Multi-stage Micropattern Gaseous Detectors for the ALICE TPC Upgrade - Studying and Modelling Charge Transfer and Energy Resolution

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Abstract

With the upgrade of the LHC (Large Hadron Collider) at CERN, the interaction rate of ALICE (A Large Ion Collider Experiment) will be increased up to 50 kHz for Pb-Pb collisions. Thus the gated and rate-limited readout technology of the TPC (Time Projection Chamber) requires a complete redesign to allow for a continuous operation. Micropattern Gaseous Detectors (MPGD) are considered a promising solution to overcome the gating required for the existing Multiwire-Propotional Chambers (MWPC) technology. Several solutions like a multi-GEM (Gaseous Electron Multipliers) stack and a hybrid detector consisting of two GEM stages and a single Micromegas were under investigation. A solution with four GEMs has been adopted as baseline solution for the upgraded chambers since the operation of multi-GEM stages was more understood and studied at this time.

Within the scope of this work an alternative approach consisting of two GEM foils and a single Micromegas (MM) has been investigated in terms of the energy resolution, the ion backflow and the gain. The hybrid 2GEM-MM detector as well as the newly developed Slow Control to operate the setup are presented in detail. A systematic study of the recorded $^{55}$Fe energy spectra is a central part which finally leads to a dedicated fit model to obtain the energy resolution. A comparison yields that fitting a single Gaussian distribution to the photo peak overestimates the energy resolution by 1 % up to 2 % (difference of absolute values). The measurements are compared to the baseline solution of the ALICE TPC upgrade program as well as to a hybrid 2GEM-MM setup which has been studied at the Yale University. The hybrid 2GEM-MM detector clearly competes with the baseline solution of the ALICE TPC upgrade and the Yale measurements can be reproduced.

A major part of this work is the investigation of the charge transfer processes in GEM stacks, as these transfer efficiencies highly determine the energy resolution, the ion backflow and the gain. Within two-dimensional electrostatic calculations of electric fluxes, analytic expressions of the electron as well as of the ion transfer efficiencies can be derived as functions of the hole size, the pitch and the thickness of a GEM. The equations are compared to simulations, allowing to immediately calculate transfer efficiencies for arbitrary electrostatic configurations and GEM geometries. A big advantage is the short calculation time compared to the time-consuming simulations. The calculations lead to a profound and detailed understanding of the formation of the characteristic transfer efficiency curves.

The transfer efficiencies are used in order to derive models to calculate the energy resolution, the ion backflow and the gain of stacks with multiple GEM stages and for arbitrary electric field configurations. The model calculations allow for in-depth studies of the processes within multiple amplification stages and to understand the contributions of the individual stages to the measured quantities of the detector, i.e. energy resolution, ion backflow and gain. The models are compared to the measurements of the Bonn and the Yale hybrid 2GEM-MM detector as well as to the quadruple GEM stack of the ALICE TPC upgrade. Finally, the developed charge transfer models as well as the energy resolution model are implemented in the new ALICE O² framework which will be used with the ongoing upgrade of ALICE for the online as well as for the offline data acquisition.
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Introduction

1.1 Historic background

Scattering experiments are the key to access and understand the inner structure of matter. By "shooting" projectiles (e.g. electrons) on targets (e.g. nucleons like protons or neutrons) under well-defined conditions (momentum, energy, polarization, angle ...), information can be inferred about the inner structure and underlying interactions by investigating the outgoing reaction products (e.g. scattered particles or newly formed matter) and comparing them to theoretical predictions.

The inner atomic structure, for instance, was studied by Rutherford in 1911 when he investigated the elastic scattering of charged particles ($\alpha$ and $\beta$ particles) in the Coulomb potential of thin atomic layers (e.g. gold foils). Based on the angular distribution of the scattered particles, he developed a model of an atom with a tiny, massive and positively charged nucleus that is uniformly surrounded by a sphere of electrons [1]. It was discovered that also the nucleus has a substructure consisting of positively and neutrally charged nucleons, i.e. protons and the neutrons. In 1932, Chadwick proved the existence of the neutron by investigating the unknown radiation which is emitted once $\alpha$ particles irradiate a beryllium target [2].

Measurements in the 60s at the Stanford Linear Accelerator Center (SLAC) - which delivered electron energies up to 25 GeV - revealed that even the nucleons have a substructure. An important quantity for scattering experiments is the four-momentum transfer $Q$ which describes the transferred energy $\nu$ and momentum $p$ during a single scattering interaction. Within deep inelastic scattering the four momentum transfer $Q$ is much higher than the dimension $R$ of the nucleus, i.e $Q^2 \gg \hbar^2 / R^2$. This allows an interaction with the substructure of the nucleons and thus to resolve the inner geometries. With the mass $M$ of the target particles, the Bjorken scaling variable is defined as $x := Q^2 / 2M\nu$. In case of elastic scattering processes the Bjorken scaling variable is given by $x = 1$. For inelastic scattering processes $x$ is given by $0 < x < 1$. The cross section for inelastic scattering of electrons on nucleons can be written according to

$$\frac{d^2\sigma}{d\Omega dE} = \left( \frac{d\sigma}{d\Omega} \right)_{\text{Mott}} \left[ \frac{F_2(x, Q^2)}{\nu} + \frac{2F_1(x, Q^2)}{M c^2 \tan^2 \theta / 2} \right]$$

(1.1)

where the structure functions $F_1(x, Q^2), F_2(x, Q^2)$ and the Mott cross section (scattering of pointlike, spin 1/2 particles on pointlike and spinless targets) have been introduced [3]. The structure functions can be correlated to the magnetic current and electric charge-density distributions of the target and thus to its
Chapter 1 Introduction

(a) Structure function $F_2$ for the proton as a function of $x$ and different values for $Q^2$ [3].

(b) Ratio of the structure functions $F_1$ and $F_2$ fulfilling the Callan-Gross relation, i.e. $\approx 1$ [4].

Figure 1.1: Measurements of the structure function $F_2$ for the proton and the ratio of the structure functions $F_1$ and $F_2$ (crosscheck of the Callan-Gross relation).

Internal structure. For pointlike charge distributions, like for electrons, the structure function $F_2$ turns out to be independent of the momentum transfer $Q^2$, i.e. $F_2(x, Q^2) \rightarrow F_2(x)$ (also called Bjorken scaling). Measurements at SLAC showed that the structure function $F_2$ of protons is mostly unaffected by the four momentum transfer $Q$ (cf. Fig. 1.1(a)) which leads to the conclusion that protons and neutrons have a substructure consisting of pointlike particles.

For spin 1/2 particles, a relation can be found between the structure functions which is also known as the Callan-Gross relation [5]: $2xF_1(x) = F_2(x)$. This relation has been confirmed as shown in Fig. 1.1(b) which leads to the conclusion that protons and neutrons have a substructure consisting of pointlike particles.

Protons and neutrons consist of pointlike particles with spin 1/2. With the knowledge of the total spin of a nucleon it can be inferred that at least three or more constituents are needed. As the total electric charge of a nucleon is either $q = 0 \, e$ (neutron) or $q = 1 \, e$ (proton), fractional charges can be assumed. These pointlike particles are quarks. In total there are six kinds of quarks (up and down, charm and strange, top and bottom) which differ in their masses and charges. Together with the leptons (electron, muon, tau and the corresponding neutrinos) they can be arranged in three generations as shown in Fig. 1.3. Generally hadrons consisting of three quarks (i.e. qqq-states) are referred to as baryons. Quark-Antiquark states (q\overline{q}) are known as mesons. As example protons consist of two up and a single down quark (uud), neutrons of two down and a single up quark (udd). These quarks which determine the quantum numbers are also called valence quarks. In addition also virtual quark-antiquark pairs (sea quarks) exist which do not affect the quantum numbers as their contributions average out to zero. The remaining bosons are the intermediating exchange particles for the fundamental forces, i.e. the weak interaction (W and Z bosons), the strong interaction (gluons) and the electromagnetic interaction (photons). The latest breakthrough in the search of the Higgs boson at CERN was rewarded with the nobel prize in 2013 for Higgs [6] and Englert [7].

The underlying relativistic field theory of the strong interaction is referred to as quantum chromodynamics (QCD) and was first introduced in the early 1970s by Fritzsch et al. [8]. Like for the electromagnetic
interaction of electrons with photons (described by quantum electrodynamics (QED)), a force carrier is required for the strong interaction which are the gluons. An additional quantum number - the color charge - has been introduced in order to explain hadronic bound states consisting of quarks and gluons \[9\]. Particle production experiments in electron-positron annihilations confirm the existence of three color charges by comparing the cross sections for hadronic production \(\sigma(e^+e^- \rightarrow \text{hadrons})\) to muon pair production \(\sigma(e^+e^- \rightarrow \mu^+\mu^-)\) processes (see \[3\] for details). In total there are three color states and the corresponding anti-colors. In QCD the gluons themselves carry color charges which leads to a self-interaction between the gluons. This is a fundamental difference to QED where photons do not interact with each other as they carry no electric charge. The coupling constant of the strong interaction \(\alpha_s\) shows a large dependence on the momentum transfer \(Q\) as a consequence of the self-interaction of the gluons (cf. Fig. 1.2). The coupling constant \(\alpha_s\) decreases for increasing momentum transfers \(Q\) or - accordingly - smaller spatial distances, i.e. quarks behave like free or weakly bound in this regime. This behavior is also referred to as asymptotic freedom. At large distances or - equivalently - lower momentum transfers \(Q\), the strong interaction between two objects becomes stronger and the coupling constant increases. Due to the strong increase of the coupling constant for low values of \(Q\) (or larger distances), free quarks and gluons have never been observed as they always appear in bound and color-singlet hadronic states. This behavior is known as confinement.

It is assumed that a deconfined state of matter has existed up to a few milliseconds after the big bang just before hadronic matter will be formed. Accordingly if hadronic matter is sufficiently heated or compressed, the finite-sized hadrons start to overlap and the quarks as well as the gluons are capable of moving freely over large space-time distances, i.e. a deconfined phase only consisting of free quarks and gluons has been formed \[12\]. This phase of matter is also called Quark-Gluon Plasma (QGP).

Nowadays ultrarelativistic nucleus-nucleus collision experiments give access to investigate the phase diagram of strongly interacting matter and to study the QGP phase. Dedicated studies have been carried out at the Super Proton Synchrotron (SPS) at CERN and at the Relativistic Heavy Ion Collider (RHIC) at...
Brookhaven (see e.g. \cite{13} and \cite{14} for summarized results of the STAR\textsuperscript{1} and the PHENIX\textsuperscript{2} experiments). Understanding the QCD phase diagram and the transition from hadronic matter to the Quark-Gluon Plasma is a major exploratory focus of ALICE (A Large Ion Collider Experiment). A detailed introduction to ALICE and recent results from Run 1 and Run 2 will be presented in Sec. 3.

At sufficiently high temperatures and / or net baryon densities, a transition to the deconfined state can be expected at a critical temperature in a region close to $\alpha_s \sim 1$, i.e. a perturbative method to solve the QCD Lagrangian for further calculations (like done in QED) is not possible. Lattice QCD offers a way to solve the QCD equations by discretizing the Lagrangian on a four-dimensional Euclidean space-time lattice with fixed lattice spacings and thus volume elements $V$. Calculations yield that a transition from hadronic matter to QGP can be found at the critical temperature $T_c = (154 \pm 9)$ MeV \cite{15}. The calculated energy density $\epsilon = E/V$ is shown in Fig. 1.4 as a function of the temperature $T$: A change can be observed at the critical temperature $T_c$ (for vanishing net baryon density) which translates to an energy density of $\epsilon_c \approx (0.34 \pm 0.16) \text{ GeV fm}^{-3}$ \cite{16}. The solid lines indicate results obtained from hadron resonance gas (HRG) model calculations \cite{17}: A model prior to QCD which describes hadronic matter as a non-interacting gas based on experimentally known resonances. The HRG model calculations are in a good agreement to the QCD calculations for lower temperatures up to the transition region from hadronic matter to QGP. The obtained thermodynamic observables of the HRG model calculations (like pressure) are at the lower error band of the QCD calculations which might be a hint that even more resonances could exist which had not been experimentally observed so far (see \cite{16} for details).

\footnote{1 Solenoidal Tracker at RHIC}
\footnote{2 Pioneering High Energy Nuclear Interactions Experiment}
1.1 Historic background

Figure 1.4: Calculated pressure $p$, energy density $\epsilon$ and entropy density $s$ in QCD as a function of the temperature (plot from [16]). Solid lines indicate results obtained from hadron resonance gas (HRG) model calculations [17].

Fig. 1.5 shows a simplified schematic view of the phase diagram and the different phases for QCD matter as a function of the temperature $T$ and the baryochemical potential $\mu_b$ (a measure of the net baryon density which correlates the number of baryons and anti-baryons in a given volume). For low baryon densities $\mu_b \approx 0$ and at high temperatures, the transition from hadronic matter to QGP is expected to happen via a smooth crossover (dashed line) at the critical temperature $T_c$. The solid line indicates a first-order phase transition for moderate temperatures $T$ and baryon densities $\mu_b > 0$. Between the crossover and the first-order transition, a second-order transition can be found at a critical end point in the phase diagram.

The QCD phase diagram can be investigated with ultrarelativistic nucleus-nucleus collisions. As the nucleons collide an initial and deconfined state of dense and hot matter is formed. The evolution of this QGP fireball is determined by strong anisotropic pressure and density distributions and thus momentum gradients of the quarks and gluons within the volume. The system starts to expand and cools down while forming hadronic matter (also known as hadronization). During the further expansion, the hadrons...
Chapter 1 Introduction

Figure 1.6: Fourier coefficient $v_2$ as a function of the transverse momentum $p_t$ for different particle species within Au-Au collision at RHIC [20].

interact inelastically until the chemical freeze-out has been reached, i.e. inelastic collisions cease and the abundance ratios of the created hadron species do not change anymore [18]. Once thermal freeze-out has been reached, even elastic scattering processes end and the momenta of the particles cease to change. A Fourier analysis of the angular distribution of the produced particles momenta from the fireball allows to introduce a Fourier coefficient $v_2$ (also known as elliptic flow) which can be expressed as a function of the transverse momentum $p_t$ of the particles. The elliptic flow is a observable which allows to study the initial spatial anisotropic geometry of the fireball in heavy-ion collisions [19]. A symmetric particle distribution is found for $v_2 = 0$, i.e. $v_2 > 0$ is a measure for the anisotropy of the distribution. Measurements from Au-Au collisions at RHIC have been compared to hydrodynamical calculations (Fig. 1.6 see [20, 21] for details) and show a good agreement for different particles, i.e. hadronic matter within the QGP phase "flows" like a liquid. As a consequence of the high density within the QGP, even quarks and gluons of high momentum can not leave the fireball without multiple scattering. This leads to a strong suppression of hadrons with high $p_t$. The suppression is expressed by the nuclear modification factor $R_{AA}$ which correlates the number of events in nucleus-nucleus collisions to that produced in pp collisions (normalized such that no energy loss will lead to $R_{AA} \sim 1$). Based on Fig. 1.7 a strong suppression by a factor of approximately 5 can be observed in case of $\pi^0$ as well as $\eta$ mesons at high transverse momenta. As photons only interact electromagnetically, no suppression can be observed in case of direct photons (photons emerging from the initially probed state).

A profound and detailed introduction to QCD phase diagrams and further signatures to probe the existence and the properties of the QGP phase can be found in [18, 22–24] and the included references.

After the ongoing upgrade of the Large Hadron Collider (LHC) at CERN during the Long Shutdown 2 (LS2), further studies will be done to probe QCD matter and to continue with the heavy-ion program of ALICE through Pb-Pb collisions in Run 3 and Run 4 (see Sec. 3). In the context of the upgrade of the Large Hadron Collider, the interaction rate of ALICE will be increased up to 50 kHz for Pb-Pb collisions. As a consequence, a continuous readout of the Time Projection Chamber (TPC) will be required which is currently limited by rates in the order of 300 Hz due to a gated readout system. A
solution with four Gaseous Electron Multipliers (GEM) foils has been adopted as baseline solution for the upgraded chambers.

1.2 Outline

The basic concepts of gaseous detectors will be introduced in Sec. 2 followed by a detailed introduction to ALICE and the upgrade program of the ALICE Time Projection Chamber (cf. Sec. 3).

