the orange spectrometer also fall in this category [Ki97].

The mini-orange spectrometer uses permanent magnets to guide electrons to a silicon detector positioned behind a central absorber. The magnets act as a filter to separate electrons from positrons and the absorber reduces the amount of \(\gamma\)-rays and other background radiation from reaching the detector. This spectrometer however only transmits electrons in a narrow energy window and in a small solid angle. This problem can be overcome by the use of smaller magnets relative to the detector size. A simple representation of a mini-orange filter is shown in Figure 3.1.

Combined \(\gamma\)-ray and electron spectrometers are very powerful tools in spectroscopy. Magnetic spectrometers have been combined with a small number of \(\gamma\)-ray detectors in the past giving many interesting results. Drastically improved results can be achieved
by combining electron spectrometers with $\gamma$-ray detector arrays.

An example of this is ICEBall [Me93], a mini-orange array that can be used in conjunction with large $\gamma$-ray detector arrays like the Spin Spectrometer and GAMMASPHERE. It consists of six mini-orange spectrometers and the same number of Si(Li) detectors. Its compact geometry allows it to be positioned inside the target chamber of the $\gamma$-ray detector arrays.

Magnetic lenses with silicon detectors were combined in many cases with germanium detectors and scintillators in different geometries as described in [Di95, Di99].

A different approach was taken by a Polish collaboration [An08] combining quasi-mini-oranges (similar to mini-oranges only without the central absorber and using less magnets) with a solenoid field transporting the electrons to five detectors (2 Si(Li) and 3 PIPS-passivated implanted planar silicon detectors). This spectrometer can be coupled to the OSIRIS-II HPGe detector array.

SAGE falls in the category of magnetic lenses combined with germanium detectors. It is built on experience from previous devices trying to optimise its resolving power and efficiency both for electrons and $\gamma$ rays. Along with a relatively high electron transmission efficiency it is a prominent representative in the field of combined
electron and γ-ray spectroscopy.

3.2 SAGE design requirements

SAGE is a spectrometer that combines γ-ray and internal conversion electron spectroscopy. This is not an easy task as many different parameters and issues need to be optimised in order for such a spectrometer to be successful.

- γ-ray detection:
  To efficiently detect γ rays with energies less than 1 MeV it is essential to have as little absorbing material as possible in the path of the γ rays to the detector.

- Electron detection:
  A silicon detector is used for the detection of ICES but due to its high sensitivity to all types of radiation it needs to be positioned far from the target.

- Electron transport:
  To transport the ICES to the detector a magnetic field is required and so solenoid coils need to be implemented in the design.

- Reduction of electron background:
  Low energy delta electrons are produced when the beam particles interact with the target and the surrounding materials. Reaction kinematics cause most delta electrons to move at forward angles [Sc92, Ke92]. Placing the detector at a backward angle and using a high-voltage barrier between the target and the detector reduces the amount of delta electrons reaching the silicon detector.

- Reduction of Doppler broadening:
  Doppler broadening has a large effect in ICE spectroscopy. Using an almost collinear geometry between the magnetic field axis and the beam axis reduces this.
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• Analysis techniques:
  To use the powerful analysis technique of recoil-decay tagging, SAGE needs to be combined with a recoil separator and a focal-plane spectrometer.

• High beam intensities:
  For the study of exotic nuclei with very low production cross-sections higher beam intensities are required. Digital electronics allow the data acquisition system to cope with the higher count rates. Rotating targets are also essential in this case.

Many compromises were made during the design as a result of the above restrictions not being independent of each other. For example an optimal magnetic field that would achieve maximum ICE transmission efficiency would require the target to lie within the solenoid coils. This would result in all the $\gamma$ rays being absorbed by the coil material, rendering the $\gamma$-ray detectors useless.

Each one of the above topics is approached and discussed in detail in the following sections.

3.3 Geometry of the setup

To make the description of the setup easier a rendered mechanical drawing of SAGE is shown in Figure 3.2, with its main parts being the germanium and silicon detectors. In the figure SAGE is combined with RITU and GREAT.

• Germanium-detector array
  The JUROGAM II germanium-detector array surrounds the target region and detects the prompt $\gamma$ rays. JUROGAM II consists of 15 Phase I Compton-suppressed germanium detectors [Be92, No94] and 24 fourfold segmented Clover detectors [Sh99]. To allow the use of magnetic coils close to the target one ring of five Phase I detectors is removed, leaving a total of 10 detectors in place (see Fig-
Figure 3.2: Schematic representation of the SAGE spectrometer, in conjunction with RITU and GREAT.

The γ-ray detection efficiency of the array with 10 Phase I and 24 Clover detectors is 5.5% at 1332 keV.

- Silicon detector

The internal conversion electrons are transmitted to the segmented silicon detector employing the magnetic field induced by solenoid coils. A near collinear geometry is used between the beam and solenoid axes to reduce Doppler broadening. The silicon detector is circular with 50 mm diameter and segmented into 90 individual segments. It is described in more detail in Section 4.1.

- Recoil separator and focal plane spectrometer

SAGE is combined with the RITU gas-filled recoil separator [Le95] and the GREAT focal-plane spectrometer [Pa03b]. RITU transports the fusion evaporation reaction products towards GREAT where they are implanted in DSSDs. Their subsequent decays are detected by the different detectors of GREAT. Using SAGE in conjunction with RITU and GREAT allows the use of the recoil-decay tagging technique [Si86, Pa95].

- RITU (Recoil Ion Transport Unit) is a QDQQ type separator using helium
at 1 mbar pressure as a filling gas. It was mainly built and designed for the study of heavy elements. It has maximum magnetic rigidity of 2.2 Tm, angular acceptance of 10 msr and dispersion 10 mm/% [Le95].

- The GREAT (Gamma Recoil Electron Alpha Tagging) focal-plane spectrometer is an arrangement of silicon, germanium and gas detectors. At the entrance of GREAT lies a multiwire proportional counter (MWPC) which is
responsible, along with the DSSDs, for distinguishing fusion reaction products from their subsequent decays and from scattered beam particles. The recoils are implanted in the two DSSDs of GREAT. Different gain settings can be applied at the DSSDs to measure the electrons, protons or alpha particles the implanted ions emit. An array of PIN diodes (p-type - intrinsic - n-type semiconductor) surrounds the DSSDs with approximately 30% geometrical efficiency and detects ICE that are emitted from the decaying recoils. Low energy $\gamma$ rays and X rays are measured by a planar double-sided germanium strip-detector situated directly behind the DSSDs. The GREAT Clover detector is positioned above the DSSDs and can detect higher energy $\gamma$ rays with greater efficiency than the planar detector. For the same purpose additional detectors (usually VEGA or Clover detectors) may be placed at either side of the spectrometer and behind the DSSDs [Pa03b].

### 3.3.1 Mechanical design of SAGE

A series of solenoid magnetic coils are used to transport the conversion electrons to the silicon detector positioned upstream of the beam. This is necessary because the silicon detector is sensitive to all types of radiation, therefore it cannot be placed near the target region otherwise the useful conversion electron signals would be submerged in the background. A compromise between maximising the transmission efficiency, reducing background noise and keeping within the available space had to be made. Other aspects such as support, cooling and powering of the coils were taken into account when finalising the design.

The design process was an iterative procedure starting with a general design of the setup, simulating the magnetic and electric fields and using these to simulate the transmission efficiency. If the outcome was not satisfactory the design was modified by altering the dimensions and position of the solenoid coils, the high-voltage barrier, the carbon foil unit and the beam line. Figure 3.4 shows a cross-section of the setup
where the different parts can be seen.

Figure 3.4: A cross-section of SAGE as seen from above. The different parts of the setup are as follows: “A” the target position, “B” the silicon detector, ”C” the carbon foil unit and “D” the high-voltage barrier. Surrounding the target region are the JUROGAM II germanium detectors and around the beam pipe are the solenoid coils. The beam (represented by a cone) passes next to the detector moving towards the target.

The space envelope available for the coils is defined by JUROGAM II, RITU and the beam-line diagnostics box which contains vacuum pumps and a Faraday cup. The JUROGAM II array is positioned at the target region, so the volume defined by the collimators of the Compton-suppression shields of the germanium detectors provides the space limits for the target chamber. The shaping of both the downstream and upstream coil (shown in Figure 3.4) depends on the space envelope outlined by the two outer rings of JUROGAM II detectors. As the target position is fixed the position of the gate valve connecting SAGE to RITU (see Figure 3.5) gives the maximum allowed length of the downstream coil. Similarly, the position of the beam-line diagnostics box restricts the length of the main and upstream coils.

The arrangement is schematically shown in Figure 3.5. Between the downstream coil and the RITU gate valve a bellows is needed to allow alignment adjustments, hence
the empty space in this region. Part of the large space between the main coil and the diagnostics table is taken by the detector chamber, but most of it is left empty. This is necessary in order for the setup to be positioned in and out of JUROGAM II.

In an ideal situation with no space restrictions, the magnetic field produced by the coils would be constant on the solenoid axis; in reality it is not so uniform and it shows valleys and peaks. In Figure 3.6 a comparison of the two is made. In both cases the detector is positioned in a region of lower magnetic flux density.

Ideally, the way to increase the transmission efficiency would be by applying a higher magnetic field between the target and RITU. This would act as a magnetic mirror reflecting some of the electrons moving towards RITU back to the detector. One such example is shown in Figure 3.6, where the single-coil geometry was modified to allow more current through some of the loops closest to RITU, thus boosting the field.

Figure 3.5: A schematic representation of the volume available for the solenoid coils. The gate valve connecting to RITU is on the right hand side and the beam diagnostics box on the left.

From the different types of magnetic coils commercially available, water-cooled copper coils were chosen for SAGE. In the first implementation of the SACRED spectrometer [Bu96] superconducting coils were employed, but were found to be difficult to work with and also require a lot more space than what was actually available. Permanent magnets were not an option either as they cannot provide the necessary
Figure 3.6: A comparison of the magnetic fields on the solenoid axis produced by a single coil, when no space restrictions apply, and by the more realistic space-restricted case. The field produced by a modified version of the single coil that boosts the field between the target and RITU is shown for comparison.

A relatively strong (0.5 T or more) and uniform magnetic field on the solenoid axis is essential for obtaining a high electron transmission efficiency. Additionally, if the magnetic field is not adequately strong the electrons will move in larger radii (see Equation A.20 in Appendix A) making the use of a bigger, and hence more expensive, silicon detector necessary. On the other hand if the field is too strong then the majority of the electrons will be concentrated in the central segments of
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Figure 3.7: A picture of the magnetic coils and target chamber during construction.

the detector. To ensure that the electron flux is more equally distributed across the detector, this is positioned outside the bore of the magnet in a region of lower magnetic flux density. This is important in increasing the maximum allowed count rate for the detector and also to reduce the pileup.

In a lower magnetic field the spiral motion of the electrons on the solenoid axis is elongated, i.e. the electrons form fewer circles in the perpendicular plane per unit of length. This means that the incident angle between the normal to the detector surface and the electron path reduces, decreasing the probability for backscattering [Wa68, Mi79]. Backscattering is angle dependent but appears to be energy independent in the energy region between 40 keV and 1200 keV [Ka57, Wa68, Ca86, Me93].

Simulated and measured fields of the magnetic coils are discussed in detail in Section 3.4. The effect of the magnetic field on the PMTs of the BGO Compton-suppression shields of JUROGAM II is described in the same section. This effect is reduced by using steel shielding around the coils.

Previous experience with the SACRED spectrometer showed that collinear geometry between the solenoid axis and the beam axis provides the best spectral response [Bu96] (see [Bu87, Di91] for more details). A perfect collinear geometry would require the beam to pass through a hole at the centre of the detector, remarkably reducing the detection efficiency. In SAGE a $3.2^\circ$ angle is adopted between the two axes. This
geometry will also decrease the high flux of background delta electrons, originating mainly from atomic collisions between beam and target particles [Ka04].

To further reduce the low energy background, a high-voltage barrier is positioned between the target and detector positions. This is discussed in more detail in Section 3.5.

The safe operation of a high-voltage barrier dictates its placement in a region of high vacuum, of the order of $10^{-6} - 10^{-7}$ mbar. A carbon foil unit is used to separate the high-voltage barrier region from the 1 mbar helium gas used in RITU (see Section 3.8).

Another effect of the collinear geometry is the Doppler shift of the electrons emitted from moving recoils. The unshifted electron energy, $E$, in the recoil rest frame can be derived using the Lorentz transformation for four-momentum and the invariance principle [Ka01]. These give,

\[
E = \frac{E' + m_e - \beta \cos \theta' \sqrt{E'^2 + 2m_eE'}}{\sqrt{1 - \beta^2}} - m_e,
\]

where $E'$ and $\theta'$ are the measured values in the laboratory frame and $m_e$ is the rest mass of the electron. If different emission angles cannot be measured (as is the case here) then an average angle $\theta'$ may be used.

The method used for defining the average angle is the same as described in [Ka01]. By solving Equation 3.1 for $\theta'$, this angle can be calculated with respect to the shifted and unshifted transition energies. Unshifted are the Doppler corrected energies measured from the $\gamma$ rays and shifted the ones measured from the conversion electrons. The Lorentz factor $\beta$ is calculated from the reaction kinematics. For each transition observed in both the electron and $\gamma$-ray spectra and therefore for each energy, a different average emission angle is calculated. A straight line is fitted through these calculated angles to obtain an average Doppler correction angle for all the energies.

A quick calculation of the average angle can be made using the Monte Carlo code SOLENOID, presented in [Bu96] and Section 3.6. This code is executed for the desired reaction and amongst other values it gives the average emission angle per energy of the electrons that are subsequently detected.
Despite the energy dependence of the Doppler correction angle and the backscattering due to the mirroring effects described in Section 3.4, the above method yielded correct electron energies within 0.1 keV or less of the calculated ones, when used for SACRED. In SACRED average angles between 150-160° were used. Simulations performed for SAGE indicate similar values in the energy region between 40 and 400 keV.

3.3.2 Support and position adjustment mechanism

A support structure was designed and built (Figure 3.8) that allows individual fine position adjustment of every major part of SAGE as well as precise placement of the whole unit inside the JUROGAM II array.

The schematic diagram in Figure 3.9 shows the different parts of the support structure and the allowed movement for each one. All the parts of the setup form a rigid unit positioned on the main support frame (indicated with the number 1 in the schematic). This is supported on caster wheels used for rolling it into position and for crude positioning adjustment. When in the final position, brass pushers are used to lift the whole structure up to adjust it to the required height. A lifting cradle can also be attached to the main support frame for craning SAGE into the experimental hall if needed.

A secondary support frame (2 in the figure) resides on the main one, with the main and upstream coil supports fixed to it. This can move parallel to the main frame, but no height adjustment is possible at this level. The downstream support structure (number three in the diagram) slides into its final position on rails attached to the secondary frame. This is necessary because of the shape of the target chamber’s back plate (drawn with dashed line in the schematic).

The main and upstream coils and detector chamber are fixed into position on the secondary frame and are used as the fixed points for adjusting the rest of the setup. The downstream coil can be adjusted precisely using a separate support plate (number 4) that allows movement on all three axes (Figure 3.10).

Alignment between the individual parts is ensured through spirit levels fixed on
the movable parts. A telescope is used to align the electron part of the setup with the beam line.

Figure 3.8: The support structure for the electron detector part of SAGE. The different adjustment planes are indicated on the figure, see Figure 3.9 for comparison.
Figure 3.9: A schematic diagram of the support structure used to allow precise position alignment of SAGE. Different parts of the support structure are numbered as follows: (1) the main support frame, (2) the secondary frame, (3) the movable support for the downstream coil and (4) the position adjustment plate for the same coil. The arrows show the allowed directions of movement. The slanted arrows indicate movement perpendicularly to the drawing.
Figure 3.10: The downstream coil adjustment plate. The upper plate can move on ball bearings relative to the lower one allowing adjustment on the horizontal plane. Vertical adjustment is achieved by jacking screws.
3.4 Magnetic field study

3.4.1 Magnetic field simulations

A three-dimensional model of the solenoid coils and of the components made from magnetic material in their vicinity was built. The coil design was optimised through simulations and measurements using a prototype. Only results from simulations using the finalised solenoid coil geometry (Figure 3.11) will be presented here.

![Figure 3.11: The finalised solenoid coil geometry. The magnetic shielding is also shown.](image)

The maximum current that will be used in the coils of SAGE is 1000 A but for most experiments 700 A is adequate to allow high transmission efficiency. The magnetic field on the solenoid axis for 700 A and 1000 A current through the coils is presented in Figure 3.12. The magnetic field strength is reduced when lower current is used, but the peak to valley ratios (mirror ratios) do not change. Using the notation of Figure 3.12 the mirror ratios are presented in Table 3.1 along with the same ratios when magnetic shielding is used; as discussed further in the text.

Knowledge of the peak to valley ratios is important as the possibility an electron will be mirrored when approaching a stronger field (e.g. magnetic mirror effect when
TABLE 3.1: Comparison of the peak to valley ratios for the magnetic field on the solenoid axis with and without magnetic shielding.

<table>
<thead>
<tr>
<th>Ratio</th>
<th>Without shielding</th>
<th>With shielding</th>
</tr>
</thead>
<tbody>
<tr>
<td>a/b</td>
<td>1.11</td>
<td>1.11</td>
</tr>
<tr>
<td>c/b</td>
<td>1.26</td>
<td>1.23</td>
</tr>
<tr>
<td>c/d</td>
<td>3.26</td>
<td>3.92</td>
</tr>
<tr>
<td>e/d</td>
<td>3.91</td>
<td>4.52</td>
</tr>
</tbody>
</table>

moving from the target towards the detector) or even trapped between two stronger field regions (e.g. magnetic bottle effect between peaks c and e) depends greatly on these mirror ratios. It can be easily proved that for a certain mirror ratio particles need to have a large component of velocity parallel to the field axis to not be mirrored. Consequently particles with insufficient energy are mirrored/trapped.

\[
\left| \frac{u_{||0}}{u_{\perp0}} \right| < \left( \frac{B_m}{B} - 1 \right)^{1/2}, \tag{3.2}
\]

where \(u_{||0}\) and \(u_{\perp0}\) are the components of velocity parallel and perpendicular to the field axis at the origin and \(B_m\) and \(B\) the maximum and minimum magnetic fields involved in the mirror ratio [Ja99].

The magnetic field flux lines when applying 1000 A current through the solenoid coils and without using magnetic shielding are shown in Figure 3.13. For higher flux densities the line spacing decreases, indicating that the field is mainly concentrated in the inner bore of the solenoids, weakening further away. The direction of the magnetic field is always tangential to the field lines. For reasons explained in Appendix B all the figures presenting magnetic lines use a totally collinear geometry between the beam and solenoid axes.

The magnetic field density equipotential lines as obtained from a simulation for 1000 A current are presented in Figure 3.14. Plots like Figure 3.12 are projections of
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Figure 3.12: A comparison of the magnetic field strength on the solenoid axis for 700 and 1000 A current through the coils.

As was mentioned in Section 3.3, the strength of the magnetic field and the field direction close to the photomultiplier tubes highly affect the propagation of electrons between the dynodes of each PMT. Some of the PMTs of the BGO shields of JUROGAM II are individually shielded by thin mu-metal cylinders. In order to weaken and redirect the magnetic field in the region of the PMTs 6 mm thick steel shielding plates are mounted as shown in Figure 3.11.

The direction of the magnetic field affects the force applied on the electrons inside the PMTs (Figure 3.15), as can be deduced from Equation A.10. The simulated effect of shielding on the direction of the magnetic field flux lines at 1000 A current

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1Mu-metal is a nickel-iron alloy with very high magnetic permeability.
Figure 3.13: Magnetic field flux lines without magnetic shielding. The density of the lines is greater where the magnetic field is stronger. The current through the coils is 1000 A.

is presented in Figure 3.16.

The maximum field at the region of the PMTs that are situated closest to the coils is reduced by an order of magnitude due to the shields. As an example the field strength at point A of Figure 3.14 is reduced from 35.08 mT to 2.94 mT when shielding is used. From tests performed using a prototype and later with the SAGE coils it was found that thicker shielding is necessary close to these PMTs. This extra shielding was designed and mounted on the JUROGAM II array.

Another purpose of magnetic shielding is to increase the field strength on the solenoid axis as shown in Figure 3.17 for 1000 A current. The field upstream of the detector chamber where the shields lie closer to the solenoid axis is enhanced. The effect of the shields is smaller on the downstream side where the shielding plates are situated further from the axis. The peak to valley ratios are different to when no shielding is used as shown in Table 3.1. The increase of the e/d mirror ratio
Figure 3.14: Equipotential lines of the magnetic field density for 1000 A current through the coils. The projection is made looking the setup from above so the bottom part of the main coil on the drawing is where the cooling outlets are positioned where no shielding is used. Point A indicates the position of the PMT closest to the solenoid coils.

leads to more electrons moving from the target towards RITU to be reflected towards the detector; even though the c/d ratio increases as well, the magnetic bottle effect reduces and both these factors increase the transmission efficiency.

The magnetic field on the solenoid axis was measured using a Hall-effect\(^2\) gaussmeter and compared to the simulated one. All the magnetic coils were used in the measurement but only the main coil shielding was present. A tool to position the Hall probe precisely on the solenoid axis was designed and the Hall probe was corrected for the magnetic field of the earth.

The very good agreement between simulated and measured magnetic fields at

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\(^2\)The production of a transverse potential gradient in a material by a steady electric current which has a component normal to the magnetic field is called the Hall effect [Sm72]
700 A current is shown in Figure 3.18. The discrepancies are mainly due to uncertainties in the positioning of the Hall probe. The error of the Hall probe is of the order of 1%. All errors are smaller than the size of the data points.
Figure 3.16: The magnetic field flux lines induced by the SAGE coils and employing the magnetic shielding plates. The current through the coils is 1000 A. Because the density of the lines is greater where the magnetic field is stronger, most of the magnetic field flux lines are directed through the shields. To show their direction outside the shield region a greater number of magnetic lines is used in this figure compared to Figure 3.13.
Figure 3.17: A comparison of the simulated magnetic field strength on the solenoid axis with and without shielding.

Figure 3.18: Measured and simulated magnetic fields on the solenoid axis.
3.4.2 Measurements using a prototype

Prototypes of SAGE were built to investigate the electron transmission efficiency and the magnetic shielding. These studies were made employing the SACRED solenoid coils.

The electron transmission efficiency measurements of the prototype are not directly comparable with SAGE as their geometries are not identical. In general, the SAGE coils are bigger and more uniform. One of the prototype setups is shown in Figure 3.19. In this setup an iron block was positioned inside the last coil (further right in Figure 3.19) to strengthen the magnetic field in the source region.

![Figure 3.19: Photograph of one of the prototype coil setups.](image)

The tests were made with a cooled to approximately -30°C, 25.4 mm² silicon PIN diode coupled to a PSC 761 preamplifier and a linear amplifier. A $^{133}\text{Ba}$ source was used and the current in the coils was between 560 A and 800 A. The main ice
energies of $^{133}$Ba are 45.0 keV, 75.3 keV, 124.6 keV, 240.4 keV, 266.9 keV, 320.0 keV and 350.5 keV. The magnetic field and transmission efficiency were simulated and then measured. The predicted and measured values follow the same trend, as shown in Figures 3.20 and 3.21. The discrepancies are due to system losses that were not taken into account in the simulations, like poor vacuum conditions, misalignment of the source and detector etc. A transmission efficiency between 10% and 5% in the 45-350 keV energy range was achieved.

![Transmission efficiency graph](image)

Figure 3.20: Measured and simulated electron transmission efficiency for the prototype presented in Figure 3.19.

The effect of the stray magnetic field induced by the prototype coils in the PMTs of the BGO shields of JUROGAM II and the amount of shielding needed to minimise it was studied. A BGO shield populated with PMTs in various positions was placed close to the coils at discrete angles allowing different distances between the PMTs and the coils.

In the worst case scenario, one of the PMTs was parallel to the coils and lying adjacent to them. In other cases the BGO shield was either at 45° or 90° with respect
to the solenoid axis. It was found that a 1 mm thick iron shield bent around the PMT significantly increased the overall performance. Additional shielding around the coils was used. A 2 cm thick iron shield was found adequate in most cases.
3.5 Electric field simulations

A large flux of low energy delta electrons is produced during the interaction of the beam and target particles. Like conversion electrons, these can be transported to the silicon detector generating a huge background in the low energy region. To reduce this flux an electric field gradient induced by a high-voltage barrier is employed between the target and the detector.

The energy of delta electrons depends on the energy of the projectile, $E_a$, the K-electron binding energy of the atoms of the target, $E_k$, and the electron and projectile masses $m_e$ and $m_a$ respectively. The maximum energy they can get for a certain target and projectile is given by [Kl69]:

$$E(\delta) = 4 \sqrt{\frac{m_e E_k}{m_a} + \frac{m_e E_a}{m_a}}.$$  \hspace{1cm} (3.3)

A typical reaction for the study of nobelium is $^{48}$Ca beam on lead target at 219 MeV. For this reaction Equation 3.3 gives maximum delta electron energy of approximately 69 keV.

The maximum voltage that can be applied to the SAGE high-voltage barrier is -50 kV, which is adequate to stop most of the low energy electrons. The high-voltage barrier geometry is presented in Figure 3.22 with the real barrier shown in Figure 3.23.

The high-voltage power supply is connected through a shielded, PVC-insulated cable to a resistor chain consisting of five 1 GΩ resistors. This in turn is connected to a ceramic high-voltage vacuum feedthrough to the beam pipe. The voltage is applied to the electrode using the horseshoe shaped connector shown in Figure 3.24. An unshielded PVC-insulated cable connects the feedthrough with the horseshoe which is placed inside the electrode.

Both the electrode and the horseshoe are made of stainless steel and the electrode has rounded edges. This is a precaution as field density increases at sharp edges and consequently increases the possibility of producing discharges. For the same reason the inner bore and edges of the electrode are highly polished.