Within the scope of this work, an alternative approach consisting of two GEM foils and a single Micromegas (also referred to as hybrid detector) has been investigated in terms of the energy resolution, the ion backflow and the gain (cf. Sec. 3). Micromegas offer an intrinsic suppression of the ion backflow of the order of a few percent at relatively high gains. As a consequence less GEM stages are required to block the back drifting ions. The GEM foils are commonly used as pre-amplification stages in order to reduce the needed gain of the Micromegas and to keep the discharge probability at a reasonable level. The obtained results (energy resolution, ion backflow and gain) will be compared to the quadruple GEM stack of ALICE as well as to an alternative hybrid setup which has been investigated at the university of Yale. A big focus lies on a profound and dedicated study of the energy spectra given by $^{55}$Fe. The background distribution will be investigated in detail which finally leads to an advanced fit model in order to describe the full spectrum. Different methods to obtain the energy resolutions will be discussed and compared.

The gains, the energy resolutions and the ion backflows of Micro-Pattern Gaseous Detectors (MPGD) highly depend on the charge transfer processes between the individual amplification stages. Within this work, an analytic model will be derived in order to describe the transfer probabilities of the charge carriers (cf. Sec. 3). The obtained equations inherently include the appropriate limits and offsets that can be seen in experiments and simulations so far. The transfer model will be discussed in detail and compared to simulations. A big advantage of the model calculations is the fast calculation time compared to the time-consuming microscopic simulations.
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The transfer probabilities give access to an in-depth study of the fundamental properties and processes within GEM stacks. Based on the transfer calculations, models will be derived in order to describe the energy resolution, the gain as well as the ion backflow for GEM stacks (cf. Sec. 6). The outcome of these calculations will be compared to measurements with the Bonn hybrid detector, the alternative hybrid setup of Yale as well as with the ALICE baseline solution based on four GEM foils.
CHAPTER 2

Gaseous detectors

Starting with the very first Geiger-Müller counter [26] in 1928, the invention of the Multiwire Proportional Chamber (MWPC) [27] in 1968, or the development of the Gas Electron Multiplier (GEM) [28] in 1997: Gaseous detectors have been developed and studied in numerous designs and patterns within the last decades. Nowadays they play an important role in many modern high energy physics experiments in order to identify charged particles or to reconstruct their trajectories. Though there are plenty of different designs, all gas-filled detectors are based on a common working principle: Once a charged particle traverses the active volume of the detector, different ionization processes can take place within the gas which create electron-ion pairs. The interaction of charged particles as well as of photons with matter will be discussed in Sec. 2.1 to Sec. 2.3. Commonly the created electrons are guided by electric fields to amplification stages. The movement of the charges will be part of Sec. 2.4. Usually the amount of created electrons is too small to be detected and requires an additional charge multiplication (see Sec. 2.6). The final charges are thereupon read out and allow to extract spatial information about the incident particles (cf. Sec. 2.7). The physics of these processes will be introduced and discussed in the following sections.

2.1 Interaction of charged particles with matter

Heavy charged particles interact electromagnetically with matter in multiple inelastic collision processes. Within each individual collision a certain amount of energy is lost through atomic ionization or excitation. Within primary ionization processes of charged particles (p) with matter, electrons (e⁻) are liberated from the shells of the interacting atom (X):

\[ p + X \rightarrow p + X^+ + e^- . \]  (2.1)

Some of the primary electrons are able to ionize further atoms as the remaining energy is still sufficient for additional secondary ionization processes. The kinetic energy distribution of the primary ionization electrons shows a less probable but long tail for higher energies. Electrons with very high energies from head-on collisions are emitted in forward direction. However most of the higher energetic electrons (energies of a few keV) are emitted perpendicularly to the direction of the incident particle and cause ionizations far from the track. These electrons are also referred to as delta electrons. As a consequence the resolution of gaseous detectors is limited by delta electrons. A further ionization process is the Penning effect. Once excited, some atoms are able to remain in a metastable state (X'). Through collisions with

[1] Also elastic processes are possible but neglected as this does not cause noticeable energy losses.


<table>
<thead>
<tr>
<th>Gas</th>
<th>W / eV</th>
<th>Gas mixture</th>
<th>W / eV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ne</td>
<td>37</td>
<td>Ne-CO₂ (90-10)</td>
<td>38.1</td>
</tr>
<tr>
<td>Ar</td>
<td>26</td>
<td>Ne-CO₂-N₂ (90-10-5)</td>
<td>37.3</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ar-CO₂ (90-10)</td>
<td>28.8</td>
</tr>
</tbody>
</table>

Table 2.1: Average energy per produced electron-ion pair W for different gases and mixtures \[30, 32\].

atoms (Y) of a different type, a de-excitation can occur which leads to an ionization:

\[
X^* + Y \rightarrow X + Y^+ + e^- .
\]

The quantity which describes the collision probability between two successive collisions for an incoming particle is the mean free path \(\lambda\). The mean free path can be correlated to the density of the electrons \(n_e\) and the collision cross section \(\sigma\) according to

\[
\lambda = \frac{1}{n_e \cdot \sigma}.
\]

The average amount of primary collisions within a length \(L\) is given by \(\mu = L/\lambda\) and is described by a Poisson distribution \[29\]. Indeed most of the electrons emerge from secondary ionization processes: For Argon and at normal temperature (20 °C) and pressure (one atm) the total amount \(N_T = 97 \text{ cm}^{-1}\) of created electron-ion pairs per centimeter of track length is larger than the amount of primary electron-ion pairs \(N_P = 25 \text{ cm}^{-1}\) (for minimum ionizing particles, also MIPs). The same holds for Neon \(N_T = 40 \text{ cm}^{-1}\) and \(N_P = 13 \text{ cm}^{-1}\) \[30, 31\] which will be the main gases used in the framework of this PhD thesis. Within each single collision only a certain amount of energy is lost in ionization processes. The total number of ionizations \(N_T\) can be expressed by the average energy to create a free electron \(W\) and the track length \(L\) according to Eq. 2.3.

\[
W \cdot N_T = L \cdot \left\langle \frac{-dE}{dx} \right\rangle
\]

Some example energies \(W\) for the investigated gas mixtures are listed in Table 2.1. The stopping power \(\left\langle -dE/dx \right\rangle\) has been introduced as the mean energy loss per unit path length and will be part of the following section.

### 2.2 Energy loss

The mean energy loss per unit path length can be described by the Bethe equation (also called Bethe-Bloch equation, cf. Eq. 2.4). The equation holds for heavy (i.e. heavier than electrons) charged particles in a kinematic range of \(0.1 \leq \beta \gamma \leq 1000\) \[30\].

\[
\left\langle -\frac{dE}{\rho dx} \right\rangle = K \varepsilon^2 Z \frac{1}{A \beta^2} \left[ \ln \frac{2m_e c^2 \beta^2 \gamma^2 W_{\text{max}}}{I^2} - \beta^2 - \frac{\delta(\beta \gamma)}{2} \right]
\]

The absorbing material is described by the atomic number \(Z\) and the atomic mass \(A\). The incident particle is described by the charge number \(z\) and the velocity \(\beta = v/c\). Accordingly the Lorentz factor is given by \(\gamma^2 = 1/(1 - \beta^2)\). The maximum possible energy transfer in a single collision is denoted by \(W_{\text{max}}\) and the mean excitation energy of the absorbing material by \(I\). Furthermore, the remaining constant \(K = 4\pi N_A r_e^2 m_e c^2 = 0.307075 \text{ MeV mol}^{-1} \text{ cm}^2\) includes \(N_A\) as Avogadro’s number, the classic electron radius \(r_e\), the electron mass \(m_e\) and the speed of light \(c\). The average energy loss is mostly independent
2.2 Energy loss

The mean energy loss for some selected gases and gas mixtures as a function of $\beta\gamma$ is shown in Fig. 2.1. The plot is based on Eq. 2.4 with values from Tab. 2.2.

The mean energy loss can be approximated by $\langle -dE/d\rho \rangle / \rho \approx 2\text{ MeV g}^{-1}\text{ cm}^{2}$. For higher energies the logarithmic term starts to dominate which leads to an increase of the stopping power. This effect is also known as the relativistic rise: With respect to the flight direction of the incident particle it can be shown that the longitudinal component of the electric field flattens and the transversal component extends (as seen from the rest frame of the absorber). Additionally, the maximum possible energy transfer $W_{\text{max}}$ increases with $\beta\gamma$ which leads to a higher average energy loss. Nevertheless the relativistic rise is partly compensated by the density effect which is described by the term $\delta(\beta\gamma)$ in Eq. 2.4. Due to the electric polarization of the absorbing medium the electric field of the incident particle is altered and leads to a limitation of the interaction range. Electrons that are far from the track are screened and do not contribute to the mean energy loss as predicted by Bethe-Bloch. The name of this effect already suggests that it depends on the density of the absorbing material. The induced polarization is larger for dense matter, i.e. the corrections of the density effect increase with the density of the absorber. Some selected gas properties, mean excitation energies and average energy losses in the minimum are listed in Tab. 2.2. In case of gas mixtures the mean excitation energy can be approximated by Bragg’s additivity rule according to Eq. 2.5 with the electron densities $n_i$ and the mean excitation energies $I_i$ of the single components.

$$\ln I = (\sum n_i \ln I_i) / \sum n_i$$

(2.5)

The densities can be rewritten as $n_i = (pV_i)/(RT)$ where the pressure $p$ and the temperature $T$ are assumed to be constant. If the total volume of the gas mixture $V$ is fixed, the mean excitation energy can...
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<table>
<thead>
<tr>
<th>Gas</th>
<th>Z</th>
<th>A / g mol⁻¹</th>
<th>Density ρ / g l⁻¹</th>
<th>Mean excitation energy I / eV</th>
<th>⟨-dE/dx⟩/ρ / MeV g⁻¹ cm²</th>
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<tr>
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<td>1.52</td>
</tr>
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<td>1.98</td>
<td>85</td>
<td>1.83</td>
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<tr>
<td>N₂</td>
<td>14</td>
<td>28.01</td>
<td>1.25</td>
<td>82</td>
<td>1.83</td>
</tr>
</tbody>
</table>

Table 2.2: Selected atomic properties, mean excitation energies and specific ionization in the minimum [34].

be written as

\[ \ln I = \sum f_i \ln I_i \]  

where the volume fraction is defined as \( f_i = \frac{V_i}{V} \). The mean excitation energy of Ne-CO₂ (90-10) is given by 130.6 eV. By adding Nitrogen the mean excitation energy is lowered to 127.8 eV for Ne-CO₂-N₂ (90-10-5). In case of Ar-CO₂ (90-10) it is given by 173.7 eV.

2.3 Interaction of photons with matter

Since photons carry no electric charge they follow different interaction mechanisms with matter compared to heavy and charged particles. Generally photons are more capable of penetrating through matter due to smaller cross sections. They interact either by absorption or by scattering processes which change the intensity but not the energy of the incident beam. The intensity of the beam falls exponentially with the thickness \( x \) of the absorber according to

\[ I(x) = I_0 \exp(-\mu x) \]  

where \( I_0 \) corresponds to the primary beam intensity. The mass absorption coefficient \( \mu \) can be expressed by the total cross section \( \sigma \) and the atom density \( N \) (cf. Eq. 2.8). Like for heavy and charged particles a mean free path \( \lambda \) can be found for the interaction of photons with matter.

\[ \mu = N \sigma = \frac{1}{\lambda} \]  

The main interaction mechanisms of photons in matter are the \textit{photoelectric effect}, \textit{Compton scattering} and \textit{pair production}. The physics of the mentioned processes will be briefly summarized. A more detailed introduction can be found in [35] or [36].

2.3.1 Photoelectric effect

The photoelectric effect describes the absorption of a photon by an atomic bound electron. The electron is thereupon kicked out of the atom. Due to energy and momentum conservation, this effect can not occur for free electrons: The recoil momentum is transferred to the nucleus and the energy to the electron. The bound electron is capable of escaping from the atom if the energy of the incoming photon \( E_\gamma = h \nu \) exceeds its binding energy \( E_B \). The kinetic energy \( E_{\text{kin}} \) of the outgoing electron is given by Eq. 2.9 [37].

\[ E_{\text{kin}} = E_\gamma - E_B \]  

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2.3 Interaction of photons with matter

Fig. 2.2 shows the cross section of the photoelectric effect for some selected noble gases and copper as a function of the incident photon energy. As the photon energy increases, a photoelectric ionization of the inner atomic structure becomes possible once the binding energy of a specific shell has been reached. In this case, the cross section increases which leads to the characteristic absorption edges (both absorption edges in Fig. 2.2 correspond to the inner K-shell). The probability for the photoelectric cross section turns out to be higher for inner shells, i.e. stronger bound electrons which are closer to the nucleus. The following cross section holds for the innermost K-shell in case of the non-relativistic limit \( E_\gamma \ll m_e c^2 \) and for photon energies above the absorption edge

\[
\sigma_{\text{p.e.}}^K (\epsilon \ll 1) = \left( \frac{32}{\epsilon} \right)^{1/2} \alpha^4 Z^{4.5} \sigma_{\text{Thomson}} \frac{Z^{4.5}}{\epsilon^{7/2}}
\]

with \( \epsilon = E_\gamma / (m_e c^2) \), the Fine-structure constant \( \alpha \) and the Thomson cross section \( \sigma_{\text{Thomson}} = \frac{8}{3\pi \epsilon^2} \) [36]. For high energies \( E_\gamma \gg m_e c^2 \) the energy dependence changes and the cross section is given by

\[
\sigma_{\text{p.e.}}^K (\epsilon \gg 1) = 4\pi \epsilon^2 Z^{4.5} \alpha^4 \frac{1}{\epsilon} \propto \frac{Z^{4.5}}{\epsilon}.
\]

The dependence on the atomic number \( Z \) turns out to be correlated to the photon energy and varies from 4 to 5 in a range between 0.1 MeV \( \leq E_\gamma \leq 5 \) MeV. Obviously a stronger shielding can be obtained by choosing materials of higher atomic number \( Z \). The energy of the photon beam is conserved but the intensity gets reduced as the interacting photons are completely absorbed.

### 2.3.2 Compton scattering

If the photon energy is much higher than the binding energy \( (E_\gamma \gg E_B) \) the electrons can be regarded as quasi-free. The incident photons are capable of scattering on the quasi-free electrons which is known as Compton scattering. The photon energy lowers from \( E_\gamma \) to \( E_\gamma' \) but the intensity remains unchanged as no
Figure 2.3: Energy spectrum of the recoil electron for Compton scattering and different energies $h\nu$ of the incident photon [35].

photons are absorbed. The energy transfer $T$ is given by Eq. (2.12) and has a maximum for a scattering angle of $\theta = 180^\circ$, i.e. the photons are scattered back. The maximum energy transfer can be seen as a sharp drop (Compton edge) in the energy spectrum of the recoil electron (cf. Fig. 2.3).

\[
T = E_\gamma - E_\gamma' = E_\gamma \frac{\epsilon (1 - \cos \theta)}{1 + \epsilon (1 - \cos \theta)}
\]  

(2.12)

The cross section was first derived and published by Klein and Nishina (see [39]) and leads to

\[
\sigma_{\text{Compton}} = \frac{3}{4} \sigma_{\text{Thomson}} \left[ \frac{1 + \epsilon}{\epsilon^2} \left( \frac{2 (1 + \epsilon)}{1 + 2 \epsilon} - \frac{1}{\epsilon} \ln (1 + 2 \epsilon) \right) + \frac{1}{2 \epsilon} \ln (1 + 2 \epsilon) - \frac{1 + 3 \epsilon}{(1 + 2 \epsilon)^2} \right].
\]

(2.13)

Following relations can be obtained for the total Compton cross section in the limits of $\epsilon \ll 1$ and $\epsilon \gg 1$:

\[
\sigma_{\text{Compton}} \propto \begin{cases} 
1 - \epsilon & \text{if } \epsilon \ll 1, \\
\frac{1}{\epsilon} (1 + 2 \ln 2 \epsilon) & \text{if } \epsilon \gg 1.
\end{cases}
\]

(2.14)

Due to energy conservation, both energies of the recoil electron and the outgoing photon add up to the total energy of the incoming photon. Accordingly the total cross section $\sigma_{\text{Compton}}$ can be split into two distributions $\sigma_{C}$ and $\sigma_{A}$ with $\sigma_{\text{Compton}} = \sigma_{C} + \sigma_{A}$. The cross section $\sigma_{C}$ belongs to the average fraction of the total energy which is contained in the scattered photon (index $s$) and the cross section $\sigma_{A}$ belongs to the fraction which is absorbed by the recoil electron (index $a$). Fig. 2.4 shows the total Compton cross section and both fractional cross sections as a function of the incoming photon energy.

### 2.3.3 Pair production

In the vicinity of a nucleus and its corresponding Coulomb field, an incoming photon is capable of converting into an electron-positron-pair. This process requires a minimum photon energy of $E_\gamma \geq$
2.3 Interaction of photons with matter

Figure 2.4: Total cross section for Compton scattering of photons on free electrons [35].

The cross section is strongly affected by the screening of the nuclear Coulomb field by the surrounding outer electrons. For low energies \( E_\gamma \ll m_e c^2 \), only photons close to the nucleus reach a sufficiently high probability to undergo pair production. As they are close to the nucleus, the screening of the outer atomic electrons can be neglected and the photon interacts with the unscreened nucleus. In this case, the cross section is given as [36]:

\[
\kappa_{\text{nuc}} = 4 \alpha r_e^2 Z^2 \left( \frac{7}{9} \ln 2 \epsilon - \frac{109}{54} \right) \propto Z^2 \ln 2 \epsilon .
\] (2.15)

At high energies \( E_\gamma \gg m_e c^2 \), also photons far from the nucleus converge in the Coulomb field. The screening of the outer atomic electrons must be taken into account and leads to the following cross section [36]:

\[
\kappa_{\text{nuc}} = 4 \alpha r_e^2 Z^2 \left( \frac{7}{9} \ln \frac{183}{Z^{1/3}} - \frac{1}{54} \right) \propto Z^2 \ln \frac{183}{Z^{1/3}} .
\] (2.16)

Although less probable, pair production might also occur in the Coulomb field of another electron, leading to a different cross section \( \kappa_e \) (see [36] for more information). The dependence on the photon energy is shown in Fig. 2.4 as example for some selected noble gases and copper: As explained, pair production takes only place for energies above a threshold of 1.022 MeV. The energy of the incident photon beam remains unchanged. Yet the intensity lowers as the amount of photon decreases.

2.3.4 Total cross section

The total cross section for photon interactions in matter \( \sigma \) can be written as the sum of the major contributions, i.e. the photoelectric effect \( \sigma_{\text{p.e.}} \), Compton scattering \( \sigma_{\text{Compton}} \) and pair production \( \kappa_{\text{nuc}} / \kappa_e \) (cf. Eq. 2.17). The multiplication with \( Z \) takes the number of electrons per atom into account as the
Compton cross section from Eq. \[(2.13)\] holds only for a single electron.

\[
\sigma = \sigma_{\text{p.e.}} + Z \cdot \sigma_{\text{Compton}} + \kappa_{\text{nuc}} + \kappa_{\text{e}}
\]  

(2.17)

The resulting superpositions of all cross sections are shown in Fig. 2.5 for some selected noble gases and copper. For low energies \(E_\gamma \ll 1 \text{ MeV}\), the total photon cross section is dominated by the photoelectric effect. For medium photon energies \(E_\gamma \approx 1 \text{ MeV}\), Compton scattering becomes more dominant. In case of high energies \(E_\gamma \gg 1 \text{ MeV}\), pair production plays the most important role.

### 2.4 Charge movement in gases

The interactions of charged particles and photons with matter lead to excitation or ionization (cf. Sec. 2.1 and 2.3). Electrons which emerge from these processes play an important role for particle identification and spatial reconstruction with gaseous detectors. Commonly the amount of primary and secondary electrons is too small to be detected and requires an additional amplification. By using electric fields the electrons can be "guided" to further amplification or readout stages. The movement of the charge carriers (electrons as well as ions) within electric and magnetic fields will be part of the following section.

#### 2.4.1 Drift of electrons in gases

The movement of the electrons can be compared to a "stop and go" motion as they scatter isotropically on the gas molecules. In each elastic collision a fractional amount of the energy is lost due to recoil or excitation and the direction of motion as well as the momentum are randomized. Between two successive collisions the moderated electrons rapidly gain kinetic energy as they accelerate in the external electric field which is again partially lost in a following collision. The electron movement is characterized by the
2.4 Charge movement in gases

Figure 2.6: Total cross sections for some selected noble gases and copper. The plots show the individual cross sections and the total cross section as a function of the photon energy $E_\gamma$ [38].
drift velocity \( u \) (gained velocity of the electron in the external electric field between collisions) and – in addition – by the instantaneous and randomly oriented velocity \( c \) after a collision (where \( u \ll c \)). In the \textit{microscopic picture} these quantities can be expressed as a function of the number density \( n \), the collision cross section \( \sigma \) and the average fractional energy loss per collision \( \Lambda \) according to \cite{29}:

\[
u^2 = \frac{eE}{n \sigma m_e} \sqrt{\frac{\Lambda}{2}},\tag{2.18}
\]

\[
c^2 = \frac{eE}{n \sigma m_e} \sqrt{\frac{2}{\Lambda}}.\tag{2.19}
\]

Both the cross section \( \sigma \) as well as the fractional energy loss \( \Lambda \) are functions of the kinetic energy of the drifting electron \( \epsilon \). Fig. 2.7 shows the cross section for neon and argon. In case of argon, the cross section has a minimum at energies near \( \epsilon \approx 0.25 \text{ eV} \) which is also known as the \textit{Ramsauer minimum}. The dip is caused by quantum-mechanical processes in the scattering of the electron with the gas molecules (see \cite{40} for details). On average this "stop and go" motion leads to an equilibrium energy which determines the macroscopic drift velocity \( \bar{u} \). In the \textit{macroscopic picture} the equation of motion can be written as

\[
m_e \frac{d\bar{u}}{dt} = e\bar{E} + e[\bar{u} \times \bar{B}] - K\bar{u}\tag{2.20}
\]

with the electric field \( \bar{E} \) and the magnetic field \( \bar{B} \). The first term in Eq. 2.20 describes the movement in the electric field. The second term corresponds to the Lorentz force in the magnetic field. The interaction of the particles with the gas is described by the frictional force which is proportional to the velocity and given by \( K\bar{u} \) where \( K \) can be expressed by the mean collision time as \( K = m_e/\tau \). In literature Eq. 2.20 is also referred to as the \textit{Langevin equation}. The general solution for the velocity is given by

\[
\bar{u} = \frac{e}{m_e} \tau E \frac{1}{1 + \omega^2 \tau^2} \left( \bar{E} + \omega \tau \left[ \bar{E} \times \bar{B} \right] + \omega^2 \tau^2 \left( \bar{E} \cdot \bar{B} \right) \bar{B} \right) \tag{2.21}
\]
2.4 Charge movement in gases

Figure 2.8: Electron drift velocity and diffusion coefficients (longitudinal as well as transversal) for different gas mixtures as a function of the electric field strength \([42]\).

where \(\hat{E}\) and \(\hat{B}\) denote the unit vectors in the directions of the fields. The circular motion in the magnetic field is characterized by the Larmor frequency \(\omega = -\left(\frac{e}{m_e}\right) B\). In case of no magnetic field Eq. \(2.21\) simplifies and leads to

\[
d \vec{u} = \frac{e\tau}{m_e} \vec{E} = \mu \vec{E}
\]

where the mobility \(\mu\) has been introduced\(^2\). A detailed derivation and discussion for different field scenarios can be found in \([29]\). The drift velocities for neon and argon mixtures are shown in Fig. \(2.8\) as a function of the external electric field. For common drift fields of 400 V cm\(^{-1}\) the drift velocity (the mobility) is given by \(u = 2.73 \text{ cm} \mu\text{s}^{-1} (\mu = 6.825 \text{ cm}^2 \text{ V}^{-1} \text{s}^{-1})\) in Ne-CO\(_2\) 90-10. Adding a small amount of nitrogen barely influences the drift velocity (the mobility) and leads to 2.58 cm\(\mu\)s\(^{-1}\) (6 450 cm\(^2\) V\(^{-1}\) s\(^{-1}\)). In case of Ar-CO\(_2\) 90-10, the drift velocity (the mobility) is given by 3.31 cm\(\mu\)s\(^{-1}\) (8 275 cm\(^2\) V\(^{-1}\) s\(^{-1}\)).

\(^2\)The mobility is also a function of the electric field, i.e. \(\mu = \mu(E)\).
2.4.2 Drift of ions in gases

The drift motion of ions differs from electrons as they have a much higher mass. Though their energy gain in external electric fields is similar to that of electrons, they lose a larger fractional amount of energy in collisions. The ion momentum is not randomized as for electrons and ions are capable of "memorizing" their former momentum and direction. Due to this the diffusion of ions is much smaller than the diffusion of electrons as it will be discussed in Sec. 5.4.1. The mobility of ions does not vary as much as for electrons. For small electric fields (where the thermal motion can not be neglected) the drift velocity is given by

\[ u_\perp = \left( \frac{1}{m_{\text{ion}}} + \frac{1}{m_{\text{gas}}} \right)^{1/2} \left( \frac{1}{3kT} \right)^{1/2} \frac{eE}{n\sigma} \propto E \]  

and the mobility turns out to be independent of the electric field (cf. Eq. 2.22, \( k \) Boltzmann’s constant, \( T \) gas temperature, \( m_{\text{ion}} \) mass of scattering ion and \( m_{\text{gas}} \) mass of gas molecules). For large electric fields it can be shown that

\[ u_\parallel = \left( \frac{eE}{m_{\text{ion}}n\sigma} \right)^{1/2} \left( \frac{m_{\text{ion}}}{m_{\text{gas}}} \left( 1 + \frac{m_{\text{ion}}}{m_{\text{gas}}} \right) \right)^{1/2} \propto E^{1/2} . \]  

Accordingly the mobility decreases as \( 1/ \sqrt{E} \). The ion mobility has been measured for gas mixtures like Ar-CO\(_2\) or Ne-CO\(_2\) for different water contents and admixtures of N\(_2\). A detailed study can be found in [43] where the reduced mobility \( K_0 \) (at normal temperature and pressure) and the mobility \( K \) (at given temperature \( T_{\text{Meas}} \) and atmospheric pressure \( p_{\text{Meas}} \)) are correlated by:

\[ K_0 = K \times \frac{237.15 \text{ K}}{T_{\text{Meas}}} \times \frac{p_{\text{Meas}}}{1013 \text{ mbar}} . \]  

Fig. 2.9 and 2.10 show the inverse reduced ion mobility for different Ar-CO\(_2\) and Ne-CO\(_2\)-N\(_2\) mixtures. For Ne-CO\(_2\) (90-10) and common drift fields like 400 V cm\(^{-1}\), the inverse ion mobility can be approximated as \( 1/K_0 \approx 0.325 \text{ V s cm}^{-2} \) and leads to \( K \approx 3.74 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1} \) (in case of \( T_{\text{Meas}} = 294.15 \text{ K} \) and \( p_{\text{Meas}} = 1033 \text{ mbar} \)). Adding a small amount of nitrogen leads to a comparable ion mobility of \( K \approx 3.56 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1} \) for Ne-CO\(_2\)-N\(_2\) (90-10-5). In case of Ar-CO\(_2\) (90-10), the ion mobility decreases and is given by \( K \approx 2.17 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1} \), i.e. neon-based gas mixtures should be preferred in terms of ion mobility. Apparently the electron mobility (cf. Sec. 2.4.1) is by a factor of \( \approx 10^3 \) higher than the ion mobility. As a consequence ions remain much longer in a given electric field than electrons. This effect can lead to an accumulation of ions and unwanted field distortions for gaseous detectors.

2.4.3 Diffusion

Electrons and ions do not strictly follow electric field lines. As introduced in Sec. 2.4, the movement is determined by elastic scattering processes which affect the momentum and the direction after each collision. These processes lead to a random "zigzag" like trajectory of the charge carriers while drifting in an electric field. A pointlike cloud of electrons will be broadened after a certain drift length. Within electric fields, the shape of the broadening turns out to be non-isotropic (electric anisotropy) and can be described by a two-dimensional Gaussian distribution [44]. Assuming that the electrons start at the same origin, the transversal and longitudinal width of the electron cloud (with respect to the macroscopic drift
2.4 Charge movement in gases

Figure 2.9: Measured inverse reduced ion mobility for different Ar-CO$_2$ mixtures as a function of the external electric field (water content 34 ppm – 98 ppm) [43].

Figure 2.10: Inverse reduced ion mobility for different Ne-CO$_2$-N$_2$ mixtures as a function of the external electric field (water content < 20 ppm) [43].
direction) after the time \( t \) is given by

\[
\sigma_{T/L} = \sqrt{2D_{T/L} t}
\]  

(2.26)

where the transversal \( D_T \) and the longitudinal \( D_L \) diffusion coefficients have been introduced. The geometric expansion of the charge cloud can be expressed as a function of the drift length \( z_{\text{Drift}} \). By introducing the diffusion constants \( D_{\text{const},T/L} = (2D_{T/L}/v_{\text{Drift}})^{1/2} \), Eq. (2.26) turns into

\[
\sigma_{T/L} = D_{\text{const},T/L} \sqrt{z_{\text{Drift}}}. 
\]  

(2.27)

Example values for the transversal as well as the longitudinal diffusion constants are shown in Fig. 2.8 as a function of the electric field and for different gas mixtures (no magnetic fields are applied). As example: In Ne-CO\(_2\) (90-10) and for common drift fields of 400 V cm\(^{-1}\), the transversal diffusion constant is given by \( D_{\text{const},T} = 225 \mu m / \sqrt{cm} \). Assuming a drift distance of 3 cm, the transversal broadening is given by \( \sigma_T \approx 390 \mu m \).

Inside a magnetic field, transversal as well as longitudinal diffusion can be observed with respect to the direction of the magnetic field \( \vec{B} \). This effect is also known as magnetic anisotropy. In fact, the longitudinal diffusion (with respect to the magnetic field) remains unchanged as the transversal diffusion \( D_T(\omega) \) is given by

\[
D_T(\omega) = \frac{D_T(0)}{1 + \omega^2 \tau^2}
\]  

(2.28)

with the cyclotron frequency \( \omega = (e/m_e)B \) of the electron and the mean collision time \( \tau \). The transversal diffusion without a magnetic field is denoted by \( D_T(0) \). Obviously, the transversal diffusion can be reduced by applying an external magnetic field.

If an electric and a magnetic field are present, both the electric as well as the magnetic anisotropies combine. In case of parallel fields, a reduction of the transversal diffusion can be achieved which allows to improve the spatial resolution of gaseous detectors. In fact, this effect will be used for the ALICE Time Projection Chamber (TPC) (see [32]).

### 2.5 Electron Attachment

As the produced electrons drift to the electrodes, they encounter molecules which are able to attach them. These impurities are commonly caused by unwanted leakages of the detector vessel or the gas system due to constructional limitations. Depending on the electric field strength, also the quenching gas might cause attachment. Drifting electrons (with kinetic energies in the order of \( \approx eV \)) are commonly attached to electronegative gases such as O\(_2\), H\(_2\)O or halogen-containing impurities. As a consequence the number of detectable charges gets reduced. The attachment probability depends on the kinetic energy of the electrons and thus on the electric field. Fig. 2.11 shows the attachment factor for different argon and neon gas mixtures. The water and oxygen contents (200 ppm and 30 ppm) have been chosen similar to the experimental conditions of this work. In Ne-CO\(_2\) (90-10) the influence of attachment starts to dominate for fields higher than 2500 V cm\(^{-1}\) and can be neglected for common drift fields in the order of 400 V cm\(^{-1}\). The amount of attached electrons \( dN \) in a small distance \( dl \) and for an electric field \( E \) is given by \( dN = -NC(E) \, dl \). With the initial amount of electrons \( N_0 \), the fractional amount of the
remaining electrons can be calculated according to

\[ \frac{N}{N_0} = \exp \left[ -C(E) l \right]. \quad (2.29) \]

As example: In a small drift gap of 4 mm and for a field strength of 3 kV cm\(^{-1}\) (Ne-CO\(_2\) (90-10)), the attachment factor is given by \(C = 0.4 \text{ cm}^{-1}\). Accordingly about 84\% of the primary electrons remain and 16\% are lost due to attachment.

## 2.6 Charge amplification and fluctuations

### 2.6.1 Primary fluctuations

The amount of created primary ionization electrons \(N_i\) differs and is determined due to statistical fluctuations. Here the Fano factor \(F\) characterizes the fluctuations of the primary ionizations for a fixed radiation energy \([46]\). As the processes for the creation of the charge carriers are statistically not independent, the Fano factor expresses the difference to a Poisson distribution. The variance of the distribution is given by \(\sigma_i^2 = FN_i\) and turns into a Poisson distribution in case of \(F = 1\). Commonly values \(F < 1\) are found for rare gases, e.g. \(F = 0.13\ [47, 48]\) for neon and \(F = 0.15 - 0.16\ [47, 49, 50]\) for argon.