A Noryl insulation sleeve surrounds the electrode and is longer than it, with
both edges extending further to ensure that no flashovers can occur between the high-voltage barrier and any grounded surfaces, such as the inside of the beam pipe. Such discharges could destroy the silicon detector or some of the preamplifiers. The Noryl insulator has dielectric constant of 2.7, breakdown voltage of 16-20 kV/mm [Ol97, Go98] and is 7.5 mm thick, more than enough to safely prevent a breakdown at maximum voltage. A thin aluminium grounding sleeve is wrapped on the outside of the insulation providing a more uniform grounding plane around the electrode.

The Noryl insulator has 0 V potential on one side and -50 kV on the other which causes a high stress potential gradient across it. Should a flashover occur either in the insulator or due to impurities in the vacuum, then the voltage will drop in the resistor chain that is attached to the high-voltage cable on the outside of the chamber thus quenching the discharge to prevent damage.

The high-voltage barrier can be considered as a cylindrical capacitor with capacitance given by:

$$ C = \frac{2 \pi \epsilon_0 \epsilon_k L}{\ln \frac{r_{\text{out}}}{r_{\text{in}}}} = 73.47 \text{ pF}, \quad (3.4) $$

where $\epsilon_0$ and $\epsilon_k$ are the dielectric constants of vacuum and Noryl respectively, $L$ is
Figure 3.23: A picture of the finalised high-voltage barrier.

the length of the barrier and \( r_{\text{out}}, r_{\text{in}} \) its outer and inner radii.

The barrier along with the resistor chain form a low-pass filter with characteristic time \( RC=367 \text{ ms} \) (Figure 3.25). This means that should a discharge occur it will take 367 ms for the barrier to charge up to 63 % of the input voltage or about 1.8 s to fully charge up.

All the individual parts of the high-voltage barrier form a rigid unit carefully machined to avoid the presence of any trapped air volumes between the individual parts. To further reduce this possibility, the unit is positioned inside the beam pipe using small protrusions on the outside of the grounding sleeve.

The high-voltage barrier is a crucial component of SAGE, since a sudden discharge of such a high voltage could destroy sensitive electronic equipment or even cause a fatality. To ensure a safe working environment everything is well shielded and insulated, and during the design process a number of simulations were conducted to recognise any faults and indicate where additional insulation would be required.

Three-dimensional simulations were made with OPERA 3D [VF07] where the high-
Figure 3.24: The horseshoe connector used to charge the high-voltage barrier. The cable plugs into the socket and is held in position by a grub screw.

Figure 3.25: The circuit formed by the resistor chain and the high-voltage barrier.

voltage barrier and the power cable were simulated along with the coil support, illustrated in Figure 3.26. With these simulations a design was chosen where high field concentrations are minimised. The resulting electric field and high voltage profile are presented in Figure 3.27, which was produced using FEMM [Me06].
Figure 3.26: A perspective view of the setup used for the OPERA 3D simulations. The high-voltage barrier, the cable and the beam pipe are shown.

Figure 3.27: The electric field potential of the high-voltage barrier. In this simulation the high-voltage supply cable is not taken into account.
3.6 Electron transmission efficiency simulations

A very important factor for the good performance of SAGE is optimisation of the electron transmission efficiency. To simulate the transmission efficiency, the Monte Carlo computer code SOLENOID [Bu96] was used. The geometry of the setup (detector, beam line, carbon foil unit, silicon detector) was integrated in the code. A cascade of electrons of various energies was produced and uniformly emitted at the target position. The amount of electrons that either reached the detector or were lost in the way (both through interactions with the electromagnetic fields or the surrounding material) was calculated.

As in the case of the magnetic field simulations, where different solenoid coil geometries were studied, different geometries were considered in the transmission efficiency’s case. Because the transmission efficiency depends on the magnetic and electric fields it was considered before performing any major changes on the solenoid coils and the high-voltage barrier.

The results presented here are for the finalised setup geometry. The solenoid coils are as described in Section 3.4 and the high-voltage barrier and carbon foil unit as in Sections 3.5 and 3.8 respectively. The silicon detector is described in Section 4.1 and is 1 mm thick. The simulated electron transmission efficiency is presented in Figure 3.28. The magnetic field was calculated for 1000 A current and the electric for -30 kV on the high-voltage barrier.

The transmission efficiency increases rapidly with increasing energy until about 160 keV, then until 360 keV it is roughly constant around 7.5%. It then gradually decreases with increasing energy. This behaviour can be explained as at lower energies more electrons are absorbed inside the target and carbon foils or are reflected by the electromagnetic fields. The magnetic field is not strong enough to constrain higher energy electrons thus the number reaching the detector decreases above 360 keV. Additionally the 1 mm thick silicon detector used in these simulations is efficient for electron detection up to roughly 500 keV, this also affects the amount of detected electrons are higher energies.
Figure 3.28: Simulated electron transmission and detection efficiency when using a 1 mm thick detector. Magnetic field calculated for 1000 A current through the coils and electric field for -30 kV on the high-voltage barrier.

The energy dependence of the major loss mechanisms is plotted in Figure 3.29. At higher energies the bottleneck is the radius of the carbon foil unit (aperture in Figure 3.29). At lower energies different mechanisms compete with the reflection of electrons from the electromagnetic fields being the most prominent. In the simulation output the same electron may be included in more than one category. For example the same electron can be reflected by the electromagnetic fields and consequently absorbed in the carbon foils. This can lead to the total losses to be presented as greater than 100%.

The electron distribution on the detector surface is also something that was con-
Figure 3.29: Percentage losses of electrons per energy as calculated from the simulations. Aperture is the carbon foil unit and EM fields the electromagnetic fields of the setup. The total losses are shown greater than 100% in some energies for the reasons explained in the text. 1000 A current through the coils and -30 kV on the high-voltage barrier were used in the simulations.

As expected at lower energies most of the electrons are measured by the inner detector rings. This is because the focusing power of the solenoid coils is proportional to the transverse momentum of the electrons \( r \propto \frac{p_\perp}{B} \), see Appendix A). At higher
energies the outer segments give higher count rates for the same reason. The central pixel acts similarly to the inner rings.

Due to the high delta-electron background higher segmentation is chosen for the central pixels. This allows more evenly distributed count rate over the detector. Rate distribution estimates are further discussed in Section 3.7.

It should be stressed that the transmission efficiency should be calculated separately for each SAGE experiment, because the calculated values depend on target thickness, recoil energy, angular distribution of the emitted electrons and different
electric and magnetic field strengths. The Monte Carlo code allows the user to do this very quickly and since the electric and magnetic fields only require scaling when different voltage and current is used the whole process is fairly straight forward.

3.7 Silicon detector rate distribution estimates

The SAGE silicon detector had to be designed in a way that allows the use of high intensity beams and offers an evenly distributed count rate throughout its surface. It was not possible to build prototype detectors because of the high manufacturing cost and long delivery times, so a set of calculations were made trying to find the best design.

As a starting point the SACRED silicon detector [Ka04] and the count rate distribution measurement from a $^{254}$No experiment [Bu02] using this detector was taken. It was found that for the SACRED detector the rate density as a function of segment radius follows the equation:

$$f(r) = \frac{a}{1 + \left(\frac{r}{b}\right)^3},$$

where $a$ and $b$ are constants. This function has no physical weight and was an educated guess. The agreement between data and function are shown better by integrating the function over the geometry of the detector. By applying each detector ring’s inner and outer radii as limits to the integration and then dividing the resulting value by the number of segments per ring the count rate per segment was calculated. All the above values are presented in Figure 3.31.

Other detector geometries were studied by integrating the function given in Equation 3.5 with different limits and the following constraints were applied to ensure realistic results. The angle of the beam and the solenoid axes was kept at 2.5° and the magnetic field properties were the same as in SACRED. The spacing between the detector segments was kept at 0.1 µm, the detector diameter at 60 mm and its distance from the target at 90 cm.

The final geometry selected gives the most evenly distributed count rate from all
Figure 3.31: Comparison between the measured and calculated count rate distributions of the SACRED detector. A fit using Equation 3.5 with $a=2.47$ and $b=3.34$ is also shown.

the studied geometries. By further dividing the inner rings and central pixel, higher count rates are achieved because the distribution does not peak in the centre, as shown in Figure 3.32.

A second set of calculations was made to study the effect of detector displacement on count rate distribution. The same method as above was used but this time a two dimensional grid was defined with each point of the grid having a relative value given by the count rate distribution function. The detector was displaced and the area covered by each segment integrated to give the new count rate. The displacement was made along the $22.5^\circ$ diagonal. This represents a worst case scenario as the count rate peaks mainly in one segment.
This exercise showed that the detector must be precisely positioned otherwise the count rate at various segments would increase drastically, as demonstrated in Figure 3.33. The results are normalised so that for a centred detector the count rate in the highest counting pixel equals one.

A fine position adjustment mechanism is integrated in the detector support to correct for any misalignment (see Figure 3.34). The support structure is mounted on rails connected to the detector chamber door and moves on the vertical and horizontal axes using stepper motors. The system allows for $\pm 6$ mm movement on both axes, restricted by limit switches.
Figure 3.33: Normalised count rate of the fastest counting segment per ring as a function of detector displacement. The displacement is made along the 22.5° diagonal. The normalisation is done so that the maximum count rate when the detector is centred equals one.
Figure 3.34: A picture of the silicon detector support frame where the fine position adjustment of the detector takes place. The frame is mounted on a dummy plate used during manufacture and testing.
3.8 The vacuum and pumping arrangement

Appropriate vacuum conditions are vital for the correct operation of SAGE. The bulk of the spectrometer can be separated into two main pumping volumes. RITU, that operates using 1 mbar helium gas and the high-voltage barrier region that requires a high vacuum for proper operation. These two volumes need to be well separated, while ensuring good electron transmission.

Good vacuum is necessary in the region of the high-voltage barrier as any impurities, mainly helium particles and water vapour, can cause discharges of the barrier. The beam can interact with residual gas molecules (contaminants) ionising them producing a flux of electrons which aided by the electric and magnetic fields are accelerated and transmitted to the detector, resulting in a large background.

In [Ka04] a study was made to find the best way to minimise the amount of helium in the region of the high-voltage barrier. It was shown that separating the two regions with a single 40 $\mu$g/cm$^2$ carbon foil gives at best $10^{-5}$ mbar pressure, which is inadequate. The carbon foil would collapse at higher pressure differences and helium gas leaked through it preventing better vacuum conditions.

This is overcome by employing a unit using two 50 $\mu$g/cm$^2$ carbon foils with intermediate pumping (Figure 3.35). Adequate pressure ($10^{-6}$ - $10^{-7}$ mbar) is achieved this way. To prevent leaks around the carbon foil unit a piston seal is used. Helium particles diffuse through the carbon foils at a slow rate but have no major effect on the pressure in the high-voltage barrier region.

Transmission efficiency simulations showed that the best position for the carbon foil unit is upstream of the target. If it is positioned downstream of the target it significantly reduces the recoil transmission into RITU and prevents gas cooling of the target.

To further improve the vacuum a cold trap (see Figure 3.36) is positioned below the detector chamber and cooled with liquid nitrogen through a cold finger to approximately -50$^\circ$C. The cold trap removes water vapour and gases from the pumping stream by sublimating the gas molecules on the trap surface. This means that the
gas molecules transform from the gas phase to a solid crystalline phase, bypassing the liquid phase.

If the cold trap was not present, any impurities would be attracted towards the cold surface of the detector. Impurities implanted on the detector surface could locally alter its electric field configuration, form conductive bridges between neighbouring pixels decreasing the inter-strip resistance [Ha09] or interact with the incident electrons, affecting the resolution of the detector.

The pumping scheme is shown in Figure 3.37. The spectrometer is separated from the beam line and RITU with gate valves. Pumping to $10^{-2}$-$10^{-3}$ mbar can be done separately for each volume using the rotary vane pump and lower pressures are reached using a 12001/s turbo pump.

In normal operation all volumes are pumped simultaneously first with the rotary vane pump through the needle valve and then with the turbo pump until equilibrium is reached throughout the spectrometer volume. Then the gate valve to RITU (that is in vacuum mode) is opened and while pumping the carbon foil unit with the rotary vane pump, helium is slowly introduced in the system.

Continuous monitoring of the pressure in each volume is made with manometers, from atmospheric pressure to a few millibar, and with Pirani/cold cathode full range gauges for pressures down to $10^{-7}$ mbar.

Figure 3.35: A schematic diagram of the carbon foil unit, showing the pumping port and the sealing groove.
By closing gate valves G1, G2 and the beam line gate valve one gains access to the detector without venting the whole system. In this case valve v2 allows continuous pumping of the high voltage region through the turbo pump. To gain access to the target region while keeping the detector chamber under vacuum gate valve G2 and the RITU gate valve are closed.
Figure 3.37: The vacuum control system for the SAGE spectrometer.
Chapter 4

Detector and Electronics

4.1 The SAGE silicon detector

4.1.1 The silicon detector properties

The silicon detector used in the SAGE spectrometer (Figure 4.1) is 1 mm thick and has a concentric ring structure. It is divided into 90 individual segments with the geometry shown in Figure 4.2. The numbering scheme of the detector pixels is presented in the same figure. The central pixel of the detector is divided into two semicircles with 1 mm radius (pixels 1 and 2). Following these are the 7 inner rings which are 1 mm wide and each one is split into 8 segments (pixels 3 to 58). The 8 outer rings are 2 mm wide and are divided into 4 segments each (pixels 59-90), making the overall diameter of the active part of the detector 48 mm. The capacitance of the larger detector segments is roughly 10 pF reducing to about 0.2 pF for the central pixel.

By taking a closer look at the detector one can see that the inter-strip gaps separating the segments are 70 µm wide. The way the separation is made is shown in Figure 4.3. The strip pitch is 1 mm measured from the end of one strip to the end of the next. For example let's consider the centre of the detector to be at the origin. The end of the central segment is at 1 mm, following this is a 70 µm wide inter-strip gap. The second strip is between 1.07 mm and 2 mm. Then an inter-strip gap and so on.
Taking into account the inactive areas that separate strips into individual segments gives a total inactive area for the detector of approximately 4%.

Surrounding the active segments of the detector are a series of guard rings [Ka67]. These shape the electric field, reducing edge effects and providing a homogeneous potential to all the strips, increase the breakdown voltage of the detector and reduce the leakage current by collecting its surface component [Br83, Mi05, Ha09]. These factors improve the overall performance of the silicon detector. The various technical design characteristics of the detector are presented in Table 4.1. This detector design was chosen for the reasons explained in Section 3.7.

Figure 4.1: Photograph of the SAGE silicon detector. The detector segmentation and signal wires are clearly visible.
Figure 4.2: The geometry of the SAGE silicon detector. The numbers and dimensions of the individual pixels are shown.

4.1.2 The detector Printed Circuit Board layout

A PCB was designed to accommodate the silicon detector and the 90 preamplifiers. The preamplifiers used are the CAEN A1422 that are described in Subsection 4.2.3. The layout is such that all the channels are equivalent with the preamplifiers positioned equidistantly from the detector and having individual power supply filtering. The circuit diagram of a single detector channel is shown in Figure 4.4 and the layout of the PCB in Figure 4.5. The outputs are divided into groups of ten channels and are taken through 9 25-way D-type connectors on the feedthrough plate of the vacuum chamber. These are connected to the PCB through flexible ribbon cables that are also
used for powering the preamplifiers. Micro coaxial cables are under consideration for replacing the ribbon cables as they are better shielded and will reduce noise pickup. The ribbon cables plug into 26-way side-latch headers on the PCB. On the outside of the chamber, small adaptor boxes connect into each of the D-type connectors to convert the D-type to SMA (sub-miniature version A) connectors.

The bias voltage is applied either directly through a bias resistor to the back side of the detector or through the preamplifiers. Both these methods are integrated in the PCB design so that if one method introduces noise in the system the other will be used. The two methods are interchangeable by means of a jumper but the default bias method is through the bias resistor.

The PCB is placed in high vacuum of the order of $10^{-7}$ mbar. The detector and preamplifiers are cooled through contact and radiative cooling using a refrigerated circulator with ethanol as the refrigerant. The ethanol circulates through the cooling plate as shown in Figure 4.6.
Table 4.1: Design technical specifications of the SAGE silicon detector as defined by the manufacturer.

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Outer diameter</td>
<td>50 mm</td>
</tr>
<tr>
<td>Detector thickness</td>
<td>1000 µm</td>
</tr>
<tr>
<td>Full depletion (FD)</td>
<td>150 V typical</td>
</tr>
<tr>
<td></td>
<td>200 V maximum</td>
</tr>
<tr>
<td>Total leakage current (FD+30 V at 20°C)</td>
<td>1 µA typical</td>
</tr>
<tr>
<td></td>
<td>3 µA maximum</td>
</tr>
<tr>
<td>Estimated capacitance</td>
<td>11 pF/cm² for Si</td>
</tr>
<tr>
<td></td>
<td>1 pF/cm² for tracks</td>
</tr>
<tr>
<td>Electron cut-off energy</td>
<td>4 keV</td>
</tr>
</tbody>
</table>

### 4.1.3 Offline detector testing

The detector was tested by the manufacturer by connecting all the segments together and measuring the overall capacitance and current at different bias voltages. The measurements were made in a light-sealed chamber, at 20-21°C temperature and humidity of 35-60%. From these, one can determine the full depletion bias voltage region and also how resistive the detector is. The results from these measurements are presented in Figure 4.7.

The detector becomes fully depleted around 128 V, as seen in Figure 4.7(a). This is explained if we consider that the depletion region of the detector simulates a charged capacitor. By increasing the reverse bias voltage the depletion region increases and the capacitance decreases [Kn00]. When the detector becomes fully depleted the capacitance is almost constant.

In Figure 4.7(b) the voltage-current characteristics of the detector are shown. Its operational region is between $V_{min}$ and $V_{max}$ which ideally should be flat. This would
allow the detector to be operated at higher bias voltages and hence have lower capacitance. This can be achieved by cooling the detector to reduce the leakage current and consequently the electronic noise and so improving the resolution. Beyond $V_{max}$ the current increases rapidly. This is due to ionisation of atoms in the detector material that produce free charge carriers. This decreases the resistance of the detector material causing an increasing current flow [Sm78].

The detector was bonded to the assembled PCB for further testing. After mounting the PCB to the detector cooling plate the preamplifiers were connected as shown in Figure 4.6. To ensure a quiet testing environment that additionally resembles the experimental conditions, the PCB was placed inside a screened box and the D-type to SMA converter boxes were used in the output.

Most of the channels were tested with cosmic radiation and all of them with the 59.54-keV $\gamma$-ray of an $^{241}$Am source. Oscilloscope screen-shots from the tests are presented in Figures 4.8 and 4.9. The tests were made in air and no detector cooling

Figure 4.4: Circuit diagram for one channel of the detector PCB, employing a CAEN A1422 preamplifier. The bias and preamplifier power supply filtering circuits can be seen.
was used. The power dissipation from the preamplifiers (276 mW per preamplifier) caused a temperature increase on the PCB which in the final implementation will be avoided by cooling.

The overall results of these tests are highly satisfactory. The completed, fully populated PCB with mounted detector and preamplifiers, has a low noise level (approximately 4 mV baseline noise) and no cross talk between the channels is observed.
Figure 4.6: Photograph of the detector PCB from the back side, with the preamplifier covers removed. The inlet and outlet of the coolant are indicated by arrows.

The random high frequency noise seen in the oscilloscope screen-shots will be filtered and averaged by the digital data acquisition system. The different amount of noise seen in the two figures is due to the different scales on the oscilloscope.
Figure 4.7: (a) Capacitance as a function of bias voltage. The detector is fully depleted when 128 V is applied. (b) Current as a function of voltage. The operational region of the detector lies between the two dotted lines. $V_{\text{max}}$ shows the point where the detector becomes resistive.
Figure 4.8: Oscilloscope screen-shot from the detector tests with cosmic radiation. A random detector channel was selected (Channel 2) where approximately 1 MeV was deposited.

Figure 4.9: Oscilloscope screen-shot from the detector tests with an $^{241}\text{Am}$ source where the 59.54-keV $\gamma$-ray was detected. A random detector channel was selected (Channel 2).
4.2 Preamplifiers tested for the silicon detector

In a silicon detector the charge produced by the incident radiation is too small to be used without further amplification [Kn00]. The first amplification stage in the signal chain is made using an FET followed by a preamplifier.

The output of each of the segments of the SAGE silicon detector is connected to a separate preamplifier. Because of the considerable number of channels (90 in total) it is essential the preamplifiers are compact so they can be placed in close proximity to the detector, on the same PCB if practicable. This will reduce the capacitive load on the detector compared to using long cables between the detector and preamplifiers and minimise the effect on signal to noise ratio. SAGE will be mainly employed in the study of low energy electrons, thus high gain preamplifiers need to be used. Additionally the preamplifiers are required to be able to operate in high vacuum.

The preamplifiers that fulfilled the above criteria were the Amptek A250F/NF, the Cremat CR-110 and the CAEN A1422 hybrid preamplifiers, as well as modified versions of the latter two with increased gain. Hybrid preamplifiers, in this case, means preamplifiers made by using printed and discrete electronic components on a ceramic interconnection substrate [Se95]. Technical specifications of these preamplifiers are described in Table 4.2. The physical dimensions of the preamplifiers are shown in Figure 4.10 and photographs of the chips in Figure 4.11.

![Figure 4.10: Schematic drawing of the tested preamplifiers with their dimensions.](image-url)
Table 4.2: Specifications table of the tested preamplifiers showing their gain, output and operational characteristics.

<table>
<thead>
<tr>
<th></th>
<th>A250F/NF</th>
<th>CR-110(61)</th>
<th>A1422(90)</th>
<th>A1422(200)</th>
<th>A1422(400)</th>
<th>A422</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gain [mV/MeV]</td>
<td>175</td>
<td>61</td>
<td>200</td>
<td>90</td>
<td>200</td>
<td>400</td>
</tr>
<tr>
<td>Noise in Si [keV]</td>
<td>&lt;2</td>
<td>&lt;2</td>
<td>&lt;2</td>
<td>&lt;2</td>
<td>&lt;2</td>
<td>&lt;10</td>
</tr>
<tr>
<td>$T_{\text{rise}}$ [ns]</td>
<td>3.80</td>
<td>6.55</td>
<td>6.55</td>
<td>12.00</td>
<td>12.00</td>
<td>12.00</td>
</tr>
<tr>
<td>$T_{\text{rise}}$ [µs]</td>
<td>250</td>
<td>140</td>
<td>1180</td>
<td>50</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>Output impedance [Ω]</td>
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<td>50</td>
<td>50</td>
<td>50</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>Power dissipation [mW]</td>
<td>20</td>
<td>±6 ±13</td>
<td>±6 ±13</td>
<td>±4</td>
<td>±12</td>
<td>±12</td>
</tr>
<tr>
<td>Operating voltage [V]</td>
<td>±15</td>
<td>±3.5</td>
<td>±7.5</td>
<td>±20</td>
<td>±30</td>
<td>±20, +30</td>
</tr>
<tr>
<td>Operating current [mA]</td>
<td>±4</td>
<td>±4</td>
<td>±4</td>
<td>±4</td>
<td>±4</td>
<td>±4</td>
</tr>
<tr>
<td>Power dissipation [mW]</td>
<td>276</td>
<td>276</td>
<td>276</td>
<td>276</td>
<td>276</td>
<td>276</td>
</tr>
</tbody>
</table>
The preamplifiers were tested using both a test pulse from a pulse generator and a detector signal. The main tests were made with a window-less silicon PIN photodiode (HAMAMATSU S3590-06), with active area of 9x9 mm and 25 pF capacitance. The preamplifiers were mounted on test PCBs placed inside a metal box, for shielding from electronic noise and light. A $^{133}$Ba source was used for testing the setup with electrons and some additional tests were made with alpha particles using an $^{241}$Am source and cosmic rays.

For the tests with the detector the setup was placed inside a vacuum chamber with an ultimate pressure of the order of $10^{-2}$ mbar and was cooled down to -30$^\circ$C (temperature at circulator) with a refrigerated circulator using ethanol as the refrigerant (photodiode temperature varied depending on cooling time). The preamplifiers were powered directly from a standard NIM crate with no additional filtering and the PIN diode was biased at +75 V.

The preamplifier output was connected to a spectroscopy amplifier and then to the analog or digital pulse processing electronics. The digital electronics used for the tests were the Lyrtech 16-channel VHS-ADC with Virtex-2 field-programmable gate array (FPGA) (a version of these cards with Virtex-4 FPGA is used with SAGE), the
4-channel Tracking Numerical Treatment (TNT2) card [Ar06] and the Gamma-Ray Tracking 4 channel (GRT4) VME card [La04]. The firmware for the Lyrtech digital electronics was still under development during the test period so the main bulk of the tests were made using a multichannel analyser with the MAESTRO-32 MCA emulation software.

The different preamplifiers tested for SAP are presented in the following subsections together with the results from the performed tests.

4.2.1 The AMPTEK A250F/NF

The A250F/NF was thoroughly tested as its high gain, small size and low power consumption made it a strong candidate for the preamplifier choice. For the first series of tests the AMPTEK PC250F test board (Figure 4.12) was used. Modifications were made to the detector bias supply and the preamp bias supplies. Low-pass bias filters and power supply decoupling capacitors (reservoir capacitors to provide low impedance for the preamplifier circuit and to filter the power supply) were added to the test board. These can be seen in more detail in the circuit diagram in Figure 4.13.

An in-house version of the AMPTEK test board that can accommodate 4 preamplifiers and uses a surface mount FET (MMBF4416) was used to further test the A250F/NF. The tests showed that the preamplifier is unable to drive a 50 Ω load over long distances as distortions to the signal occur. This was later verified by the manufacturer by confirming it has an output impedance of 100 Ω.

To overcome this problem buffer amplifiers were used for impedance matching. The quad channel Cadeka CLC4600 and the single channel Texas Instruments OPA820 and OPA842 amplifiers were tested. The buffers proved to have an effect on signal quality. In some cases overshooting or undershooting was observed, in others the exponential character of the signal was lost. Various methods were used to filter the output signal and compensate for the changes produced from the buffer amplifiers.