\(^1\) Also further electrons might be created within secondary ionization processes. Speaking about "primary" ionization electrons takes all initial electrons into account.
2.6.2 Amplification factor / gain

Commonly the amount of primary ionization electrons is too small to detect and requires further charge multiplication. Generally this can be achieved by using high electric fields in the order of several kV/cm as a single electron obtains sufficient energy for further ionization processes. The avalanche formation can be described by the first Townsend coefficient \( \alpha = n \sigma_{\text{ion}} \) where \( n \) describes the density of the gas molecules and \( \sigma_{\text{ion}} \) the cross section for ionization collisions. No general expression exists for \( \alpha \) and it has to be measured for every gas mixture. The Townsend coefficient depends on the specific excitation or ionization cross sections of the gas, the environmental condition (temperature, pressure) and on the electric field strength.

The single electron gain is a statistical quantity and the (average) gain as defined by Eq. 2.30 does not describe the individual amplification process for each primary electron. For non-uniform fields the Townsend coefficient depends on the Townsend coefficient, the gas mixture, the Townsend coefficient and the Townsend coefficient for ionization, i.e where \( E \) is the drift velocity (cf. Sec. 2.4.1) and the diffusion coefficient (cf. Sec. 5.4.1) are barely influenced. This allows to optimize the charge transfer properties by using higher electric fields before gas amplification starts. In case of Ar-CO\(_2\) (90-10) much higher electric fields are required before gas amplification starts. The following relation holds for the amplification factor (also known as gain) \( G \) if a primary amount of electrons \( N_1 \) drifts through an inhomogeneous electric field along a path \( s \):

\[
G = \frac{N}{N_1} = \exp \left( \int_{s_0}^{s_1} \alpha(s) \, ds \right) = \exp \left( \int_{E(s_0)}^{E(s_1)} \frac{\ln 2}{\Delta V} \exp \left( -\int_{E(s_0)}^{E(s_1)} \frac{ds}{E} \right) dE \right) . \tag{2.30}
\]

The points \( s_0 \) and \( s_1 \) denote the integration limits where the electric field strength exceeds the threshold for ionization, i.e where \( E(s_1) \geq E_{\text{min}} \). The field strength \( E_{\text{min}} \) scales with the gas density \( \rho \) by \( E_{\text{min}}(\rho) = E_{\text{min}}(\rho_0) \cdot \rho/\rho_0 \) where \( \rho_0 \) is the normal gas density. Here the linear relation \( \alpha = \beta E \) has been used which was first introduced by Diethorn who derived a formula for the gain in case of thin wires \([51]\). The required potential \( \Delta V \) to create an electron-ion pair has been introduced as \( \beta = \ln 2/\Delta V \). Example values for Ar-CO\(_2\) (90-10) are given by \( E_{\text{min}}(\rho_0) = 20 \text{ kV cm}^{-1} \) and \( \Delta V = 43 \text{ V} \)[52].

2.6.3 Single electron gain fluctuations

The single electron gain is a statistical quantity and the (average) gain as defined by Eq. 2.30 does not describe the individual amplification process for each primary electron. For non-uniform fields the variance of the avalanche distribution \( P(n) \) is given by \( \sigma_n^2 = \bar{n}^2 f \), where the single gain fluctuation factor \( f \) and the mean number of amplified electrons \( \bar{n} \) have been introduced. A detailed study can be found in \([53]\). The avalanche distribution can be described by a Polya distribution, leading to

\[
P(n) = \frac{1}{\bar{n}} \left( \frac{\theta + 1}{\theta + 1} \right)^{\theta+1} \left( \frac{n}{\bar{n}} \right)^{\theta} \exp \left[ -\left( \frac{\theta + 1}{\theta + 1} \right) \frac{n}{\bar{n}} \right] \tag{2.31}
\]

with \( f = 1/ (\theta + 1) \). The value of \( f \) and the shape of the Polya distribution depends on the electric field strength. For low fields the avalanche distribution is found to be exponential, i.e. \( f = 1 \) (for high fields \( 0 \leq f < 1 \)). Fig. 2.13 shows the avalanche distribution for different values of \( f \). As \( f \) is a function of the field strength it also depends on the Townsend coefficient and thus on the gain. Fig. 2.14 shows the single gain fluctuation factor \( f \) as a function of the Townsend coefficient for different gas mixtures. Exponential functions have been used in order to describe the data points.
2.6 Charge amplification and fluctuations

Figure 2.12: Reduced Townsend coefficient $\alpha$ / gas pressure for different gas mixtures as a function of the electric field strength (plot by [42] using Garfield++ [45]).

Figure 2.13: Polya distributions for different values of $f$ and $\bar{n} = 10$. 
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\[ \alpha \mathrm{cm}^{-1} \]

Figure 2.14: Single gain fluctuation factor \( f \) for Ne-CO\(_2\) 90-10 (squares) and Ar-CO\(_2\) 90-10 (circles) at atmospheric pressure and \( T = 293.15 \) K. Open symbols correspond to data from Magboltz not including Penning effect and closed symbols with [54].

2.7 Signal induction

Detecting the passage of particles is a major role of gaseous detectors. The basic interaction mechanism of charged particles and photons with matter (i.e. with the gas-filled volume of the detector) have been introduced in Sec. 2.1, 2.2 and 2.3. These processes lead to the creation of electron-ion pairs. The detection of these ionization electrons can be used in order to reconstruct the passage of a particle. As the amount of initial ionization electrons usually is too small to detect, further amplification is required. Thus electric fields are used to guide the electrons to following amplification stages. The movement of the electrons as well as the ions has been introduced in Sec. 2.4. The initial and avalanche electrons (cf. Sec. 2.6) are now capable of creating signals with sufficient amplitude for further readout electronics.

The amplitude of the signal \( S \) is proportional to the average number of created avalanche electrons \( \bar{n} \) for each of the initial electrons \( N_i \). The fluctuation of the signal is determined by the primary electron fluctuation and the single gain fluctuation according to

\[
\frac{\sigma_S}{S} = \left( \frac{\sigma_i}{N_i} \right)^2 + \frac{1}{N_i} \left( \frac{\sigma_n}{\bar{n}} \right)^2 + \left( \frac{\sigma_{ENC}}{S} \right)^2
\]

(2.32)

where the Equivalent Noise Charge (ENC) has been introduced in order to describe the electronic noise. By taking the variances of the primary electron fluctuation \( F \) and the single gain fluctuation \( f \) (cf. Sec. 2.6.1 and 2.6.3) into account, the signal fluctuation can be rewritten as

\[
\frac{\sigma_S}{S} = \frac{F + f}{N_i} + \left( \frac{\sigma_{ENC}}{S} \right)^2.
\]

(2.33)

The electrons and ions drift in opposite directions due to opposite electric charges, i.e. ions drift to the cathode and electrons drift to the anode. The movement of the charges induces electric current at the electrodes which can be used as readout signals. Commonly the grounded anode is used for readout purposes. Placing a charge \( q \) at \( z = z_0 \) in front of a grounded electrode will induce a surface charge distribution \( \sigma(x, y) \) as depicted in Fig. 2.15(a). To obtain the charge distribution, a mirror charge with opposite load \(-q\) can be assumed at \( z = -z_0 \). This leads to an equivalent scenario which fulfills the aspired boundary condition of a grounded electrode. Using this method of a mirror charge gives access to
obtain the charge distribution $\sigma(x,y)$. More information can be found in [55]. In order to obtain a spatial resolution the electrode can be segmented as shown in Fig. 2.15(b). The load $q$ will now induce charges $Q_n$ on each pad but with different amplitude, allowing to measure the corresponding currents $I_n$ and thus to calculate a central position and to deduce spatial information. As the charge drifts to the electrode with velocity $v$, the amplitudes of the induced charges $Q_n$ will change and currents flow between the pads and ground. A theory to calculate the induced signal on a grounded electrode by a moving point charge is given by Shockley [56] and Ramo [57]. A detailed introduction to this topic can also be found in [29].

### 2.8 Micro-Pattern Gaseous Detectors (MPGD)

An important progress in the field of gaseous detectors was the invention of the Multiwire Proportional Chamber (MWPC) by Charpak et al. in 1968 [27], rewarded with the Nobel Prize in 1992. Till then spatial measurements with gaseous detectors were in fact only possible by taking single photographs of the tracks (e.g. with the bubble [58] or cloud chamber [59]). This technique is obviously limited by low rates of optical imaging and restricted by the spatial resolution of the photographs. Physical events of interest had to be identified by eye which was time-consuming and inefficient. The MWPC allowed to analyze the presence of particles electronically and much higher rates of data acquisition as well as spatial resolutions were now reachable. The possibility to digitize and to store the recorded data gave access to use electronics to search for events of interest. In principle several anode wires were centrally placed between two parallel cathode planes and the volume was filled with different argon mixtures. An incident particle would ionize the gas, leading to avalanche processes at the anode wires and thus measurable signals. Various gaseous detectors based on wire structures have been developed and used for decades. Nevertheless the capacities of theses detectors turn out to be limited by the growing experimental requirements in terms of spatial resolution and rate capabilities. Since the strength of the electric field increases with decreasing distance to the wire, most electron-ion pairs would be produced close to the anode wire. The drift motion of the ions is much smaller than for electrons, i.e. the ions "remain" for a certain time in the vicinity of the wire and screen the electric field. This leads to a reduction of the effective electric field strength and thus to a drop of the gain for high rates. Fig. 2.16 shows the limiting drop of the gain for a MWPC at rates exceeding $\approx 10^4 \text{mm}^{-2} \text{s}^{-1}$. Spatial limitations in the order of $\approx 1 \text{ mm}$ are caused by mechanical tolerances or electrostatic forces between the wires (repulsion or attraction).

Photolithographic etching and coating processes allow to go for smaller detector geometries and are the fundamental techniques for Micro-Pattern Gaseous Detectors (MPGD): As example Gas Electron Multiplier (GEM) [28] foils (see Sec. 2.8.1) allow to operate at higher rates ($>10^6 \text{mm}^{-2} \text{s}^{-1}$, Fig. 2.16).
and improve the spatial resolution to $< 100 \mu m$.

### 2.8.1 Gaseous Electron Multiplier (GEM) foils

Gaseous Electron Multiplier (GEM [28]) foils have become a widely spread and common gas amplification technology to develop detectable signals from single electrons. The investigated GEM foils consist of a 50 $\mu m$ thin polyimide layer with 5 $\mu m$ of copper coated on top of each side. Using photolithographic and chemical processes a hexagonal hole structure is etched inside of the foil. The holes ideally have a biconical shape with an outer hole diameter of 70 $\mu m$ and an inner hole diameter of 50 $\mu m$. For a standard GEM (S) the pitch of the holes is 140 $\mu m$ (cf. Fig. 2.18). Other geometries used in this thesis are the medium-pitch GEM (MP, 200 $\mu m$) and the large-pitch GEM (LP, 280 $\mu m$). Usually a potential difference about 200 V to 500 V is applied between the top and the bottom side of a GEM which leads to a strong electric field inside the holes in the order of several kV / cm. Electrons can be guided to the GEM by using external electric drift fields: Fig. 2.19 shows a single electron starting between the cathode and the GEM while it drifts inside the GEM hole (green line, starting above the GEM). The field strength inside the GEM hole is strong enough to create further electron-ion pairs by gas ionization processes (black points). Finally an amplified electron avalanche can be extracted at the bottom side of the GEM which drifts to the anode. The absolute gain $G_{abs}$ of a GEM is defined as the number of electrons at the exit of the hole divided by the initial number of electrons entering the hole. However, not all electrons might be collected by the GEM hole as they could end on the top side. In addition not all electrons might be extracted as they could end on the bottom side of a GEM. The effective gain $G_{eff}$ takes the collection probability $\epsilon^{-c}$ and the extraction probability $\epsilon^{-e}$ (also referred to as collection / extraction efficiency) of the electrons into account:

$$G_{eff} = \epsilon^{-c} G_{abs} \epsilon^{-e}. \quad (2.34)$$

Due to opposite electric charges, some of the ions (Fig. 2.19, red lines) will drift back and end at the top side of the GEM where they get blocked. As the diffusion of the ions is much smaller than for electrons, they tend to follow the electric field lines while drifting. This can be used in order to intrinsically suppress the amount of back drifting ions by the electrostatic field configuration. Nevertheless some ions still...
drift back to the region where the primary electrons originated from. The amount of back drifting ions is characterized by the ion backflow. As the ion backflow might cause unwanted space-charge distortions and inhomogeneities in the drift region, a suppression of the ion backflow is usually wanted. This can be achieved by stacking up multiple GEM stages on top of each other. Here different GEM pitch geometries (geometric blocking) and specific field configurations allow to enhance the blocking of ions. Fig. 2.20 shows the resulting avalanches for a single incoming electron in a quadruple GEM stack (S-LP-LP-S). As the avalanche ions drift back they partially get blocked by previous GEM stages. Furthermore the stability in terms of discharges can be improved by splitting up the total gain to the individual GEM stages: A high gain can be achieved by keeping the GEM voltages low and the discharge probability of each stage can be kept at a reasonable level. The total effective gain of a stack with \( J \) GEMs \((J \geq 1)\) is given by

\[
G_{\text{eff,stack}} = \prod_{j=1}^{J} G_{\text{eff},j}
\]

where the single effective gains of the stages contribute as \( G_{\text{eff},j} \). Losses of electrons due to attachment (between the GEM stages) are not taken into account by the transfer efficiencies. The ion backflow \( IB \) of a GEM stack is given by

\[
IB = \frac{1 + \epsilon}{G_{\text{eff,stack}}}
\]

with \( \epsilon \) as the total number of back-drifting ions per incoming electron. The charge transfer probabilities of the ions can be expressed by the ion collection \( \epsilon^+_{\text{c}} \) and the ion extraction \( \epsilon^+_{\text{e}} \) efficiency. The number of back drifting ions per incoming electron \( \epsilon \) can be expressed by the electron as well as the ion efficiencies and will be part of Sec. 6.2.1.

An important aspect which influences the ion backflow of a GEM stack is the alignment of two successive GEM foils, i.e the rotation or the displacement. Due to the underlying hexagonal structure of the GEM holes, various Moiré patterns with different areas of optical opaqueness can be observed (see Fig. 2.17). These spatial inhomogeneities of hole overlaps lead to non-uniform distributions of the ion backflow. An optimum and almost uniform distribution can be achieved for a rotation of 90°. Accordingly two successive GEM foils should be rotated by 90° within a GEM stack.

### 2.8.2 Micromegas

Micro-Mesh-Gaseous detectors (Micromegas) were first presented in 1996 by Giomataris et al. \([63]\) and thus appeared approximately at the same time as GEM foils. Historically it consisted of a two-stage parallel plate avalanche chamber which was separated by a micromesh in a small amplification region of 100 µm thickness and a conversion drift region of 3 mm thickness. Fig. 2.21 illustrates the working principle of a Micromegas: Once a charged particle traverses the drift gap it will ionize the gas. The ionization electrons drift towards the anode passing the micromesh. Within the amplification region amplification processes lead to detectable signals at the anode strips. Commonly a strong asymmetric field configuration is applied to the Micromegas, e.g. \( E_1 \approx 1 \text{ kV cm}^{-1} \) in the drift gap and \( E_2 \approx 100 \text{ kV cm}^{-1} \) in the amplification gap. Such high field ratios lead to an intrinsic and efficient collection of the avalanche ions by the micromesh. Only a small fraction of ions manages to enter the drift gap which is inversely proportional to the electric field ratio \( \zeta = E_2/E_1 \). Fig. 2.22 shows the ion and electron transmission
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Figure 2.17: Two rotated GEM foils placed on a light table. Local spatial inhomogeneities of hole overlaps can be observed (Moiré patterns) which lead to non-uniform distributions of the ion backflow \cite{62}.

Figure 2.18: Microscopic picture of a standard GEM foil.
probability for the mesh as a function of the field ratio $\zeta$: Indeed most of the ions are blocked for high ratios and almost all electrons are collected.