To take advantage of the full dynamic range of the digital electronics, additional gain and offset adjustment is required, as explained in Section 4.3. A version of the
Figure 4.12: The AMPTEK PC250F test board used for testing the A250F/NF preamplifier.

test board that takes these requirements into account and also filters the output was built (see schematic representation in Figure 4.14). This test board has two different gain stages. In the first one a CMOS operational amplifier allows high gain without greatly affecting the signal shape; a buffer amplifier is used in the second stage as a line driver. The offset adjustment is made by changing resistor values, so if this design was to be used in the final SAGE PCB an offset should have been decided beforehand. This test setup produced the best results observed using the A250F/NF. A typical output pulse from this is shown in the oscilloscope screen-shot in Figure 4.15.

The overall performance of the A250F/NF was acceptable but some of the better test results were not reproducible. If this preamplifier was to be used in SAGE, an additional buffering circuit needed to be accommodated on the detector PCB to drive the 50Ω load on the output of the preamplifiers. This preamplifier uses an external FET (unlike the ones presented next that have integrated FETs) that should also have been integrated in the detector PCB circuit. These would have made the PCB more difficult to manufacture and more prone to reliability issues.
4.2.2 The Cremat CR-110

Cremat provides the CR-150 evaluation board (shown in Figure 4.16) for testing this preamplifier. The tests with the Cremat preamplifier proved to be more straightforward than those described earlier for the A250F/NF. The CR-110 has a 50 Ω output impedance and so it can drive a 50 Ω load over long distances. No modifications were made to the CR-150 test board, as from the circuit diagram one can see that both the bias and the preamplifier power supplies are very well filtered.

The CR-110 performed consistently better than the A250F/NF. Non-exponential behaviour (similar to the one observed for the A250F/NF) was present in some cases.
in the output pulses of a CR-110 preamplifier. The oscilloscope screen-shot from Figure 4.17 is an example of such a pulse. These pulses are not desirable but were often seen with the CR-110 and A250F/NF.

4.2.3 The CAEN A422 and A1422

The first preamplifier received from CAEN was the A422 hybrid charge-sensitive preamplifier with a fast timing output. This preamplifier has adjustable gain of 1 mV/MeV, 45 mV/MeV and 90 mV/MeV and requires ±12 V and ±24 V to operate. The performance of the preamplifier was very poor and it was also found to overheat. This was attributed to two Zener diodes that were used as voltage regulators to provide the ±12 V from the ±24 V inputs (see Figure 4.18). The Zener diodes were removed and the ±12 V were externally supplied to the preamplifier, ultimately reducing the produced heat. The performance of this preamplifier was found to be insufficient showing two components in the rise time and very poor resolution.

To solve the problems of the A422, CAEN provided the A1422 hybrid charge-sensitive preamplifier. This preamplifier was smaller in size than the A422 and had a fixed gain of 90 mV/MeV. Its decay time constant was more than five times that of the A422 (1180 µs compared to 220 µs). The PCB shown in Figure 4.19 was specifically built to test this preamplifier which performed very well. Simple filtering was applied at the preamplifier power supplies, and no filtering used for the detector
bias, as the photodiode was powered through the preamplifier that has an on-board bias filtering circuit. This product was still under development from CAEN thus additional modifications were requested. The decay time constant was reduced to 50 µs (to avoid pileup events at high rates) and the gain was increased to 200 mV/MeV for two test units and 400 mV/MeV for two others. All the tested A1422 units have 200 pf input capacitance.

The performance of the higher gain version was very good, making it a strong candidate over the A250F/NF and the CR-110. A typical example of a pulse from the output of an A1422 preamplifier is shown in Figure 4.20, where the very good signal quality is evident. The major issue with this device is its size, but from a detailed
study it was found that 90 of these preamplifiers could be accommodated on the SAGE PCB. Their power dissipation is also higher than the other tested preamplifiers but this is unlikely to be an issue as they will be sufficiently cooled.

A summary of the results from the preamplifier tests can be seen in Tables 4.3 and 4.4. The first table shows the test results using an open $^{133}$Ba source and the second the outcome of the $^{241}$Am tests that were made to see how the preamplifiers perform at the low energy range.

From these results one can observe that the A1422 is consistently better than the CR-110 and A250F/NF. In fact the A250F/NF showed the poorest performance. The results also show that the TNT2 cards worked better than the Lyrtech, but this is not unexpected as the development of the TNT2 cards was almost complete at the time of the tests whereas the Lyrtech firmware was still under development, the
main outstanding issue being the non finalised constant fraction discriminator (CFD) algorithm. The A1422 is also the only one of the three preamplifiers that gave good results with the Lyrtech in its unfinished state. After consideration of all the above mentioned factors the CAEN A1422 with gain 400 mV/MeV and decay time constant of 50 $\mu$s was chosen as the preamplifier to be used with the silicon detector of SAGE.
Figure 4.18: The Zener voltage regulator diodes, mounted on the A422 preamplifier, used to provide ±12 V from the ±24 V inputs.

Figure 4.19: Circuit diagram of the test board built for testing the A1422 preamplifier.
Figure 4.20: The output from an A1422(400) preamplifier on the oscilloscope, taken using cosmic radiation. The setup was in atmospheric conditions and double screened within two metal boxes.

Table 4.3: Preamplifier test results using analogue and digital electronics with a $^{133}\text{Ba}$ source. The full width at half maximum (FWHM) for the 320.03 keV electron peak is shown for each case.

<table>
<thead>
<tr>
<th>Preamplifier</th>
<th>A250F/NF</th>
<th>CR-110(61)</th>
<th>A1422\textsuperscript{a}</th>
</tr>
</thead>
<tbody>
<tr>
<td>Analogue [keV]</td>
<td>4.84</td>
<td>4.41</td>
<td>4.17</td>
</tr>
<tr>
<td>TNT2 [keV]</td>
<td>4.97</td>
<td>4.53</td>
<td>4.67</td>
</tr>
<tr>
<td>Lyrtech [keV]</td>
<td>7.93</td>
<td>7.16</td>
<td>5.44</td>
</tr>
</tbody>
</table>

\textsuperscript{a}TNT2 results taken with the 90 mV/MeV version and Lyrtech with the 400 mV/MeV.
Table 4.4: Preamplifier test results using analogue and digital electronics with an $^{241}$Am source. The FWHM for the 59.54-keV $\gamma$-ray peak is shown for each case.

<table>
<thead>
<tr>
<th>Preamplifier</th>
<th>A250F/NF [keV]</th>
<th>CR-110(61) [keV]</th>
<th>A1422(400) [keV]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Analogue</td>
<td>3.20</td>
<td>2.85</td>
<td>2.61</td>
</tr>
<tr>
<td>Lyrtech</td>
<td>7.73</td>
<td>Not available(^a)</td>
<td>3.03</td>
</tr>
</tbody>
</table>

\(^a\)The internal trigger of the Lyrtech VHS-ADC was not finalised at the time of these tests and the software could not trigger correctly making the measurements with this low gain preamplifier impossible at low energies.
4.3 Gain and offset Cards

To be able to use the full dynamic range of ±1.1 V of the Lyrtech VHS-ADC, gain and offset adjustments are needed for both the silicon and germanium signals. Different options were investigated and it was decided that a Gain and Offset card (GO-card) should be used.

The two different GO-cards tested were the SmartPET GO-card (SGO-card) and the Liverpool GO-card (LGO-card). These cards were tested with signals from germanium and silicon detectors and a pulse generator.

The SGO-card is a NIM module developed and built by the Rutherford Appleton Laboratory (RAL). It is remotely controlled via Ethernet using a control interface implemented in the Multi Instance Data Acquisition System (MIDAS) [Pu]. Each module consists of 32 individually controlled channels, but the option of reducing them to 16 per card exists, in order to suppress electronic noise.

The SGO-card is a non-inverting unit that allows the buffering of detector signal, the removal of input offset, and the adjustment of gain and offset on the output signal. From the two offset level adjustments, one precedes (input bias) and one follows (output offset) the gain stage and a combination of the two allows very precise offset adjustment. The card has an input impedance of 50 Ω, thus if a preamplifier signal is to be directly connected to it, the preamplifier needs to have 50 Ω output impedance. The bandwidth of the SGO-card is independent of gain and limited to about 26 MHz.

A gain calibration of this card was performed using a 60 mVpp tail pulse, with 200 ns rise time and 1 ms fall time. The outcome of the calibration shown in Figure 4.21 indicates the mainly logarithmic character of the gain stage. Hexadecimal values are used on the gain axis as the SGO-card control interface requires input in hexadecimal values.

The LGO-card (a prototype unit at the time of the tests mentioned here) is designed by the University of Liverpool. It consists of a mother board, accommodating the connector sockets, and daughter boards with the gain and offset circuits. It is
Figure 4.21: Gain calibration of SGO-card using 60 mVpp tail pulse, indicated by the red dashed line. Both decimal and hexadecimal values are shown on the gain axis.

an inverting unit with all the channels having fixed gain but adjustable offset via a potentiometer. The gain of each channel can be modified by changing the gain adjustment components, or alternatively daughter boards with different gains can be built, a more preferable option. The input impedance of the card is 1 kΩ allowing the direct connection of signals from sources that are unable to drive 50 Ω loads e.g. the A250F/NF preamplifier. The bandwidth of the LGO-card is not dependent on gain over the gain ranges of interest and is limited to about 20 MHz.

The finalised version consists of two different types of motherboards. One with SMA connectors, for the silicon detector, that can accommodate three 4-channel daughter boards, and one with MMCX (micro-miniature coaxial) connectors, for the
germanium detectors, in which four 4-channel daughter boards can be connected. Up to four mother boards (48 channels in the SMA version and 64 channels in the MMCX one) can be placed in an individual, purpose-built box containing all the electronics needed (including power supplies), thus a NIM crate is not required.

Both the GO-cards were thoroughly tested mainly using a p-type germanium detector with a \(^{133}\)Ba source. The main \(^{133}\)Ba γ-ray energies are 53.2 keV, 79.6 keV, 81.0 keV, 160.6 keV, 223.2 keV, 276.3 keV, 302.8 keV, 356.0 keV and 383.8 keV [Tr90]. The detector signal was fed through the GO-card to a spec-amp and then to an MCA. Analogue electronics were used in most of the cases as the Lyrtech digital electronics were still under development. The gain on the spec-amp was adjusted so that the 356.0 keV γ ray of \(^{133}\)Ba was always at channel 7000 on the MCA. FWHM values were measured for the 53.16 keV and 356.01 keV γ rays.

The next set of tests was made using a tail pulse, obtained from a pulse generator. The tail pulse amplitude was adjusted so that in each case the resulting peak on the MCA was positioned at approximately the same position as the 53.16 keV and the 356.01 keV peaks from the germanium tests using the same spec-amp settings. This ensured that the MCA and the spec-amp settings did not affect the test results.

To check how the resolution depends on the gain, the offset of the buffer cards was set to 0 V. For the SGO-card both the offset settings were adjusted to setpoint 0x8000 (the middle of the range) which gave the required 0 V offset. The gain was adjusted using a tail pulse and comparing the output with that of the relevant LGO-card channel. The cards were considered to have the same gain when their output pulses had the same amplitude when measured on the oscilloscope. To make the two cards directly comparable the input signal of the LGO-card was 50 Ω terminated, but measurements were also made using 1 kΩ input impedance. A reference measurement was made without using a GO-card, i.e. the germanium detector or the pulse generator signals were directly connected to the spec-amp. The input of the spec-amp was 50 Ω terminated, to match the input impedance of the Lyrtech system.

A summary of the results of the tests is shown in Tables 4.5 and 4.6 and Fig-
ures 4.22 and 4.23. The LGO-card performs better than the SGO-card independent of energy, throughout the examined gain range. In Figure 4.24 a comparison of the 356.01 keV γ ray from the reference measurement and the SGO-card x8, LGO-card x8 50 Ω and 1 kΩ measurements is made.

It should be noted that for the same gain on the LGO-card with 50 Ω input impedance the input signal is attenuated to approximately half of that with 1 kΩ impedance. To compensate for this the gain on the spec-amp was increased, introducing additional noise in the system. This makes the LGO-card x1 1 kΩ measurement directly comparable with the LGO-card x2 50 Ω rather than the LGO-card x1 50 Ω and so on.

It was also investigated whether for a set gain the resolution achieved is offset dependent. Both GO-cards were tested at gain factors of 1 and 8 (LGO-card 1 kΩ input impedance). The germanium detector with a $^{133}$Ba source was used and the FWHM for the 356.01 keV γ ray was measured. The gain settings on the spec-amp were kept constant for each module.

For the LGO-card it was found that the FWHM is constant within 2% for the x1 gain and 3.5% for x8 gain throughout the offset range. For the SGO-card it varied up to 20% for both gains. Both GO-cards have an offset range of ±2 V for gain greater than x1. When the SGO-card is studied in the attenuation region the offset range reduces gradually down to ±1 V. If the LGO-card is used as an attenuator then the offset range can be kept at ±2 V if required.

It should be noted that the two offset stages of the SGO-card cannot be individually monitored. A combination of the two stages gives the desired offset in each case, hence the same offset can be achieved using different settings. To further investigate this, more thorough testing was made by using the SGO-card at x4 gain and setting the offset at -800 mV using different offset settings.

The output signal of the SGO-card was fed to a spec-amp adjusted so that the 356.01 keV γ ray was positioned at channel 7000 of the MCA; when both SGO-card offset settings were adjusted at the centre of their range (0x8000). From Table 4.7
Table 4.5: Performance of the GO-cards for a 53.16 keV $\gamma$ ray compared to a reference measurement in which no GO-card was used. Due to the good peak-to-background ratio the error on the FWHM calculation is not greater than 5% in any case.

<table>
<thead>
<tr>
<th>GO-card</th>
<th>Germanium detector</th>
<th>Pulse generator</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>FWHM [keV]</td>
<td>FWHM [ch]</td>
<td>$\approx$ FWHM [keV]</td>
</tr>
<tr>
<td>Reference</td>
<td>0.73</td>
<td>14.30</td>
<td>2.23</td>
</tr>
<tr>
<td>SGO-card x1</td>
<td>3.28</td>
<td>64.51</td>
<td>3.38</td>
</tr>
<tr>
<td>LGO-card x1 50 $\Omega$</td>
<td>1.39</td>
<td>27.33</td>
<td>3.57</td>
</tr>
<tr>
<td>LGO-card x1 1 k$\Omega$</td>
<td>0.76</td>
<td>14.92</td>
<td>1.10</td>
</tr>
<tr>
<td>SGO-card x2</td>
<td>2.20</td>
<td>43.28</td>
<td>2.63</td>
</tr>
<tr>
<td>LGO-card x2 50 $\Omega$</td>
<td>0.91</td>
<td>17.80</td>
<td>1.36</td>
</tr>
<tr>
<td>LGO-card x2 1 k$\Omega$</td>
<td>0.68</td>
<td>13.33</td>
<td>0.43</td>
</tr>
<tr>
<td>SGO-card x4</td>
<td>2.06</td>
<td>40.56</td>
<td>2.37</td>
</tr>
<tr>
<td>LGO-card x4 50 $\Omega$</td>
<td>0.92</td>
<td>18.17</td>
<td>0.74</td>
</tr>
<tr>
<td>LGO-card x4 1 k$\Omega$</td>
<td>0.71</td>
<td>13.99</td>
<td>0.95</td>
</tr>
<tr>
<td>SGO-card x8</td>
<td>2.33</td>
<td>45.79</td>
<td>2.45</td>
</tr>
<tr>
<td>LGO-card x8 50 $\Omega$</td>
<td>0.65</td>
<td>12.77</td>
<td>0.62</td>
</tr>
<tr>
<td>LGO-card x8 1 k$\Omega$</td>
<td>0.70</td>
<td>13.72</td>
<td>0.59</td>
</tr>
</tbody>
</table>

can be seen that the resolution varies a lot with the offset. The output offset seems to introduce most of the noise. A higher offset level, obtained by keeping the output offset at 0x8000 and increasing the input bias, gave a FWHM of around 2.6 keV in all cases.

Noise is introduced to the system when the offset levels are set further away from the middle of the range. To avoid this and to try to make the overall performance of the SGO-card better, filtering capacitors can be implemented at both offset stages.

To test how the GO-cards work with digital electronics an additional test was
Table 4.6: Performance of the GO-cards for a 356.01 keV $\gamma$ ray compared to a reference measurement taken without a GO-card. Due to the good peak-to-background ratio the error on the FWHM calculation is not greater than 5% in any case.

<table>
<thead>
<tr>
<th>GO-card</th>
<th>Germanium detector</th>
<th>Pulse generator</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>FWHM [keV]</td>
<td>FWHM [ch]</td>
</tr>
<tr>
<td>Reference</td>
<td>1.03</td>
<td>20.18</td>
</tr>
<tr>
<td>SGO-card x1</td>
<td>3.59</td>
<td>70.60</td>
</tr>
<tr>
<td>LGO-card x1 50 $\Omega$</td>
<td>1.57</td>
<td>30.84</td>
</tr>
<tr>
<td>LGO-card x1 1 k$\Omega$</td>
<td>1.06</td>
<td>20.80</td>
</tr>
<tr>
<td>SGO-card x2</td>
<td>2.82</td>
<td>55.56</td>
</tr>
<tr>
<td>LGO-card x2 50 $\Omega$</td>
<td>1.16</td>
<td>22.79</td>
</tr>
<tr>
<td>LGO-card x2 1 k$\Omega$</td>
<td>1.01</td>
<td>19.90</td>
</tr>
<tr>
<td>SGO-card x4</td>
<td>2.67</td>
<td>52.60</td>
</tr>
<tr>
<td>LGO-card x4 50 $\Omega$</td>
<td>1.15</td>
<td>22.69</td>
</tr>
<tr>
<td>LGO-card x4 1 k$\Omega$</td>
<td>1.02</td>
<td>20.00</td>
</tr>
<tr>
<td>SGO-card x8</td>
<td>2.50</td>
<td>49.14</td>
</tr>
<tr>
<td>LGO-card x8 50 $\Omega$</td>
<td>0.99</td>
<td>19.51</td>
</tr>
<tr>
<td>LGO-card x8 1 k$\Omega$</td>
<td>1.00</td>
<td>19.69</td>
</tr>
</tbody>
</table>

made with a germanium detector and $^{60}$Co source (observing the 1332 keV $\gamma$-ray) using the LGO-card at x8 gain. The results from this test are shown in Table 4.8. The measurements made with the LGO-card at x8 gain where consistently better than the reference measurements.

Another issue of importance is whether the pulse characteristics change because of the GO-cards. This was tested mainly using square or tail pulses from a pulse generator and also with a germanium detector. The results of this study are presented in Table 4.9. The fast fall time of a square pulse is affected by both the GO-cards
Figure 4.22: The FWHM of the 53.16 keV $\gamma$ ray for the measurements employing the different GO-cards with a germanium detector. The LGO-card outperforms the SGO-card throughout the examined gain range and performs equally well or even better than the reference. The errors are smaller than the data points in most cases so they were not included in the graph.

but the longer fall times of tail pulses remain mainly unaffected. The rise times of the output signals increase slightly more for the LGO-card than for the SGO-card.

Precise rise times were measured with the square pulse. The rise times of the germanium detector are approximations because the signal quality was not optimum due to signal reflections.

The overall results of the tests showed that the LGO-card outperforms the SGO-card throughout the gain range over which they were tested, so it was selected to be
Figure 4.23: Comparison of the FWHM of the 356.01 keV $\gamma$ ray for the measurements employing a germanium detector with different GO-cards. The LGO-card outperforms the SGO-card throughout the examined gain range is equally good as the reference. The errors are smaller than the data points in most cases so they were not included in the graph.

used in SAGE. For that purpose an adequate number of GO-boxes was build that can fully accommodate all the silicon and germanium channels of SAGE. Because these cards perform so well other groups will also use them for gain and offset adjustments or signal buffering.
Figure 4.24: A comparison of the 356.01 keV $\gamma$ ray from the reference measurement and the SGO-card x8, LGO-card x8 50 $\Omega$ and 1 k$\Omega$ measurements.
Table 4.7: Resolution with respect to offset settings for the SGO-card.

<table>
<thead>
<tr>
<th>Input bias</th>
<th>Output offset</th>
<th>Peak position</th>
<th>FWHM</th>
</tr>
</thead>
<tbody>
<tr>
<td>0x</td>
<td>0x</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9d00</td>
<td>0000</td>
<td>6946.39</td>
<td>3.39</td>
</tr>
<tr>
<td>8fff</td>
<td>8000</td>
<td>6984.97</td>
<td>2.65</td>
</tr>
<tr>
<td>8d00</td>
<td>a000</td>
<td>6990.82</td>
<td>3.42</td>
</tr>
<tr>
<td>8500</td>
<td>f000</td>
<td>6999.43</td>
<td>3.33</td>
</tr>
<tr>
<td>8390</td>
<td>ffff</td>
<td>7000.05</td>
<td>3.08</td>
</tr>
</tbody>
</table>

Table 4.8: Results from tests using the Lyrtech VHS-ADC in combination with the LGO-card x8 and a $^{60}$Co source.

<table>
<thead>
<tr>
<th>Setup</th>
<th>ADC</th>
<th>Energy [keV]</th>
<th>FWHM [keV]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ge</td>
<td>Lyrtech</td>
<td>1332</td>
<td>3.82</td>
</tr>
<tr>
<td>Ge + LGO-card x8</td>
<td>Lyrtech</td>
<td>1332</td>
<td>2.43</td>
</tr>
<tr>
<td>Ge</td>
<td>MCA</td>
<td>1332</td>
<td>2.29</td>
</tr>
<tr>
<td>Ge + LGO-card x8</td>
<td>MCA</td>
<td>1332</td>
<td>2.20</td>
</tr>
</tbody>
</table>
Table 4.9: Effect of GO-card on pulse characteristics. A comparison is made with different types of pulses studied either straight from the source or through one of the GO-cards. The tabulated values for the pulse generator signals are accurate within 1%.

<table>
<thead>
<tr>
<th></th>
<th>( t_{\text{rise}} )</th>
<th>( t_{\text{fall}} )</th>
<th>( t_{\text{rise}} )</th>
<th>( t_{\text{fall}} )</th>
<th>( t_{\text{rise}} )</th>
<th>( t_{\text{fall}} )</th>
</tr>
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<tbody>
<tr>
<td><strong>Square pulse</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Directly from the</td>
<td>4.73 ns</td>
<td>4.59 ns</td>
<td>201.7 ns</td>
<td>358.0 ( \mu s )</td>
<td>300.0 ( \pm 50 ) ns</td>
<td>130.0 ( \pm 10 ) ( \mu s )</td>
</tr>
<tr>
<td>pulse generator</td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Through the</td>
<td>12.47 ns</td>
<td>12.17 ns</td>
<td>202.4 ns</td>
<td>368.0 ( \mu s )</td>
<td>330.0 ( \pm 50 ) ns</td>
<td>150.0 ( \pm 10 ) ( \mu s )</td>
</tr>
<tr>
<td>SGO-card x4</td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Through the</td>
<td>17.20 ns</td>
<td>17.10 ns</td>
<td>203.7 ns</td>
<td>351.0 ( \mu s )</td>
<td>350.0 ( \pm 50 ) ns</td>
<td>120.0 ( \pm 10 ) ( \mu s )</td>
</tr>
<tr>
<td>LGO-card x4</td>
<td></td>
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<tr>
<td>Tail pulse</td>
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<tr>
<td>Directly from the</td>
<td>4.73 ns</td>
<td>4.59 ns</td>
<td>201.7 ns</td>
<td>358.0 ( \mu s )</td>
<td>300.0 ( \pm 50 ) ns</td>
<td>130.0 ( \pm 10 ) ( \mu s )</td>
</tr>
<tr>
<td>pulse generator</td>
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</tr>
<tr>
<td>Through the</td>
<td>12.47 ns</td>
<td>12.17 ns</td>
<td>202.4 ns</td>
<td>368.0 ( \mu s )</td>
<td>330.0 ( \pm 50 ) ns</td>
<td>150.0 ( \pm 10 ) ( \mu s )</td>
</tr>
<tr>
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<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Through the</td>
<td>17.20 ns</td>
<td>17.10 ns</td>
<td>203.7 ns</td>
<td>351.0 ( \mu s )</td>
<td>350.0 ( \pm 50 ) ns</td>
<td>120.0 ( \pm 10 ) ( \mu s )</td>
</tr>
<tr>
<td>LGO-card x8</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Germanium detector</td>
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<td>Directly</td>
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<td>Directly</td>
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<td></td>
<td>300.0 ( \pm 50 ) ns</td>
<td>300.0 ( \pm 50 ) ns</td>
<td>330.0 ( \pm 50 ) ns</td>
<td>330.0 ( \pm 50 ) ns</td>
<td>350.0 ( \pm 50 ) ns</td>
<td>350.0 ( \pm 50 ) ns</td>
</tr>
</tbody>
</table>
4.4 Signal chain

In SAGE there are 196 outputs from individual detectors/detector segments, 90 from the silicon and 106 from the germanium detectors. All of these channels are connected to a digital data acquisition system employing the Lyrtech 16-channel VHS-ADC cards.

The signal chain from the preamplifier output of the detectors to the Lyrtech VHS-ADC is shown in Figure 4.25. In this figure output for the germanium detectors is considered to be the preamplifier output and for the silicon detectors the output of the D-type to SMA adaptor boxes on the detector chamber.

![Figure 4.25: Schematic diagram of the signal chain from the output of the detectors to the input of the Lyrtech VHS-ADCs.](image)

The germanium detectors have BNC outputs that are connected to a patch panel on the JUROGAM II support frame using short double-screened RG223 cables. Other connections, like bias shutdown, bias voltage and temperature monitoring are made through the same patch panel.