Historically the Micromegas was manufactured by stretching and gluing a mesh on a frame made of quartz fibers which then defined the thickness of the amplification gap between the mesh and the readout plane. In 2005 the bulk Micromegas was introduced by Giomataris et al. [64]. Using Printed Circuit Board (PCB) technology and photolithographic methods, the whole sensitive part of the Micromegas (i.e. the anode plane with copper strips/pads, a spacer structure (pillars) for the mesh and the micromesh itself) is laminated together as one object in a single process. The novel technique allowed to produce Micromegas in larger scales and improved the robustness and accuracy of the electrode materials and the thickness of the amplification gap. Fig. 2.23(a) and 2.23(b) show microscopic pictures of such a mesh with 160 lines/cm or approximately 60 $\mu$m per grid pattern. The rectangular structures in the background of Fig. 2.23(a) corresponds to the quadratic readout pads (1 cm $\times$ 1 cm) of the pad plane. The pillars which hold the mesh structure can be seen as circular dots.
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Figure 2.20: Simulation of a quadruple GEM stack (S-LP-LP-S) with Magboltz and Garfield++ [42]. Electron drift paths indicated as green lines, ions as red lines. Field configuration according to ALICE baseline solution (see Tab. 3.2).
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Figure 2.21: Schematic view of a Micromegas detector [63].

Figure 2.22: Electron and ion transmission probability for the mesh as a function of the field ratio [63].
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Figure 2.23: Microscopic pictures showing the mesh and the pad structure of the Micromegas.
CHAPTER 3

The ALICE experiment

3.1 Physics program

The first intentions to build a detector dedicated to heavy-ion physics at the Large Hadron Collider (LHC) go back to the early 1990s and were officially approved in 1996: Investigating the QCD phase diagram and the transition from hadronic matter to the Quark-Gluon Plasma in ultra-relativistic heavy-ion collisions is a major exploratory focus of ALICE (A Large Ion Collider Experiment). The following selected results were obtained from the Run 1 (2010 to 2013) and the Run 2 (2015 to 2017) data taking periods. A full and detailed review can be found in [65–66].

The spatial extension of the fireball at freeze-out can be estimated via an intensity interferometry analysis also know as Hanbury Brown-Twiss analysis (HBT) [67]. Here the extension can be correlated to three radii ($R_{\text{out}}$, $R_{\text{long}}$, and $R_{\text{side}}$) and the volume of the fireball can be estimated as the product of them. Fig. 3.1 shows the product of the radii versus the average charged-track multiplicity density as obtained in central Pb-Pb collisions at ALICE and other lower energetic experiments. The obtained ALICE results indicate that the obtained fireballs in nuclear collisions at LHC are hotter, more stable (lifetime $\tau \approx 10 \text{ fm/c}$, approximately 20% longer than measured at RHIC [65]) and expand to larger scales.

![Figure 3.1: Estimated extension of the fireball at freeze-out as a function of the average charged-track multiplicity density for different center-of-mass energies [68].](image-url)
Figure 3.2: Nuclear modification factor $R_{p\text{Pb}} / R_{\text{PbPb}}$ of charged particles as a function of the transverse momentum $p_t$ for p-Pb / Pb-Pb collisions at ALICE [69].

(volume $V \approx 300 \text{ fm}^3$, larger by a factor of two compared to RHIC [68]) as compared to experiments with lower energies.

The nuclear modification factor (cf. Sec. 1.1) has been measured for charged particles to probe the suppression of hadrons with high $p_t$ at ALICE. Fig 3.2 shows the measured nuclear modification factors $R_{p\text{Pb}} / R_{\text{PbPb}}$ for p-Pb / Pb-Pb collisions. For $p_t \geq 2 \text{ GeV/c}$, a strong suppression of hadrons can be observed in central Pb-Pb collisions which hints at the existence and the high density of the QGP, i.e. quarks and gluons of high momentum can not leave the hot matter created in collisions of heavy ions without multiple scattering [69].

Fig. 3.3 shows the coefficient of the elliptic flow $v_2$ as a function of the traverses momentum $p_t$ for different particle species, measured in Pb-Pb collisions at center-of-mass energies of 5.02 TeV. For lower transverse momenta, similar elliptic flows can be found for the $\phi$-meson and the proton / antiproton which is based on a similar effect of the radial flow on particles with comparable masses ($m_\phi \approx 938 \text{ MeV}/c^2$, $m_p \approx 1020 \text{ MeV}/c^2$ [10]). The measurements extend the $p_t$ range of the RHIC data (cf. Fig. 1.6) to intermediate transverse momenta $p_t \geq 2.5 \text{ GeV/c}$ where the predictability of the hydrodynamic approach is limited. Here other processes start to dominate where the behavior of $v_2$ is affected by the quark contents and the elliptic flows of the $\phi$-meson and the proton / antiproton start to deviate (cf. [65]).

After the ongoing upgrade of the Large Hadron Collider at CERN, further studies will be done within the heavy-ion program of ALICE in Pb-Pb collisions to probe QCD matter at vanishing chemical potentials $\mu_b$ and at much higher initial temperatures. The scientific goals of the upgraded ALICE detector are described in detail in [71] and the included references. Major aspects of the upgraded physics program are:

- Investigate the thermalization of the QGP with focus on charm and beauty quark production.
- Low-momentum quarkonium dissociation and - possibly - regeneration as probe of deconfinement and access to the temperature evolution of QGP.
3.2 Setup and detector

ALICE is located at Point 2 of the LHC in St. Genis-Pouilly, France. The main cavern of the experiment is located approximately 50 m underground and can be accessed via a shaft from the surface facilities. The overall dimensions of the detector are 16 m × 16 m × 26 m with a total weight of approximately 10 000 t.

Fig. 3.4 shows the schematic layout of ALICE and the subdetectors during Run 2 (2015 - 2018). The setup of the experiment will be briefly summarized in the following part. A more detailed and profound introduction to ALICE and the single detectors can be found in [24, 73] and the included references. The central barrel of the experiment houses the Inner Tracking System (ITS) which consists of six cylindrical layers of high-resolution silicon detectors: Two planes of Silicon Pixel Detectors (SPD) record the innermost vertices, followed by two layers of Silicon Drift Detectors (SDD) and two outer layers of Silicon Strip Detectors (SSD). Together with the Time Projection Chamber (TPC, see Sec. 3.3 for a detailed introduction), both detectors can be used for particle tracking and momentum...
reconstruction of charged particles. The Transition Radiation Detector (TRD) can be used for particle identification as well as the High Momentum Particle Identification Detector (HMPID) which is a Ring Imaging Cherenkov (RICH) counter. A Time Of Flight (TOF) detector allows to measure the flight time of particles from the collision point. The masses of the particles can be deduced from the momentum and the time of flight. Together with two electromagnetic calorimeters (PHOS and EMCal) for photon and jet measurements, the central barrel is covered by a huge solenoid magnet (maximum central field 0.5 T) which was formerly used in the L3 experiment at the Large Electron-Positron collider (LEP, dismantled in 2000 for the construction of the LHC) [74]. Two front doors of the magnet allow to completely enclose the inner subdetector system. A forward muon spectrometer is located at one side behind the central L3 magnet. It consists of several absorbers to shield the spectrometer from other reaction products than muons. Several tracking chambers allow to determine the muon tracks before and after passing the field of a dipole magnet (maximum field 0.7 T). Further chambers behind a 300 tonne iron wall contribute as trigger signals for the Central Trigger Processor (CTP).

After the Long Shutdown 2 (LS2 in 2019/2020), the LHC will be operated with an increased luminosity for Pb-Pb collisions up to $\mathcal{L} = 6 \times 10^{37}$ cm$^{-2}$ s$^{-1}$ which corresponds to an interaction rate of 50 kHz. The ALICE experiment will undergo a major technical upgrade including several subdetector systems in order to handle the new run conditions of the LHC. The new ITS will consist of seven high-resolution layers of active pixel sensors with less material budget and an increased data acquisition (DAQ) rate of 100 kHz for Pb-Pb and 1 MHz for pp collisions [72]. An additional Muon Forward Tracker (MFT) based on silicon pixel sensors will be added between the ITS and the absorber of the muon arm to improve the performance of the present muon spectrometer and to access new physics measurements such as open charm / beauty separation [75]. Further improvements will be made for the readout and triggering system, including detectors such as the Muon Tracking Chambers (MCH), Muon Identifier (MID), TRD, TOF and many more (see [76] for details). The present readout system of the Time Projection Chamber will be replaced to allow a continuous data acquisition during Run 3. A detailed introduction to the upgrade of the TPC can be found in [32] and will be part of the following Sec. 3.3.1.

### 3.3 Time Projection Chamber (TPC)

The concept of a Time Projection Chamber (TPC) was first introduced by Nygren in 1987 [78] and will be explained for the ALICE TPC; with its low material budget and excellent pattern recognition capabilities, the ALICE TPC is an ideal gaseous detector for three-dimensional tracking and identification of charged particles. Fig. 3.5 shows the central Time Projection Chamber of ALICE (see [79] for technical details of the TPC design). The outer and inner cylindrical field cages have a diameter of 5.0 m and 1.2 m respectively. With a length of 5.5 m and an active volume of about 90 m$^3$, the ALICE TPC is the largest TPC of the world. Several gas mixtures have been used in the TPC like Ar-CO$_2$ (88-12), Ne-CO$_2$ (90-10) or Ne-CO$_2$-N$_2$ (90-10-5). The gas system is a closed-loop system where the gas recirculates through the detector with approximately 15 m$^3$ h$^{-1}$. Only a small amount of about 50 L h$^{-1}$ is continuously added to the system. Copper catalyzers are used in order purify the gas by reducing the oxygen as well as the water content. A central HV electrode divides the gas volume in two half volumes of equal size and drift length of 2.5 m. As the TPC is operated with a drift field of 400 V cm$^{-1}$, a potential of $-100$ kV is applied to the central electrode. A field cage composed of several concentric cylinders (inner field cage and outer field cage) is used to form and to improve the homogeneity of the electric drift field. Strips are used to define the electric field as the applied potentials successively decrease from the central HV electrode to nearly ground potential close to the readout chambers. Once a charged particle traverses the active volume of
the TPC it will ionize the gas along its trajectory (black line in Fig. 3.5). The ions drift to the central HV electrode and the electrons to the outer endplates. With the gas choice of Ne-CO$_2$-N$_2$ (90-10-5) for Run 3 the maximum drift time is given by $\approx 100\,\mu$s. A space-resolved readout technology allows to detect the ionization electrons and thus to reconstruct the two-dimensional projection (x-y plane) of the trajectory (green curve). The third and missing z-component can be inferred from the drift time once the drift velocity is known. Therefore an external trigger signal is required to obtain the timing information.

A fundamental requirement for the successful operation of the TPC is a homogeneous electric field in the drift region, as any static or dynamic deteriorations of the electric field might influence the electron drift motion and thus distort the reconstructive capabilities of the detector. Reasons for static non-uniform electric fields are given by mechanical or electrical imperfections in the field cage or readout chambers. To correct for these field deteriorations, an ultraviolet laser calibration system has been implemented in the TPC. A sophisticated optical network consisting of several components (e.g. prism, beam splitter or beam monitors) allows to generate tracks at predefined and well known positions. Spatial deviations of the reconstructed tracks can be traced back to the field deteriorations and corrections can be applied (see [80] for details). An important contribution to dynamic field deteriorations are space-charge distortions caused by ions, as they remain by a factor of $\approx 10^3$ longer in the drift field (cf. Sec. 2.4). Detailed studies about the influence of space-charge distortions and the corresponding corrections can be found in [32, 81, 82].

Commonly the amount of primary ionization electrons is too small to detect at the endplates and an additional amplification is required. Before the ongoing upgrade of the ALICE TPC this was realized by 72 Multiwire Proportional Chambers [27], followed by a segmented pad plane with approximately 570,000 channels (see Fig. 3.6). The primary electrons drift to the anode wires. In the vicinity of the anode wires the electric field strength becomes sufficiently high to induce further ionization processes. The avalanches are capable of creating detectable signals at the readout pads. In order to prevent the
avalanche ions to drift back to the active volume of the detector where they could contribute to unwanted space-charge distortions, a bipolar gating grid (GG) is used. In open mode, the potential $U_{GG}$ of the gating grid is chosen such that it becomes transparent for electrons as well as ions. In closed mode, an alternating potential $U_{GG} \pm \Delta U$ is added to the gating grid wires. For this particular field configuration the gating grid becomes opaque for ions but also for electrons. After a fixed time to collect the ions, the gating grid is opened again. The drift motion of the avalanche ions determines this time interval which limits the theoretical readout rate to the order of 3 kHz. Together with the present readout system, the rate capabilities for central Pb-Pb collisions is limited to 300 Hz. As the interaction rate of the Large Hadron Collider will be increased up to 50 kHz for Run 3, the expected amount of events can not be handled by the current readout technology which demands a complete redesign of the TPC (see Sec. 3.3.1).

### 3.3.1 ALICE TPC upgrade: Quadruple GEM stack

After the Long Shutdown 2 (LS2), the LHC will be operated with an increased luminosity of $\mathcal{L} = 6 \times 10^{32} \text{ cm}^{-2} \text{ s}^{-1}$ which translates to an expected interaction rate of 50 kHz. As the current readout capability of the TPC is limited to 300 Hz, a redesign is required to allow for a continuous and ungated readout of the detector. At the same time the upgraded detector has to preserve the existing performance in
3.3 Time Projection Chamber (TPC)

Figure 3.7: Schematic view (a) and actual image (b) of a ALICE TPC readout chamber (dimensions in mm) \[79\].

In terms of the momentum and dE/dx resolution of about 5.5% and 7% in pp and central Pb-Pb collisions, respectively. Spatial deteriorations due to space-charge distortions must be corrected to the level of the intrinsic track resolution of a few hundred micrometer.

The gated and MWPC-based readout technology will be replaced by a continuous readout technology based on GEM foils. Previous studies showed that a configuration of at least four GEM foils is required to fulfill the aspired goals \[83, 84\]. Within an extensive R&D program, various quadruple GEM stack configurations have been systematically studied in order to find the best stack configuration in terms of energy resolution, ion backflow and reasonable discharge probability. During the upgrade, all MWPC-based readout chambers will be replaced by 18 segments of quadruple GEM stacks on each side of the TPC. Fig. 3.7 shows the schematic view and an image of a single segment: The innermost stack (IROC) is followed by three outer GEM stacks (OROC1, OROC2 and OROC3). Each stack will have a S-LP-LP-S \[1\] configuration. In total 640 GEM foils (plus 100 - 200 spare foils) had to be produced, commissioned and tested in a production time of about 20 - 24 months before they will be mounted and installed during the LS2. A detailed description of the upgrade plan can be found in \[32\] and \[85\].

In order to crosscheck the final detector performance of the upcoming TPC with small prototypes, the main properties - such as the energy resolution - had to be determined by quantities which can be easily measured with test setups in laboratories.

\[1\] S: Standard pitch 140 𝜇m, LP: Large pitch 280 𝜇m
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(a) Simulated dE/dx resolution for MIP tracks as a function of the electron transmission efficiency in the GEM stack for pp and Pb-Pb collisions (errors unknown) [32].

(b) Measured energy resolution with $^{55}$Fe and ion backflow for quadruple S-LP-LP-S stack as a function of the GEM1 potential and different GEM2 settings (errors unknown) [32].

Figure 3.8: Simulation of the dE/dx resolution for MIP tracks and measured energy resolution $\sigma/\mu$ with $^{55}$Fe as well as ion backflow for the quadruple S-LP-LP-S configuration [32].

- **Energy resolution:**
  The energy resolution of a GEM detector depends highly on the (primary) electron transmission through the stack. Fig. 3.8(a) shows the simulated dE/dx resolution of the ALICE stack as a function of the electron transmission efficiency for pp and Pb-Pb collisions. The aspired resolution of about 5.5 % and 7 % in pp and central Pb-Pb collisions translates to a lower limit of about 0.5 for the electron transmission. Lower transmissions will lead to a strong degradation of the resolution.

On the other hand, Fig. 3.8(b) shows the measured energy resolution ($\sigma/\mu$ for $^{55}$Fe) as well as the ion backflow of a quadruple S-LP-LP-S GEM detector for different electric fields and GEM potentials in Ne-CO$_2$-N$_2$ (90-10-5). The voltage of the first GEM stage was increased in a range of $225 \text{ V} \leq U_{\text{GEM1}} \leq 315 \text{ V}$ from left to right. To keep the total gain fixed at 2,000, the potentials $U_{\text{GEM3}}$ and $U_{\text{GEM4}}$ have been tuned while keeping the ratio $U_{\text{GEM3}}/U_{\text{GEM4}}$ constant. The transfer and induction fields are $4 \text{ kV cm}^{-1}$, $2 \text{ kV cm}^{-1}$, $0.1 \text{ kV cm}^{-1}$ and $4 \text{ kV cm}^{-1}$. In the limit of high GEM1 potentials, an energy resolution of approximately 8.5 % can be observed. In [32] it is assumed that this corresponds to an electron transmission of 100 % and a degradation of the measured $^{55}$Fe energy resolution can be expected for an electron transmission of 0.5 (given by the lower limit of the dE/dx resolution in case of pp and central Pb-Pb collisions). Based on Sec. 2.7 the following relation can be obtained

$$\left(\frac{\sigma}{\mu}\right) \propto \frac{1}{\sqrt{N}} \propto \frac{1}{\sqrt{\epsilon}} \tag{3.1}$$

where the electron transmission is given by $\epsilon$. In case of the lower limit of $\epsilon = 0.5$, an upper limit of $\sigma/\mu = 8.5 \% / \sqrt{0.5} \approx 12 \%$ can be expected for the measured $^{55}$Fe energy resolution.

- **Ion backflow:**
  Simulations of the TPC performance [32] indicate that the ion backflow should be lower than 1 % in order to be able to completely correct all effects from space-charge distortions, i.e. the number of back drifting ions per incoming electron $\epsilon$ has to be lower than 20 (at a gain of 2,000).

Fig. 3.9 shows the measured energy resolution ($\sigma/\mu$ for $^{55}$Fe) as a function of the ion backflow in case of...
3.3 Time Projection Chamber (TPC)

### Table 3.1: Foreseen GEM voltages for the ALICE TPC (baseline solution) and optimized settings.

<table>
<thead>
<tr>
<th>Settings</th>
<th>$U_{\text{GEM1}}$</th>
<th>$U_{\text{GEM2}}$</th>
<th>$U_{\text{GEM3}}$</th>
<th>$U_{\text{GEM4}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline [86]</td>
<td>270 V</td>
<td>230 V</td>
<td>288 V</td>
<td>359 V</td>
</tr>
<tr>
<td>Optimized baseline [42]</td>
<td>270 V</td>
<td>230 V</td>
<td>288 V</td>
<td>359 V</td>
</tr>
</tbody>
</table>

### Table 3.2: Foreseen electric field configuration for the ALICE TPC (baseline solution) and optimized settings.

<table>
<thead>
<tr>
<th>Settings</th>
<th>$E_{\text{Drift}}$</th>
<th>$E_{T1}$</th>
<th>$E_{T2}$</th>
<th>$E_{T3}$</th>
<th>$E_{\text{Ind}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline [86]</td>
<td>0.4 kV cm$^{-1}$</td>
<td>4 kV cm$^{-1}$</td>
<td>4 kV cm$^{-1}$</td>
<td>0.1 kV cm$^{-1}$</td>
<td>4 kV cm$^{-1}$</td>
</tr>
<tr>
<td>Optimized baseline [42]</td>
<td>0.4 kV cm$^{-1}$</td>
<td>1.685 kV cm$^{-1}$</td>
<td>0.289 kV cm$^{-1}$</td>
<td>2.912 kV cm$^{-1}$</td>
<td>3.5 kV cm$^{-1}$</td>
</tr>
</tbody>
</table>

the quadruple S-LP-LP-S GEM detector for different electric fields and GEM potentials in Ne-CO$_2$-N$_2$ (90-10-5). The requirements for the upgrade of the ALICE TPC ($\sigma/\mu \leq 12\%$ and $\text{IB} \leq 1\%$) can be fulfilled for some specific configurations, e.g. $U_{\text{GEM3}}/U_{\text{GEM4}} = 0.8$ and $U_{\text{GEM2}} = 235$ V. The final settings to operate the ALICE TPC (also referred to as baseline solution, see Tab. [3.1] and [3.2]) lead to an energy resolution of $\sigma/\mu \approx 12.0\%$ for $^{56}$Fe and an ion backflow of $\approx 0.7\%$ at a gain of 2 000 [86]. Commonly most of the gain is contributed by the last GEM stage as most of the ions will be created there. The following GEM stages for the ions drifting to the cathode can be used in order to suppress the ion backflow. However this stack configuration (optimized to reduce the ion backflow) is in contrast to settings which optimize the discharge probability of GEM stacks where the GEM potentials (and thus the gains) are decreased from the first stage to the readout of the stack [87, 88].

Detailed Magboltz and Garfield++ simulations of GEM stacks indicate that even more optimized field configurations might exist for the ALICE quadruple GEM stack [42]. Within the simulations, the GEM potentials are fixed and set according to the ALICE baseline solution (see Tab. [3.1]), i.e. the absolute gain of each stage remains unchanged. However, the electric fields (see Tab. [3.2]) are varied in order to optimize the electron transfer within the stack and lead to comparable values of the energy resolution at an improved gain of about 12 000. Changes of the ion backflow are ignored as only the electron transfer and thus the effective gain is optimized. Measurements in Ar-CO$_2$ (90-10) are in a good agreement to the simulations [89]. An experimental verification is still open in case of neon-based gas mixtures.

The discharge behavior has been studied during a two-week test beam campaign at the Super Proton Synchrotron (SPS) at CERN. For this purpose a high-intensity pion beam with a momentum of 150 GeV c$^{-1}$ was guided on a 30 – 40 cm thick iron absorber which was followed by an IROC prototype chamber. The total number of accumulated particles can be compared to the expected number of particles during a yearly Pb-Pb run at a collision rate of 50 kHz. The discharge probability is defined as the ratio of the number of detected discharges to the total number of accumulated particles. Therefore all readout pads were connected together and an oscilloscope recorded all signal above a certain threshold which were associated with the occurrence of a discharge. The measured discharge probability of $(6.4 \pm 3.7) \times 10^{-12}$ per incoming hadron is of the same order of magnitude as the discharge probability of the triple GEM detectors at the Large Hadron Collider beauty experiment (LHCb) [85, 90].
Chapter 3 The ALICE experiment

Figure 3.9: Energy resolution and ion backflow of a S-LP-LP-S GEM stack for different electric field configurations and GEM potentials (plot from [32], no error bars given). The aspired goals of the planned ALICE TPC upgrade are indicated by the green box.

3.3.2 Alternative approach: Hybrid Detector

A Time Projection Chamber based on a triple GEM stack was successfully operated by the GEM TPC collaboration for the FOPI spectrometer at the "Helmholtzzentrum für Schwerionenforschung" (GSI) in Darmstadt [83, 91, 92] and with the upgrade of the ALICE TPC, a GEM system has been chosen as the concept was more studied at this time.

A hybrid detector consisting of two GEM foils and a single Micromegas could be an alternative approach to the quadruple GEM stack. Micromegas already offer an intrinsic suppression of the ion backflow in the order of a few percent at relatively high gains $10^3 - 10^4$ [63, 93]. As a consequence less GEM stages are required to block the back drifting ions. The GEM foils are commonly used as pre-amplification stages in order to reduce the needed gain of the Micromegas and to keep the discharge probability at a reasonable level [94–96].

Several 2GEM+MM prototypes have been investigated in terms of discharge behavior during the test beams at the Super Proton Synchrotron (SPS) at CERN. The HIROC detector was constructed using an aluminum body and a pad plane of an IROC. Two standard-pitch GEM foils were followed by a Micromegas with 400 lines per inch and $128\mu m$ pillars attached to the mesh. The active area was similar to the quadruple GEM stack. In addition two identical hybrid detectors were constructed at Yale university but with Micromegas produced using bulk technology. The obtained discharge probabilities are listed in Tab. 3.3 and 3.4 for typical 2GEM+MM settings with low ion backflow and a gain of 2 000. The discharge probabilities are in the region of $10^{-9} - 10^{-7}$ and by orders of magnitudes larger than for the quadruple GEM stack $10^{-10} - 10^{-12}$ [85].

Within the scope of this PhD thesis a hybrid 2GEM+MM detector has been set up and studied in terms of energy resolution and ion backflow with respect to the aspired goals of the upgrade program of the ALICE TPC. A detailed introduction to the investigated setup will be given in Sec. 4.1. Systematic scans of the ion backflow and the energy resolution can be found in Sec. 4.3. The obtained results will be
3.3 Time Projection Chamber (TPC)

### Table 3.3: Discharge probabilities of the HIROC prototype in Ne-CO₂-N₂ (90-10-5).

<table>
<thead>
<tr>
<th>$U_{\text{GEM1}}$ / V</th>
<th>$U_{\text{GEM2}}$ / V</th>
<th>$U_{\text{MM}}$ / V</th>
<th>Gain</th>
<th>Discharge probability</th>
</tr>
</thead>
<tbody>
<tr>
<td>232</td>
<td>210</td>
<td>460</td>
<td>2000</td>
<td>$(3.0 \pm 0.3) \times 10^{-9}$</td>
</tr>
<tr>
<td>250</td>
<td>210</td>
<td>440</td>
<td>1800</td>
<td>$(3.1 \pm 0.7) \times 10^{-10}$</td>
</tr>
<tr>
<td>260</td>
<td>220</td>
<td>420</td>
<td>1600</td>
<td>$(1.5 \pm 1.1) \times 10^{-11}$</td>
</tr>
</tbody>
</table>

Field configurations for all settings are $E_{\text{Drift}} = 390 \text{ V cm}^{-1}$, $E_{T1} = 3875 \text{ V cm}^{-1}$ and $E_{T2} = 150 \text{ V cm}^{-1}$ [85].

### Table 3.4: Discharge probabilities of the Yale prototypes in Ne-CO₂-N₂ (90-10-5).

<table>
<thead>
<tr>
<th>$U_{\text{GEM1}}$ / V</th>
<th>$U_{\text{GEM2}}$ / V</th>
<th>$U_{\text{MM}}$ / V</th>
<th>Gain</th>
<th>Discharge probability</th>
</tr>
</thead>
<tbody>
<tr>
<td>250</td>
<td>210</td>
<td>440</td>
<td>2050</td>
<td>$(2.0 \pm 0.6) \times 10^{-9}$</td>
</tr>
<tr>
<td>260</td>
<td>220</td>
<td>420</td>
<td>2000</td>
<td>$(3.5 \pm 0.6) \times 10^{-10}$</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>420</td>
<td>450</td>
<td>$(1.7 \pm 0.5) \times 10^{-10}$</td>
</tr>
</tbody>
</table>

Field configurations of the first two settings are $E_{\text{Drift}} = 400 \text{ V cm}^{-1}$, $E_{T1} = 3875 \text{ V cm}^{-1}$ and $E_{T2} = 90 \text{ V cm}^{-1}$. For the last settings $E_{\text{Drift}} = 21 \text{ V cm}^{-1}$, $E_{T1} = 3 \text{ V cm}^{-1}$ and $E_{T2} = 2 \text{ V cm}^{-1}$ [85].

compared to results of the Yale university in Sec. 4.3.3. Finally the measurements will be compared to dedicated model calculations which will be part of Sec. 6.
Hybrid detector

4.1 Setup of the hybrid detector

The investigated hybrid detector as shown in Fig. 4.1 consists of two standard-pitch GEM foils (pitch of 140 µm, active area of 10 cm × 10 cm) followed by a bulk Micromegas (cf. Sec. 2.8.2). The foils as well as the Micromegas were manufactured at CERN. One GEM foil is rotated by 90° with respect to the other to prevent unwanted overlapping effects of the holes (see Moiré patterns, Sec. 2.8.1). The investigated Micromegas has an amplification gap of 128 µm, formed by a small mesh (160 lines/cm) and the readout plane (11 cm × 11 cm). The setup has been assembled within a gas-tight vessel with a volume of about 6 liters (see Fig. 4.2(a)). A Kapton window on the lid of the vessel allows for an external irradiation of the drift region with an X-ray tube (Mini-X from Amptek) or an 55Fe source from top through the cathode. A side window can be used in order to irradiate the drift region without passing the cathode foil. The whole setup is placed in a Faraday cage made of copper and lead to reduce electromagnetic influences and to shield the system (see Fig. 4.2(b)).

The detector is flushed using a premixed Ne-CO₂ (90-10) gas bottle (accuracy of CO₂ is (10.09 ± 0.10) %, given by manufacturer Air Liquide). Mass flow controllers allow to operate the setup with gas flows of about 2 – 6 L h⁻¹. The pressure inside the detector, the water as well as the oxygen content can be monitored with a Rapidox 3100ZD device which is connected to the gas output of the vessel.

Measuring currents in the regime of nano- or even picoamperes turns out to be quite sophisticated and requires precise and well calibrated equipment. The currents at the cathode, the different GEM stages or the anode / pad plane can be measured by pA-meters which were developed at the TU Munich [97, 98] and which are sensitive for currents down to the pA level before systematic uncertainties become dominant. They are calibrated in dependence on the temperature using a Keithley pA-meter as reference [99]. The principle of operation is based on the voltage drop over a known shunt resistor. A 16-bit analog-to-digital converter (ADC) is used to digitize the voltage drop which is then converted into a corresponding current. The pA-meters are read out using a wireless connection and send the data to a receiver which is connected to a PC. In a series of N data points, each point can be obtained approximately every 8 s. During this interval, 128 single measurements are taken into account to calculate the average current \( \bar{I}_n \) and the standard deviation \( \sigma_n \). The average current for all data points is then given by

\[
\bar{I} = \frac{1}{N} \sum_{n=1}^{N} I_n
\]  

(4.1)
Chapter 4 Hybrid detector

Figure 4.1: Schematic view of investigated hybrid setup.

and the standard deviation according to

\[(\sigma_I)^2 = \frac{1}{n-1} \sum_{n=1}^{N} (\bar{I} - I_n)^2.\]  

This of course requires a set of \(N\) independent measurements \(\bar{I}_1, ..., \bar{I}_N\) which follow a constant probability density function, e.g. no drift of currents over time. Usually GEM detectors show a drift of the currents once they are set in operation and voltages are applied. Reasons for these effects are for instance charge-up processes of the polyimide layers: Electrons might get stuck to the polyimide in the GEM holes which changes the electrostatic configuration. This on the other hand affects the drift of further charge carriers and the gain until a state of equilibrium has been reached. Dedicated studies have been carried out to understand and to quantify the charge-up effect in measurements and simulations \([100, 101]\), showing that the saturation of the currents and the gains strongly depends on the rate of the irradiation as well as on the water content inside the detector. To compensate for these effects, the hybrid detector has been operated for several hours (usually 2 – 3 hours) until a stabilization of the currents has been observed, followed by the actual measurements for the following investigations.

The readout plane consists of 128 pads in total as shown in Fig. 4.3. Each of the inner quadratic pads has a dimension of 1 cm × 1 cm, the outer pads of 0.5 cm × 1.5 cm. A summation card has been designed in order to sum up the pads as four concentric rings as shown in Fig. 4.3 (inner ring \(R_1\) to outer ring \(R_4\)). The card is directly connected to the pad plane of the Micromegas and allows to use several configuration of the sum (see Tab. 4.1). Channels which are not read out are connected to ground potential. Based on the Micromegas, the segmented pad plane has been reproduced (without the mesh) and can now be used in other setups (like GEM stacks) in combination with the summation card.
4.1 Setup of the hybrid detector

(a) Top view of the hybrid stack inside the vessel: (1) First GEM stage, (2) top cathode, (3) HV connectors, (4) LEMO readout, (5) gas input, (6) gas output.

(b) Vessel housing the hybrid detector with pA-meters on top inside the Faraday cage: (1) pA-meters from Bonn, (2) preamplifier, (3) HV cables, (4) output ring R1, (5) gas input.

Figure 4.2: Assembled and investigated hybrid 2GEM-MM detector opened (a) and mounted inside of the Faraday field cage (b).

(a) Image of the used Micromegas showing the pad plane (total size of 128 pads is 11 cm × 11 cm, without PCB board). The ledge on the left is connected to the mesh.

(b) Schematic view of the readout pads, labeling scheme and summed up rings of pads (inner ring R1 as red, followed by R2 in green, R3 in blue and R4 in gray). Dimensions of pads from ring R1, R2 and R3 are 1 cm × 1 cm. Dimensions of outer pads from ring R4 are 1 cm × 0.5 cm.

Figure 4.3: Pad structure of the used Micromegas.