From this patch panel longer super-screened MM11/50 cables take the signals to a second patch panel inside the air-conditioned electronics cabinets. The GO-boxes for the germanium detectors are positioned inside these cabinets and the connection from the last patch panel to them is made through short single-screened RG174 cables. The same type of cables is used to connect the output of the GO-boxes to the input of the Lyrtech VHS-ADCs.
From the silicon detector’s preamplifier outputs the signal is taken to the feed-through flange using 26-way ribbon cables (one of the 26 cables from each ribbon cable is not used). The 25-way D-type connectors on the flange are converted to SMA connectors through the use of custom made adaptor boxes. The SMA outputs are connected to the GO-boxes through short double-screened RD316 cables, meaning the GO-boxes are not located inside the electronics cabinets. Longer cables could be used but this would mean that the preamplifiers might be affected by the 50Ω impedance of the long cable at their output. Their performance is not influenced by the 1kΩ input impedance of the GO-cards. The longer cable would require more output current to charge up its greater capacitance than a shorter one, putting a higher load on the output of the preamplifier. This would cause increased heating and affect performance and stability as some amplifiers do not perform well with large capacitive loads on their output. For these reasons the GO-boxes are preferred for driving the longer line.

From the GO-boxes, RG223 cables take the signal to a patch panel inside the electronics cabinets. Short RG174 cables connect the patch panel to the Lyrtech VHS-ADC.

Table 4.10 lists the technical specifications of the cables used in the signal chain. The block diagram in Figure 4.26 shows the structure of the data acquisition system after the signals enter the Lyrtech VHS-ADCs.

The triggerless Total Data Readout (TDR) method [La01] is used in SAGE. In this system no hardware trigger is applied to start the data collection but all the channels run independently and are associated in software to reconstruct the events. This virtually eliminates the dead time issues arising when a common hardware trigger is used and when wide time gates are applied at the focal plane electronics.

Sixteen detector signals can be connected to each Lyrtech VHS-ADC. All the signals are timestamped using a universal clock produced by a Metronome (not shown in the figure) connected to the total data readout interface card (TDRi card) and are tagged with an address to identify which ADC channel they originate from and thus
Table 4.10: Technical specifications of cables.

<table>
<thead>
<tr>
<th></th>
<th>RG174</th>
<th>RG223</th>
<th>RD316</th>
<th>MM11/50</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inner conductor</td>
<td>Copper clad steel</td>
<td>Silver plated copper</td>
<td>Silver plated copper</td>
<td>Silver plated copper</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>copper clad steel</td>
<td></td>
</tr>
<tr>
<td>Dielectric</td>
<td>PE</td>
<td>PE</td>
<td>PTFE</td>
<td>PE</td>
</tr>
<tr>
<td>Outer conductor 1</td>
<td>Tinned copper braid</td>
<td>Silver plated copper braid</td>
<td>Silver plated copper braid</td>
<td>Silver plated copper braid</td>
</tr>
<tr>
<td>Outer conductor 2</td>
<td>None</td>
<td>Silver plated copper braid</td>
<td>Silver plated copper braid</td>
<td>Silver plated copper braid</td>
</tr>
<tr>
<td>Shielding</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>mu-metal</td>
</tr>
<tr>
<td>Jacket</td>
<td>PVC</td>
<td>PVC</td>
<td>FEP</td>
<td>PVC</td>
</tr>
</tbody>
</table>
which associated detector signal they represent.

Up to seven ADC cards are connected to a CPCI CPU (Compact Peripheral Component Interconnect Central Processing Unit) which is used for setup and control of the ADCs and readout of the parameters. Two CPCI crates are used to fully accommodate the ADC cards needed for SAGE. One TDRi card is responsible for providing the initial clock signal and a synchronisation pulse to all the ADC cards of one crate. The synchronisation pulse is used to ensure that the individual clock of each ADC card is synchronised with the rest. Each ADC card has an external trigger input that in the case of SAGE is used for the sync input from the TDRi card since a separate software trigger is responsible for triggering each channel.

The timestamped parameters from all the individual ADC channels are sent from the CPCI CPUs to the Merge. Here the timestamped data words are merged into

Figure 4.26: Block diagram of the digital electronics used in SAGE.
chronological order. The data are then filtered using a software trigger to reduce their size so they can be saved on disk or passed through to the on-line sort code. From the Merge the parameters can be optionally read into the Event Builder where the events are reconstructed in real time using temporal and spatial correlations defined by the physics of the experiment.

The software trigger is applied either to the focal plane or to the prompt detectors. In the first case whenever any data are detected in any of the focal plane detectors then any data that are present within a certain time window are recorded. In the latter case the trigger is placed on the multiplicity of a prompt event. For example it can be required that only fold-two or fold-three events are recorded. The two different triggers can be used in conjunction.
Chapter 5

Summary and Conclusion

The SAGE spectrometer was designed and built by a collaboration from the Universities of Liverpool in the UK and Jyväskylä in Finland and the STFC Daresbury Laboratory. SAGE allows cross-coincidence measurements of $\gamma$ rays and internal conversion electrons by combining a germanium detector array with a silicon detector and an electron transport system. The spectrometer is coupled with the RITU gas-filled recoil separator and the GREAT focal-plane spectrometer for recoil-decay tagging studies.

The individual parts of the setup, such as the solenoid coils, high-voltage barrier and detector electronics, are optimised for electron transmission efficiency, recoil acceptance in RITU and spectral response. The spectrometer employs fully digital front-end electronics and high-gain hybrid preamplifiers.

SAGE was successfully commissioned in the University of Jyväskylä and electron-$\gamma$ cross-coincidences were observed. Figure 5.1 shows SAGE during the final preparations before getting beam on target for the first time. In Figure 5.2 the collaborators right after the first electron spectrum was observed.
Figure 5.1: The SAGE spectrometer during the final preparations before being fully commissioned.

Figure 5.2: After the first electron spectrum was observed.
One reaction tried during the commissioning was $^{168}$Er$(^{16}$O,4$n)^{180}$Os. Osmium has electron binding energies of $B_K = 73.87$ keV, $B_{L_1} = 12.97$ keV, $B_{L_{II}} = 12.38$ keV and $B_{L_{III}} = 10.87$ keV. The decay scheme of the lower transitions of the ground-state band in $^{180}$Os is shown in Figure 5.3. The energies are a weighted average of the values reported in [Li99] using a $^{150}$Nd$(^{36}$S,6$n\gamma)$ reaction, [Dr82, Li88] using $^{166}$Er$(^{18}$O,4$n\gamma)$ and [Dr82, Dr90] where the reaction $^{168}$Er$(^{16}$O,4$n\gamma)$ was used. The ICCs for the E2 transitions under study in osmium are presented in Table 5.1.

Figure 5.3 shows the ICE spectrum obtained when gating on the 386.4 keV and 510.1 keV $\gamma$ rays, as shown in the inset. Gating on the 132.1 keV L electrons the $\gamma$-ray spectrum shown in Figure 5.4 is observed. No Compton suppression or recoil gates were used in these spectra.

Table 5.1: ICCs of the ground-state band transitions of $^{180}$Os [Ki08].

<table>
<thead>
<tr>
<th>$E_\gamma$ [keV]</th>
<th>$\alpha$</th>
<th>$\alpha_K$</th>
<th>$\alpha_{L_1}$</th>
<th>$\alpha_{L_{II}}$</th>
<th>$\alpha_{L_{III}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>132.1</td>
<td>1.468</td>
<td>0.473</td>
<td>0.0527</td>
<td>0.390</td>
<td>0.308</td>
</tr>
<tr>
<td>276.5</td>
<td>0.1175</td>
<td>0.0732</td>
<td>0.00918</td>
<td>0.01552</td>
<td>0.00896</td>
</tr>
<tr>
<td>386.4</td>
<td>0.0445</td>
<td>0.0315</td>
<td>0.00414</td>
<td>0.00391</td>
<td>0.00192</td>
</tr>
<tr>
<td>462.2</td>
<td>0.0278</td>
<td>0.0205</td>
<td>0.00275</td>
<td>0.00192</td>
<td>0.000866</td>
</tr>
<tr>
<td>510.1</td>
<td>0.0217</td>
<td>0.01637</td>
<td>0.00221</td>
<td>0.001306</td>
<td>0.000564</td>
</tr>
<tr>
<td>541.2</td>
<td>0.0188</td>
<td>0.01435</td>
<td>0.00194</td>
<td>0.001042</td>
<td>0.000438</td>
</tr>
<tr>
<td>566.1</td>
<td>0.01690</td>
<td>0.01299</td>
<td>0.001758</td>
<td>0.000878</td>
<td>0.000362</td>
</tr>
</tbody>
</table>

A different reaction studied during the commissioning was $^{144}$Sm$(^{40}$Ar,4$n)^{180}$Hg. For $^{180}$Hg the lower transitions of the ground-state band are shown in Figure 5.5 [Ko00]. The electron binding energies of mercury are $B_K = 83.10$ keV, $B_{L_1} = 14.84$ keV, $B_{L_{II}} = 14.20$ keV and $B_{L_{III}} = 12.28$ keV, and the ICCs for the ground-state band transitions are listed in Table 5.2.

An ICE recoil-gated spectrum obtained when gating on the 272.4 keV and 434.3 keV
Figure 5.3: An ice spectrum of $^{180}$Os obtained when gating on the 386.4 keV and the 510.1 keV $\gamma$ rays of the ground-state band as shown in the inset. On the right-hand side a partial level scheme of $^{180}$Os is displayed with the gated transitions highlighted.

$\gamma$ rays, is shown in Figure 5.5. Gating on the 272.4 keV K electrons gives the $\gamma$-ray recoil-gated spectrum shown in Figure 5.6.

Table 5.2: ICCs of the ground-state band transitions of $^{180}$Hg [Ki08].

<table>
<thead>
<tr>
<th>$E_\gamma$ [keV]</th>
<th>$\alpha$</th>
<th>$\alpha_K$</th>
<th>$\alpha_{L_i}$</th>
<th>$\alpha_{L_{i+1}}$</th>
<th>$\alpha_{L_{i+2}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>434.3</td>
<td>0.0382</td>
<td>0.0266</td>
<td>0.00375</td>
<td>0.00358</td>
<td>0.001512</td>
</tr>
<tr>
<td>272.4</td>
<td>0.1426</td>
<td>0.0797</td>
<td>0.01075</td>
<td>0.0238</td>
<td>0.01276</td>
</tr>
<tr>
<td>326.0</td>
<td>0.0835</td>
<td>0.0517</td>
<td>0.00712</td>
<td>0.01132</td>
<td>0.00554</td>
</tr>
<tr>
<td>404.5</td>
<td>0.0460</td>
<td>0.0312</td>
<td>0.00438</td>
<td>0.00475</td>
<td>0.00208</td>
</tr>
</tbody>
</table>

When studying electron-$\gamma$ cross coincidences in both the reactions described here,
applying different electron or $\gamma$-ray gates yields spectra showing the desired behaviour. That is by gating on one of the ground state band electrons, in the resulting $\gamma$-ray spectrum the corresponding $\gamma$-ray peak is suppressed whereas the other ground state band decays are clearly seen. Additionally transitions belonging to side bands feeding into the ground-state band can be seen. Equivalently gating on a ground-state $\gamma$ ray gives ICE spectra where the corresponding electron peaks are suppressed but other ground-state band decays are present.

For example in the spectrum in Figure 5.4 where a gate was placed on the 132 keV L electrons, the 132 keV peak is not present in the $\gamma$-ray spectrum but the other ground state-band transitions are clearly present. The same behaviour is observed in the spectrum of Figure 5.6 when gating on the 272 keV K electrons. The same is true for electron spectra obtained when gating on $\gamma$ rays.
Figure 5.5: The recoil-gated ice spectrum in $^{180}$Hg obtained when gating on the 272.4 keV and 434.3 keV $\gamma$ rays of the ground-state band. The gates are shown in the inset and the level scheme on the right.

When gates are placed on the background in the $\gamma$ or ice spectra the resulting spectrum has no apparent structure. This proves that the peaks in the presented spectra are from true cross-coincidences and not random events.

The spectra presented in Figures 5.3, 5.4, 5.5 and 5.6 are preliminary and cleaner spectra with increased statistics can be obtained from detailed offline analysis. It is important to note that the observed ground-state band decays agree well with the previously published values.

The purpose of these spectra is to highlight that the electron and $\gamma$-ray parts of the SAGE spectrometer work well both independently and in conjunction, without the one interfering with the other. Recoil gating using GREAT and RITU was successfully employed during the commissioning.
The outcome of the commissioning experiments is that the main requirements and design goals set for SAGE have been achieved. Checking the list presented in Section 3.2 one can see that the $\gamma$-ray detection efficiency and resolution have not been affected by the electron part, the low-energy electron background is successfully suppressed and the electron transmission efficiency agrees within error bars with the predicted values. The measured transmission efficiency is not presented in this thesis as not all the required measurements were performed by the time of publication.

On the data acquisition front the digital front-end electronics operate almost flawlessly allowing measurements at high rates with no energy deterioration. Rates of up to 45 kHz at the Phase I detectors were achieved without difficulty.

Even though the spectrometer works according to the design specifications there are still some electronics issues that need addressing. A course of action has been
decided and a series of tests and optimisations will be performed before the next SAGE experimental campaign.

A number of experiments have been approved related both with the superheavy nuclei project and the search for shape coexistence in lead and mercury nuclei. Already before submitting the final version of this thesis two SAGE experiments were performed to look for shape coexistence in radon and lead nuclei. No results are presented here from these experiments as the analysis is ongoing. A brief analysis of the data proves that SAGE works well within the design criteria and that by further improving the stability and resolution characteristics of the electron part some wonderful experimental results can be expected in the future.
Appendices
Appendix A

Electromagnetism basics related to the simulations

This section does not try to explain electromagnetism but rather to present the equations useful for simulating electric and magnetic fields from given charge or current distributions, respectively\(^1\).

Electromagnetism is based on Maxwell’s equations that in SI units are expressed as:

- **Coulomb’s Law**
  \[ \nabla \cdot \vec{D} = \rho_f, \tag{A.1} \]

- **Absence of free magnetic poles**
  \[ \nabla \cdot \vec{B} = 0, \tag{A.2} \]

- **Faraday’s Law**
  \[ \nabla \times \vec{E} = -\frac{\partial \vec{B}}{\partial t}, \tag{A.3} \]

- **Ampère’s Law with Maxwell’s correction**
  \[ \nabla \times \vec{H} = \vec{j}_f + \frac{\partial \vec{D}}{\partial t}. \tag{A.4} \]

\(^1\)Information from books [Gr90, Ja99, Gr99, Gr00, St01] were used in this appendix.
APPENDIX A. EM BASICS RELATED TO THE SIMULATIONS

Where \( D \) is the electric flux density or electric displacement, \( \rho_f \) is the free charge density, \( B \) is the magnetic flux density, \( E \) the electric field intensity, \( H \) the magnetic field intensity and \( j_f \) the current density.

In the case where static electric and magnetic fields are considered Maxwell’s equations are reduced to:

\[
\nabla \cdot \vec{D} = \rho_f, \tag{A.5}
\]
\[
\nabla \cdot \vec{B} = 0, \tag{A.6}
\]
\[
\nabla \times \vec{E} = 0, \tag{A.7}
\]
\[
\nabla \times \vec{H} = \vec{j}_f \tag{A.8}
\]

and the solution to a general static electromagnetic problem can be derived simply by solving the electrostatic and magnetostatic equations individually and combining the results.

The equation of continuity (Equation A.9) relates the total macroscopic charge and current densities at each point and it expresses the conservation of charge in differential form,

\[
\frac{\partial \rho}{\partial t} + \nabla \cdot \vec{j} = 0. \tag{A.9}
\]

When Maxwell’s equations are combined with the Lorentz force equation (Equation A.10) and Newton’s second law of motion the classical dynamics of interacting charged particles and EM fields can be completely described.

\[
\vec{F} = q(\vec{E} + \vec{v} \times \vec{B}), \tag{A.10}
\]

with \( q \) being the charge of a particle moving with velocity \( \vec{v} \).

Besides Maxwell’s equations other useful formulae can be used in the study of electromagnetic fields. For example, the magnetic field of a steady line current is given by the Biot-Savart Law,

\[
\vec{B}(\vec{r}_0) = \frac{\mu_0 I}{4\pi} \oint \frac{\vec{dl}' \times \vec{r}}{r^3}, \tag{A.11}
\]
where the integration is along the current path in the direction of the flow, $d\vec{l}'$ is an element of length along the path at $\vec{r}_c$ and $\vec{r}$ is the vector from the source to the point $\vec{r}_0$, where $\vec{r} = \vec{r}_0 - \vec{r}_c$, as shown in Figure A.1.

![Figure A.1: Geometry used to determine the magnetic field of a current loop with the Biot-Savart Law.](image)

The electric scalar potential of a given stationary charge distribution can be calculated using Poisson’s equation (Equation A.12) or if no charge is present in the area of interest this is reduced to Laplace’s equation (Equation A.13),

\[
\nabla^2 V = -\frac{\rho}{\epsilon}, \quad (A.12) \\
\n\nabla^2 V = 0. \quad (A.13)
\]

The constant $\epsilon$ in Equation A.12 is the dielectric constant of the material of interest. Using the dielectric constant the electric displacement $\vec{D}$ is related to the electric field $\vec{E}$:

\[
\vec{D} = \epsilon \vec{E}. \quad (A.14)
\]

In the case of magnetostatic problems, Equations A.6 and A.8 must be obeyed. To achieve this easily a magnetic vector potential $\vec{A}$ is defined as:

\[
\vec{B} = \nabla \times \vec{A}. \quad (A.15)
\]

As for the dielectric constant, the magnetic permeability, $\mu$, can be defined that combines the magnetic field flux density, $\vec{B}$, with the field’s intensity, $\vec{H}$:

\[
\vec{B} = \mu \vec{H}. \quad (A.16)
\]
From the above equations for the magnetic field an equation similar to Poisson’s equation is derived:

\[ \nabla \times \left( \frac{1}{\mu} \nabla \times \vec{A} \right) = \vec{j}, \]  

(A.17)

and if we assume \( \nabla \cdot \vec{A} = 0 \) then Equation A.17 reduces to:

\[ \nabla^2 \vec{A} = \mu \vec{j}. \]  

(A.18)

### A.1 Electron motion in electromagnetic fields

To show how electrons move inside an electromagnetic field, the simple example of an electron moving in a uniform, static, magnetic field \( \vec{B} \) will be used. The equations of motion in this case can be written as:

\[ \frac{d\vec{p}}{dt} = \frac{e}{c} \vec{v} \times \vec{B} \quad \text{and} \quad \frac{dE}{dt} = 0, \]  

(A.19)

where \( \vec{v} \) is the particle’s velocity. Since the energy is constant in time, the magnitude of the velocity is constant as well.

Solving these equations shows that the particle moves in a circle of radius \( r \) perpendicular to the magnetic field. It also has a velocity component parallel to the field meaning that it will move in a helix along the field lines, as the circular and the linear motions combine.

The gyration radius \( r \) of the particle is given by:

\[ r = \frac{cp_{\perp}}{eB}, \]  

(A.20)

where \( p_{\perp} \) is the transverse momentum of the particle.

In the case where the particle moves in combined electric and magnetic fields then the motion is the same as described above with the difference that the electric field will add additional acceleration to the particle.

In the case of SAGE the electrons move along a helical path in the direction of the magnetic field lines. The less energetic ones are deflected by the high-voltage barrier.
and the more energetic are decelerated when approaching it and then accelerated with an equal amount when moving away from it.

A detailed analysis on the motion of charged particles in electric and/or magnetic fields is made in [Ja99].
Appendix B

The simulation packages

The Vector Fields OPERA 3D simulation environment [VF07] contains a series of analysis programs including “TOSCA magnetostatics” and “TOSCA electrostatics” that were used in the simulations made for SAGE. Electromagnetic simulations are provided by independently solving the electric and magnetic equations and combining the results.

A modeller is available in OPERA 3D for the construction of a model in three-dimensional space using geometric primitive volumes and Boolean operations. A model can also be extracted from a two-dimensional cross-section. The coils of SAGE were modelled using conductors, which are available for modelling in different shapes and sizes. Any symmetries present in the model can be exploited for simplifying the model and making the simulation faster.

The user can define properties for each type of material used in the model. If a magnetic field analysis is desired then a series of options for the permeability is available. For non-linear permeability a curve of magnetic flux density versus the magnetic field intensity (BH curve) can be assigned from the available library which allows the user to add new materials if needed. When analysing an electric field problem the conductivity and relative permittivity of each material can be defined.

Before proceeding to the analysis the model needs to be meshed. A two step procedure is followed where initially the surface is meshed and then the volume. The
user can define for the surface the maximum mesh element size, maximum angle between elements and maximum deviation from the surface, and for both surface and volume mesh the absolute tolerances.

When the model is fully defined and meshed the analysis database is prepared and solved. The results of the simulation are viewed in the “Post-Processor” where one can project a number of parameters on different surfaces or axes or output the values for analysis or representation using different software.

A similar procedure (building of model, meshing, defining border and material properties and setting up and solving the problem) is used in the Finite Element Method Magnetic (FEMM) [Me06] simulation package. FEMM is a suite of programs for solving low frequency electromagnetic problems on two-dimensional planar and axisymmetric domains. A problem can be solved quickly in two dimensions with FEMM and the results have been shown to agree with the ones from OPERA 3D.

For comparison Figure B.1 shows plots of the magnetic field on the solenoid axis for the same geometry using OPERA 3D and FEMM. In these simulations a 700 A current through the coils was used. The 3.2° angle between the beam and solenoid axes was incorporated only in the OPERA 3D simulation. This is because in FEMM calculations are axisymmetric, so the angle would produce conical coils instead of solenoids. This leads to a difference in the calculated fields between the target and detector positions. This effect disappears at the region where the two axes are collinear.

FEMM was used to quickly see effects of changes in coil geometry, whereas OPERA 3D was used to simulate almost finalised geometries or cases where no symmetries could be applied to extend the two dimensional space to three dimensions in FEMM.

For solving electrostatic problems both OPERA and FEMM solve Poisson’s equation (Equation A.12) in the frame of the user-defined conditions. Following this the electric field intensity and flux density are calculated.

For magnetostatic problems FEMM solves Equation A.17 and by differentiating $\vec{A}$; $\vec{B}$ and $\vec{H}$ are deduced. This process is more complicated in the case of OPERA. The program uses a combination of total (for regions were currents do not flow) and
Figure B.1: A comparison of the magnetic field on the solenoid axis calculated using OPERA 3D and FEMM.

reduced scalar potentials (in regions where currents flow i.e. conductors) to overcome any difficulties.

In the regions where currents flow, the total field intensity can be defined by summing the reduced ($H_m$) and conductor ($H_s$) field intensities, namely:

$$\vec{H}_m = -\nabla \phi$$  \hspace{1cm} (B.1)

and

$$\vec{H}_s = \int_{\Omega_J} \frac{\vec{J} \times \vec{R}}{\vec{R}^3} \, d\Omega_J.$$  \hspace{1cm} (B.2)

In the volumes where no currents are flowing a total field intensity is represented
using the total magnetic scalar potential as:

\[ \vec{H} = -\nabla \psi, \quad (B.3) \]

where the total magnetic scalar potential satisfies:

\[ \nabla \cdot \mu \nabla \psi = 0. \quad (B.4) \]
Appendix C

Technical specification of magnetic coils

The coils are made from winding glass/epoxy impregnated copper conductors. There is 1 mm of epoxy between two conductors (0.5 mm around each conductor) and 1.5 mm on the outside of each coil (0.5 mm from the external conductors plus 1 mm around the whole coil). Individual conductors have the geometry shown in Figure C.1 and specifications as in Table C.1. The diameter indicated by “d” in the drawing is the central hole used for water cooling.

Figure C.1: Cross-section of the solenoid coil conductors.

Figure C.2 shows the cross-section of the SAGE solenoid coils. In this figure the green squares demonstrate where one conductor crosses-over to the next loop. In the case of the main coil they denote how neighbouring conductors are coupled. The
Table C.1: Specifications of the solenoid coils and the individual conductors dimensions as shown in Figure C.1.

<table>
<thead>
<tr>
<th></th>
<th>Main coil</th>
<th>Upstream coil</th>
<th>Downstream coil</th>
</tr>
</thead>
<tbody>
<tr>
<td>A [mm]</td>
<td>9</td>
<td>9</td>
<td>8</td>
</tr>
<tr>
<td>B [mm]</td>
<td>9</td>
<td>9</td>
<td>8</td>
</tr>
<tr>
<td>d [mm]</td>
<td>Ø5</td>
<td>Ø6</td>
<td>Ø5</td>
</tr>
<tr>
<td>R [mm]</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Weight [kg]</td>
<td>79.4</td>
<td>39.5</td>
<td>33.1</td>
</tr>
<tr>
<td>Resistance at 20° C [Ω]</td>
<td>0.05</td>
<td>0.02</td>
<td>0.02</td>
</tr>
</tbody>
</table>

arrows show the individual cooling circuits. The effect of water cooling in the main and upstream coils is clearly visible in the infrared camera shot shown in Figure C.3.
Figure C.2: Cross-section of the magnetic coils of SAGE. Individual cooling circuits are indicated with continuous lines. Clockwise from top: Upstream, downstream and main coil. The arrows indicate where the cool water (blue) enters the cooling circuit and where the warm water (red) exits. The green squares show cross-overs or coupling of different conductors.
Figure C.3: Infrared camera picture showing the effect of water cooling on the main and upstream coils. 600 A current was used through the coils. Temperature scale is shown in the vertical bar on the right.
Appendix D

List of publications

1. Towards combining in-beam $\gamma$-ray and conversion electron spectroscopy

2. Spectroscopy of Very Heavy Elements

3. $\gamma$-Ray Spectroscopy at the Limits: First Observation of Rotational Bands in $^{255}$Lr
   S. Ketelhut, P.T. Greenlees, D. Ackermann, S. Antalic, E. Clment, I.G. Darby,

4. Bridging the nuclear structure gap between stable and super heavy nuclei

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