<table>
<thead>
<tr>
<th>Configuration</th>
<th>Number of pads</th>
<th>Active area</th>
</tr>
</thead>
<tbody>
<tr>
<td>R1</td>
<td>4</td>
<td>2 cm × 2 cm</td>
</tr>
<tr>
<td>R1+R2</td>
<td>36</td>
<td>6 cm × 6 cm</td>
</tr>
<tr>
<td>R1+R2+R3</td>
<td>100</td>
<td>10 cm × 10 cm</td>
</tr>
<tr>
<td>R1+R2+R3+R4</td>
<td>128</td>
<td>11 cm × 11 cm</td>
</tr>
</tbody>
</table>

Table 4.1: Common configurations for the summation card.
4.2 Development of a slow-control system

In order to control and to monitor the hardware components of the setup (like high voltage modules, pA-meters, X-ray devices), a slow-control system has been developed in LabView. As the monitoring of the parameters (e.g. voltages, currents, states of devices) is not time-critical, a fast update of the parameters is of lower priority. The parameters are updated in intervals of a few seconds which makes the system "slow". The developed slow control gives full access to control and operate Micro-Pattern Gaseous Detectors (MPGD) such as GEM stacks or the Hybrid detector. The operation can be fully automated by user-defined script files which can be loaded and executed within the slow control. This allows to run dedicated measurements over long time scales, e.g. ramp to specific voltage settings, start or stop current measurements, record and save MCA spectra and more. Fig. 4.4 shows the graphical user interface. The most important features of the slow control are:

1. Display of channel voltages / currents, ramp states or error states for MPOD high voltage modules.
2. Set trip behavior of MPOD high voltage channels and modules.
3. Support of Zagreb pA-meters "PicoLogic PA125" which are used for ALICE QA measurements.
5. Check interlock of and operate Amptek Miniature X-Ray source (Mini-X).
6. Record and save MCA spectra using Amptek "Pocket MCA MCA8000D".
7. Control stepper motors as X-ray beam shutters.
8. Calculate required potentials based on user-defined stack settings and channel mappings.
9. Load and execute user-defined script files which allow to fully automate the operation of the slow control.
A detailed manual / user guide for the slow control can be found in the appendix [A,3]. The principle of operation and the technical background will be discussed in the following.

The Slow Control has been developed in LabView 2014 and requires a 32-bit architecture in order to work with the Application Programming Interface (API) of the hardware components (MPOD high voltage modules, Amptek Mini-X and Multichannel Analyzer). The Slow Control has been designed as a three-states machine: Once initialized (state 1), the software idles and allows to configure all relevant operational information (e.g. MPOD controller, pA-meter stations, database connections). By connecting to the MPOD crate (state 2), the Graphical User Interface (GUI) is locked. Once connected (state 3), operational panels are activated and the Slow Control enters the main loop which continuously communicates with the hardware components.

Based on the specific IP and the port of the MPOD controller, an ethernet connection is established to the high voltage crate by using the Simple Network Management Protocol (SNMP) and the provided Dynamic Link Libraries (DLL) from Wiener. Within the main routine of the Slow Control, the status of each channel and module is polled every half of a second and user commands are sent (e.g. ramping to specific settings or voltages).

The communication with the Mini-X is based on the provided API by Amptek, allowing to access the USB port and thus the X-Ray device. Once connected, the Slow Control checks if a valid session is running and reads out the Serial number of the device. Within an implemented routine, the status registers and the interlock are checked repeatedly. The readout of the Multichannel Analyzer (MCA) uses a specific API by Amptek which allows to make use of the serial port for data transfer. Here a RS232-USB converter has been used in order to connect the MCA to the USB port instead of the serial port.

The operation of the pA-meters from Bonn is based on a pSQL database (cf. Fig. 4.5). All relevant information of the pA-meters (mean value, error value, mode of operation, battery state, station id and timestamp) are continuously sent to the readout PC via bluetooth and stored to a database as long as the continue flag (CFlag) of a specific device is activated. The Slow Control is capable of reading out the database in order to fetch the requested data. In addition, the continue flag can be set for each specific device, which is then polled by the readout system. The usage of an ethernet connection between the readout system (blue), the pSQL server (gray) and the Slow Control (green) allows remote operation and to run several pA-meters at different places simultaneously.

With the upgrade of the ALICE Time Projection Chamber and the Quality Assurance (QA) program developed in this context, a communication with the pA-meters manufactured in Zagreb (PicoLogic PA125) had to be implemented as well. These pA-meters are optically linked to a common FPGA board which is thereupon connected to a readout PC (see Fig. 4.6). A further LabView program stores the recorded data to the National Instruments Distributed System Manager (NI-DSM). This server can either run locally (on the same machine) or on a remote system and reached via ethernet. The Slow Control connects to this server and obtains the data of the pA-meters. Again, a spatial separation can be realized using ethernet connections.

1 https://www.postgresql.org
2 http://www.picologic.hr
Chapter 4 Hybrid detector

Each user input on the GUI can be automated within the automation panel (see Fig. 4.4). The operations are organized as jobs and processed consecutively “from top to bottom”. Depending on the task, each job (e.g. ramp to specific settings, start current measurements with the pA-meters, record a MCA spectrum or simply wait and idle) requires up to three parameters which contain further information (such as voltages, channels, times). A documentation of all possible commands can be found in the appendix A.3. Script files (text files which contain the jobs line by line) can be loaded inside the Slow Control which allows to program longer measurements. These files can be stored together with the recorded data, allowing to reproduce and comprehend the measuring procedure at later times.

Although a slow control has already been in operation in Bonn[102], a redesign was required since the ALICE collaboration decided to work with pA-meters from Zagreb. The communication with pA-meters in general was not implemented at this time. In addition, the operation was limited to only three GEM foils and further changes (such as a flexible mapping of the used HV channels) were desired. With the flexibility given by the automation scripting and the above-mentioned features, the developed slow control has become the new standard for MPGD operation in the laboratories.

4.3 Characterization of the detector

4.3.1 Gain scans

The properties of the hybrid detector (such as energy resolution, ion backflow and total effective gain of the stack) highly depend on the charge multiplications (the gains) of the individual amplification stages. The knowledge of the effective and the absolute gains is therefore required to fully characterize

Figure 4.5: Communication with the pA-meters from Bonn via pSQL.

Figure 4.6: Communication with the pA-meters from Zagreb via National Instruments Distributed System Manager (NI-DSM).
4.3 Characterization of the detector

Figure 4.7: Procedure to measure the effective as well as absolute gain of a GEM foil. Arrows indicate the movement of the electrons to (a) the top side of the GEM and (b) inside the GEM hole, to the bottom side of the GEM / to the next stage (either a GEM or anode).

Generally the effective / absolute gain of a GEM can be determined as indicated in Fig. 4.7. In case of a low GEM potential ($U_{GEM} = 0$ V) and a vanishing extraction field ($E_{below} = 0$ V cm$^{-1}$), all electrons are expected to end on the top side of the GEM$^3$. This gives access to measure the primary current $I_0$ (Fig. 4.7(a)). Once an appropriate electric potential is applied to the GEM, the electrons will be guided into the holes and start to ionize the gas due to the high electric field. The electrons will then either end on the bottom side of the GEM ($I_{bot} = I_0 \epsilon_c G_{abs} \epsilon_{bot}$) or might get extracted to the next amplification / readout stage ($I_{extr} = I_0 \epsilon_c G_{abs} \epsilon_{e}$). The possible outcomes are quantified by the according transfer efficiencies. The efficiency to end at the bottom side has been introduced as $\epsilon_{bot}$ where the relation $\epsilon_{bot} + \epsilon_e = 1$ holds$^4$. The absolute gain is therefore given by

$$G_{abs} = \frac{1}{\epsilon_c} \left( \frac{I_{bot} + I_{extr}}{I_0} \right).$$

(4.3)

and requires - in addition to the currents - the knowledge of the electron collection efficiency. As a consequence only the product $\epsilon_c G_{abs}$ can be measured by the currents. The effective gain can be obtained by using the relation

$$G_{eff} = \frac{I_{extr}}{I_0}.$$

(4.4)

Commonly the amount of primary charges is too small to induce measurable currents at the top side of the first GEM stage. As a consequence the first GEM stage has been used as a preamplifier and the gain curves have been obtained from the second GEM for the different field configurations. Since both GEM foils are of the same type, the curves can be assumed to hold for the first GEM. A $^{55}$Fe source has been

$^3$ A crosscheck of the currents at the bottom side of the GEM and at the next stage show that a possible reach-through of electrons can be neglected.

$^4$ Possible electron losses at insulators - such as the polyimide layer of the GEM - have been neglected.
placed on the top side of the vessel in order to irradiate and ionize the drift volume of the detector. The effective gain as well as the product of the absolute gain and the electron collection efficiency are shown in Fig. 4.8 in case of Ne-CO_{2} (90-10) and for different field configurations.

The interpretation of the gain curves requires a detailed knowledge of the electron transfer efficiencies (see Sec. 5 for a profound discussion): For low electric fields above a GEM (e.g. TF1 = 0.4 kV cm^{-1}), all field lines from the previous stage enter the GEM hole. As the electrons macroscopically drift according to the electric field lines nearly all electrons are collected, i.e. \( \epsilon_c \approx 1 \). Since \( G_{\text{eff}} \propto \epsilon_e \epsilon_c \), the effective gain curve for TF1 = 0.4 kV cm^{-1} (red points, Fig. 4.8(a)) is higher compared to TF1 = 3.0 kV cm^{-1}. With an increased TF1 = 3.0 kV cm^{-1}, field lines above the GEM start to reach the top side of the GEM and less field lines are entering the hole. This causes a loss of collected electrons and a decrease of the collection efficiency to \( \epsilon_c \approx 0.5 - 0.6 \) (depending on \( U_{\text{GEM}} \)). As a consequence the effective gain drops (black points, Fig. 4.8(a)). The electron extraction efficiency decreases for TF2 = 0.4 kV cm^{-1} which once more causes a drop of the effective gain (blue points, Fig. 4.8(a)). The absolute gain is not affected by TF2 (red and blue points are comparable, Fig. 4.8(b)) since it only depends on the electron collection efficiency (\( G_{\text{abs}} \propto 1/\epsilon_c \), see Eq. 4.3), i.e. the difference between the black and red / blue points in Fig. 4.8(b) is only caused by the collection efficiency of the GEM.

Putting the mesh and the pad structure of the Micromegas on ground potential gives access to measure the primary current \( I_{\text{MM},0} \) at the mesh of the Micromegas. The amplified current \( I_{\text{MM}} \) can thereupon be measured at the readout pads (R1+R2+R3+R4 configuration) once set in operation. As almost all electrons are collected (see Sec. 2.8.2), the gain of the Micromegas is given by \( G_{\text{MM}} = I_{\text{MM}}/I_{\text{MM},0} \). Fig. 4.9 shows the obtained gain curve in Ne-CO_{2} (90-10) for different Micromegas voltages.
4.3 Characterization of the detector

Commonly the GEM stages are operated in a range from 200 – 300 V for the hybrid detector. This translates to effective GEM1 gains of $\approx 10 – 100$ (low collection field $E_{\text{drift}} = 400 \text{ V cm}^{-1}$, high extraction field TF1 = 3 kV cm$^{-1}$) and effective GEM2 gains in a region $\lesssim 10$ (high collection field TF1 = 3 kV cm$^{-1}$, low extraction field TF2 = 80 – 400 V cm$^{-1}$). Voltages like 300 V to 400 V are usually applied to the Micromegas. As a consequence most of the gain is contributed by the Micromegas which operates in a region of $\approx 100 – 1000$. The previous GEM foils are commonly used as preamplification stages and to keep the discharge probability of the Micromegas at a reasonable level.

4.3.2 Energy resolution studies

Influence of the K$\beta$ peak in argon and copper for $^{55}$Fe spectra

The isotope $^{55}$Fe is often used in order to estimate the energy resolution in detector physics. It is unstable and decays by an electron capture to an excited state of $^{55}$Mn*. The dominating photon emissions of $^{55}$Mn* are the K$\alpha_1$, the K$\alpha_2$ and the K$\beta_1$ lines. Both the K$\alpha_1$ line (energy 5 898.8 eV, relative intensity 100 [103]) as well as the K$\alpha_2$ line (energy 5 887.7 eV, relative intensity 50 [103]) have comparable energies which allows to combine them as a single K$\alpha$ line with an energy of 5 895 eV (relative intensity 150). The energy of the K$\beta_1$ line (in the following called K$\beta$) is given by 6 490.4 eV with an relative intensity of 17 [103]. Thus the emission probability of the K$\alpha$ / K$\beta$ line is given by 89.82 % / 10.18 %, i.e. the decay is dominated by the K$\alpha$ line.

Commonly the K$\alpha$ line is referred to obtain the energy resolution as $\sigma/\mu$ by fitting a single Gaussian function to the energy spectrum. However, the energy resolution of gaseous detectors is usually not sufficient to separate the K$\alpha$ and the K$\beta$ lines. Consequently fitting a single Gaussian to the peak will ignore the influence of the K$\beta$ line and leads to an overestimated (higher) energy resolution. The influence
Chapter 4 Hybrid detector

of the $K_\beta$ peak to the simple approach of fitting a single Gaussian will be studied in the following. Furthermore the interaction with copper is taken into account as the cathode and the GEM foils are covered with copper layers. Irradiating the detector from top side might also lead to copper excitation and influence the energy spectrum.

The probability to excite copper or argon has been estimated according to Fig. 4.10. The $K_\alpha$ and the $K_\beta$ lines are assumed to irradiate the detector perpendicularly to the pad plane in this approach. As not all electrons from copper excitations will enter the sensitive drift volume due to their limited range, only electrons emitted close to the surface of the copper layers next to the drift volume are taken into account. Assuming that a photon completely transfers its energy to an electron (most conservative approach) in a copper layer (cathode or GEM1 top electrode), the range is approximately given by $R \approx 0.25 \mu m$ for the $K_\alpha$ as well as for the $K_\beta$ line [104]. Consequently only copper excitations from surface layers with a thickness of $0.25 \mu m$ will be taken into account. While passing through matter, the photon intensity $I$ decreases in a given length $x$ exponentially according to

$$I = I_0 \exp (-\mu \rho x)$$ (4.5)

where $I_0$ is the initial intensity, $\mu$ the total mass absorption coefficient and $\rho$ the density of the passed material. In case of 6 keV photons (for the $K_\alpha$ and the $K_\beta$ lines), the total mass absorption coefficient for copper is given by $\mu_{Cu} = 1.156 \times 10^2 \text{cm}^2 \text{g}^{-1}$ and for argon by $\mu_{Ar} = 2.594 \times 10^2 \text{cm}^2 \text{g}^{-1}$ [38]. Due to this only 61.14% of the initial photons reach the inner boundary of the relevant surface layer of the cathode (dashed line, Fig. 4.10). Once passed through the surface layer of the cathode, only 59.57% of the initial intensity remains, i.e. 1.57% of the photons are lost due to relevant copper excitation in the cathode [5]. Approximately 17.88% of the initial intensity remains after passing through the sensitive drift volume filled with argon which translates to a loss of 41.69% of photons in the drift volume. At the surface of the GEM1 top electrode (neglecting the holes), again 0.46% of photons are lost due to copper excitations. In total 1.57% + 0.46% + 41.69% = 43.72% of the intensity losses are considered to be relevant as the electrons enter the drift volume (copper surfaces and the full drift volume). Only 1.57% + 0.46% = 2.03% are caused by the copper surfaces which translates to an estimated copper-excitation probability of approximately 2.03%/43.72% $\approx 0.05$, i.e. 5%. Consequently about 95% of the relevant excitations are due to argon.

Fig. 4.11 shows the involved energy lines as well as the photo and Auger electrons which have been considered in order to describe the interaction of the $K_\alpha$ line with pure argon as well as with copper (all references are indicated as well). The interaction is assumed to be dominated by argon processes (95%) and copper is only slightly taken into account (5%). Unknown branching ratios are indicated by stars and defined based on equally distributed probabilities or assumptions. As example: The emission of the $K_\alpha$ line (probability of 89.82%) leads to an argon interaction with a probability of 95%. The ionization of the innermost K-shell dominates with 89% and leads to the emission of a photo electron with an energy of 2 689 eV. With a probability of 13.5% the hole of the inner K-shell will be filled with an electron from the L$_{\text{3}}$-shell (emission of $K_{\alpha1}$ photons) or the L$_{\text{2}}$-shell (emission of $K_{\alpha2}$ photons). The absorption length of the $K_{\alpha1}$ and $K_{\alpha2}$ photons in argon usually exceeds the spatial dimension of the conversion volume, i.e. the photons "escape" and cause lower energetic lines in the energy spectrum (also referred to as escape peaks). However with a probability of 86.5% the ionization is followed by inner atomic

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5 An angular distribution of the emitted electrons within the relevant copper layer is not taken into account. All electrons are assumed to enter the drift region which makes this estimation even more conservative.
4.3 Characterization of the detector

Figure 4.10: Estimation of the probability for argon and copper excitations. The K$_{\alpha}$ and the K$_{\beta}$ photons of the $^{55}$Fe source enter the drift volume perpendicularly from above while passing through the copper cathode. Dashed lines indicate the surface layers of the copper electrodes where electrons could enter the drift region from. The ratio of the photon intensity $I/I_0$ decreases exponentially according to Eq. 4.5.

Auger processes, i.e. the K-shell is filled by an electron of higher order but instead of emitted a photon the energy is internally transferred to another electron which finally escapes from the atomic shell. New holes in higher orders appear which again will be filled by the following shells. As example for KL$_{1}$L$_{23}$, an emissions of multiple Auger electrons can be observed: One 2 574 eV, two 203 eV and one 47 eV Auger electron.

As indicated in Fig. 4.11 all reactions are also possible in case of the K$_{\beta}$ line but with shifted energies. The resulting relative intensities (taking the absorption probabilities as well as the emission probabilities into account) and deposited energies are listed in Tab 4.2. Fig. 4.12(a) shows the relative intensity as a function of the number of created electrons (deposited energy / mean ionization energy of pure argon) in case of K$_{\alpha}$ and K$_{\beta}$. The lines with less than 100 electrons are responsible for the argon escape peak which is dominated by the (K$_{\alpha}$ → Ar → K → K$_{\alpha1}$ / K$_{\alpha2}$) branch. The main peak (also photo peak) at approximately 160 electrons is dominated by the (K$_{\alpha}$ → Ar → K → KL$_{23}$L$_{23}$) branch. Each electron in each line experiences a gain which is described by a Polya distribution (cf. Eq. 2.31 here $\bar{n} = 2 000$ and $\theta = 0.5$). The corresponding and smeared distribution is shown in Fig. 4.12(b) and allows to investigate the influence of the K$_{\beta}$ peak. Fitting a single Gaussian function to the tip of the K$_{\alpha}$ peak (blue histogram) yields a resolution of $\sigma/\mu = (6.36 \pm 0.01)$%. Taking the K$_{\beta}$ into account leads for a single Gaussian to a resolution of $\sigma/\mu = (6.62 \pm 0.01)$% (green histogram), i.e. the (real) energy resolution is overestimated by approximately 0.25 % (difference of absolute values) if only a single Gaussian function is fitted to the (measured) peak. The K$_{\beta}$ peak should be taken into account.

Fig. 4.13(a) shows the contributing lines in case 5 % of copper and 95 % of argon excitations are assumed. The resulting spectrum is shown in Fig. 4.13(b) Fitting a single Gaussian function to the
Figure 4.11: Possible emission lines and Auger electron contributions in Ar and Cu for the K\textsubscript{\alpha} and K\textsubscript{\beta} lines of 55Fe. Data from [103, 105–110] as indicated in the figure.

The influence of the background to the recorded MCA spectra of 55Fe in Ne-CO\textsubscript{2} (90-10) will be discussed in the following. A proper description of the spectra includes not only both the K\textsubscript{\alpha} and the K\textsubscript{\beta} lines but also the distribution of the background. A representative spectrum as obtained with the hybrid detector is shown in Fig. 4.14 where the inner ring R1 has been read out. The prominent photo peak at channel 450 is clearly sitting on a broad background distribution which influences the measured energy resolution. Due to different internal excitation processes for neon gas mixtures, no energy lines are emitted which are able to leave the sensitive volume, i.e. no escape peak can be found.

To investigate the different contributions to the background, a 55Fe source has been used in different orientations in order to irradiate the drift region: Collimated from top, uncollimated from top and collimated using the side window. The angular distribution of the emitted radiation (see Fig. 4.15(a)) has been measured using a silicon PIN diode X-ray detector (Amptek XR-100T). The uncollimated
### 4.3 Characterization of the detector

<table>
<thead>
<tr>
<th>Process</th>
<th>rel. intensity / %</th>
<th>Energy / eV</th>
</tr>
</thead>
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<td>2892</td>
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</table>

Table 4.2: Investigated processes which describe the interaction of \(^{55}\)Fe with pure argon (probability of 95 %) and copper (probability of 5 %). Data from [103, 105–110] as indicated in Fig. 4.11.
Chapter 4 Hybrid detector

Figure 4.12: Influence of the $K_\beta$ peak to the $^{55}$Fe spectrum in pure argon gas.

Figure 4.13: Influence of 5% copper excitation to the $^{55}$Fe spectrum in pure argon gas.
Figure 4.14: Example MCA spectrum of $^{55}$Fe in Ne-CO$_2$ (90-10) for the hybrid detector ($E_{\text{Drift}} = 400$ V cm$^{-1}$, $E_{\text{TF1}} = 3$ kV cm$^{-1}$, $E_{\text{TF2}} = 80$ V cm$^{-1}$, $U_{\text{GEM1}} = 256$ V, $U_{\text{GEM2}} = 250$ V and $U_{\text{MM}} = 320$ V).

Figure 4.15: Measurement of the angular rate dependence of the collimated and uncollimated $^{55}$Fe source.
source has an opening angle of approximately 80 – 90° (Fig. 4.15(b)). With the given geometry of the hybrid detector, the pad plane is almost completely irradiated in case of the uncollimated source from top. A hollow cylinder made of aluminum with an inner diameter of 10 mm and a length of 65 mm was used as a collimator. The collimator limits the cone angle to approximately 10 – 15° which corresponds to a drop of the normalized rate from 40 % to 10 %. This reduces the irradiated area at the readout plane to the size of the inner ring R1, i.e. 2 cm × 2 cm. Only a minor fraction of photons converges next to ring R1 (Fig. 4.16). Consequently nearly all photons convert above the central ring R1 and most of the primary charges as well as ionization charges are collected. In case of the uncollimated source, charges will get partially or completely lost as they convert far off the inner ring R1. This causes a lower-energetic tail at the (left) side of the photo peak and will be referred to as sharing.

Fig. 4.17(a) shows the recorded MCA spectra for the different settings of the $^{55}$Fe source. The peaks for channels lower than 200 are caused by conversions within the amplification region of the Micromegas and can only appear for irradiation from top. The position of the photo peak at channel 1 350 is mostly unaffected for the different scenarios. However the contribution of the broad background changes significantly and shows a minimum at channel 600.

The measurements indicate that the background BG is composed of several different contributions: A small sharing contribution $S_0$ due to the remaining irradiation of the pad plane for the collimated source from top, an additional dominating sharing $S$ caused by the broad irradiation of the pad plane in case of the uncollimated source from top (cf. Fig. 4.16) and a long-tailed exponential component called $X$. Placing the uncollimated source on top causes background $X$ as well as both sharing contributions ($BG_{uncoll,top} = S + S_0 + X$). With the collimated source on top the background is given by the background component $X$ and the small sharing component $S_0$ ($BG_{coll,top} = X + S_0$). Irradiating the drift region from the side causes sharing but not background $X$ ($BG_{coll,side} = S + S_0$). The influence of the components can be estimated by looking at the counts within the minimum (Fig. 4.17(b)): Polynomial functions of third order have been fitted to determine the counts within the minimum approximately at channel 600. In case of the uncollimated (collimated) source on top, the minimum is found at approximately $BG_{uncoll,top} = 574 \pm 60$ counts ($BG_{coll,top} = 190 \pm 60$ counts). Irradiating the detector with a collimated source from the side leads to a minimum at approximately $BG_{coll,side} = 389 \pm 50$ counts. With
4.3 Characterization of the detector

(a) Full spectra showing the photo peaks at channel 1350 followed by the broad investigated background. The peaks for channels lower than 200 are caused by conversions within the amplification region of the Micromegas.

(b) Detail view of background for different $^{55}$Fe settings. As a cross check of the background relations (cf. Eq. 4.6) the differences of two MCA spectra (magenta and cyan data points) have been calculated.

Figure 4.17: MCA spectra for different $^{55}$Fe settings.

\[ S = BG_{uncoll, top} - BG_{coll, top}, \quad X = BG_{uncoll, top} - BG_{coll, side} \]
\[ S_0 = BG_{coll, top} + BG_{coll, side} - BG_{uncoll, top}. \]

The background is dominated by the broad sharing component $S$ with $(67 \pm 16)\%$, followed by the contribution of $X$ with $(32 \pm 14)\%$. The sharing component $S_0$ contributes with approximately $(1 \pm 17)\%$.

Consequently the component $S_0$ can be neglected and the following relations should hold for the backgrounds:

\[ BG_{coll, side} = BG_{uncoll, top} - BG_{coll, top}, \]
\[ BG_{coll, top} = BG_{uncoll, top} - BG_{coll, side}. \] (4.6)

Indeed these relations can be observed if the recorded MCA spectra are subtracted from each other (see Fig. 4.17(b)) which motivates the interpretation of the additive background components.

A displacement $\Delta x$ of the collimated $^{55}$Fe source (top side) from the center of the inner ring R1 allows to investigate the contribution of the sharing. Fig. 4.18(a) shows the obtained MCA spectra. The relative change of the sharing contribution can be studied by looking at the minimum of the background distribution at approximately channel 1000. Fig. 4.18(b) shows the relative change of the background amplitude (compared to $\Delta x = 20$ mm) and thus the relative change of the sharing contribution. As expected, the influence of sharing increases as the collimated source is moved from the central position of ring R1. The contribution of sharing increases approximately by a factor of 5 once the collimator has been displaced by $\Delta x = 20$ mm from the center of ring R1.

To investigate the background component $X$ a highly collimated $^{55}$Fe source was placed on the top
Figure 4.18: Contribution of sharing for different displacements of the collimated $^{55}$Fe source. (a) Obtained MCA spectra for different displacements of the collimated $^{55}$Fe source. (b) Relative sharing contribution for different displacements $\Delta x$ (normalized to $\Delta x = 20$ mm). Solid circles indicate the position of the collimator with respect to the inner ring R1. Dotted circles indicate the irradiated area at the anode due to the geometry of the detector and the collimated source (drop of rate to 10 %).

Side of the detector. Here the inner diameter of the collimator was limited to 1 mm in order to completely suppress sharing and to assure that only the inner ring R1 was irradiated. With a reduced rate of about 6 Hz, the MCA spectrum shown in Fig. [4.19] was recorded during 12 hours. The background can be described with a rather broad exponential function and disappears once the detector is detached from the MCA. In this case only the exponential electronic noise for channels lower than 100 remains. The integrated exponential background $X$ leads to a total amount of approximately 21 000 counts (from channel 0 to 2 000). With the given area of the pad plane (ring R1, size 2 cm $\times$ 2 cm) and for 12 hours of data acquisition, this translates to a background intensity of $I_X \approx 7.3 \text{ cm}^{-2} \text{ min}^{-1}$. A comparison to the approximated intensity of vertical muons for horizontal detectors at sea level ($I_\mu \approx 1 \text{ cm}^{-2} \text{ min}^{-1}$ [10]) yields that the background $X$ is not caused by cosmics. The origin of the background $X$ is not completely understood yet. It is assumed that the background might be caused by intrinsic electronic noise of the hybrid detector itself. For the upcoming investigations the background $X$ will be taken into account as a broad exponential function.

A fit model for the energy resolution

Taking all investigations about the energy spectrum of $^{55}$Fe and the background constituents together leads to the following fit model in case of neon gas mixtures:

$$\text{Counts (CH)} = \text{Exp}_{\text{Noise}} + \text{Exp}_X + \text{Sharing} + \text{Gaus}_{K_x}(\sigma_x, \mu_x, A_x) + \text{Gaus}_{K_{\beta}}(\sigma_{\beta}, \mu_{\beta} = \mu_x \cdot 6490/5895, A_{\beta} = A_x / 8.8).$$

(4.7)
4.3 Characterization of the detector

Both the electronic noise as well as the background component X are described by exponential functions. The lower-energetic tail of the photo peak caused by sharing is modeled as a product of a linear function and an error function according to

\[
\text{Sharing} = a \cdot \text{CH} \cdot \text{erfc} \left( \frac{\text{CH} - \mu_\alpha}{b} \right).
\]  

(4.8)

As seen in Fig. 4.17(b), the background distribution first decreases (described by the exponential decay of background X), shows a minimum and finally increases again. The linear increase of the sharing function has been introduced to describe this observed behavior together with the exponential background X. As sharing causes a lower-energetic tail of the photo peak, an error function has been introduced which smoothly suppresses the sharing function for channels exceeding the mean position \( \mu_\alpha \) of the K\( \alpha \) peak. The K\( \alpha \) and the K\( \beta \) peak are described by Gaussian functions. Their mean positions as well as amplitudes are correlated due to the ratios of the emitted energies and the relative intensities. An example MCA spectrum including the individual contributions to the fit model is shown in Fig. 4.20(a). The small bump at channel 50 is caused by photo conversions within the amplification region of the Micromegas and can be neglected as it does not influence the main photo peak. Fig. 4.20(b) shows the energy resolutions for different GEM1 potentials as obtained by fitting a single Gaussian to the photo peak or by applying the full fit model to the spectrum. Increasing the GEM1 potential creates more electrons \( N \) at the first GEM stage. Since \( (\sigma/\mu) \propto 1/\sqrt{N} \) (see Eq. 2.33) this improves the energy resolution as more charges drift to the read out. Fitting a single Gaussian clearly overestimates the energy resolution by approximately 0.5% (difference of absolute values) and leads to larger uncertainties of the fit results. The introduced fit model will be used in all following measurements of the energy resolutions.
4.3.3 Ion backflow and energy resolution scans

Bonn hybrid detector

The hybrid detector has been investigated with respect to the goals of the upgrade of the ALICE Time Projection Chamber (see Sec. 3.3.1), i.e. energy resolution of $\sigma/\mu \leq 12\%$ and ion backflow less than 1% at a total gain of 2000 in Ne-CO$_2$(90-10-5). The setup has been operated using Ne-CO$_2$ (90-10) as the admixture of nitrogen mostly influences the drift velocity $v_{\text{drift}}$ but keeps properties such as diffusion $D_L$, $D_T$ or the number of primary electrons per MIP $N_{\text{total}}$ unaffected (Tab. 4.3). The data will be compared to measurements from Yale university (see Sec. 4.3.3) which have been done in Ne-CO$_2$ (90-10) as well. The following results have been obtained within GEM1-Micromegas scans, i.e. the potential of GEM1 has been scanned while adjusting the potential of the Micromegas in order to keep the total gain constantly at 2000.

To determine the ion backflow IB, a collimated X-ray source (Amptek Mini-X) has been operated (30 kV, emission current of 5 $\mu$A) to irradiate the detector from the top side. All rings have been read out (R1+R2+R3+R4). With the anode current $I_{\text{anode}}$ and the cathode current $I_{\text{cathode}}$, the ion backflow can be
obtained by:
\[ \text{IB} = \frac{|I_{\text{cathode}}|}{I_{\text{anode}}} \]  
(4.9)

The following statistical uncertainties of the ion backflow measurements are determined by the accuracy of the pA-meters, cf. Sec. 4.1.

The total effective gain of the stack has been determined according to Sec. 4.3.1 as the fraction of the current at the readout plane and the initial current within the drift region. Again, the collimated X-ray source has been used to irradiate the full pad plane (R1+R2+R3+R4). Two methods were used in order to measure the primary current: As ions and electrons are created pairwise within the drift region, both the current at the drift cathode \( I_{0,\text{cathode}} \) (due to ions) as well as the current at the top side of the first GEM \( I_{0,\text{G1T}} \) (due to electrons) can be used as initial currents (where \( |I_{0,\text{cathode}}| \approx I_{0,\text{G1T}} \)). For this purpose all GEM and Micromegas potentials were set to ground and only a drift field of \( E_{\text{drift}} = 400 \text{ V cm}^{-1} \) has been applied. Since electrons might also convert in the region between GEM2 and the Micromegas (TF2), an offset current \( I_{\text{anode,MMonly}} \) has been measured at the readout plane. In this case only the Micromegas has been operated and the according transfer field TF2 has been applied. Conversion between GEM1 and GEM2 has been neglected\(^6\). Both methods lead to the following effective gains of the stack
\[ G_{\text{eff,cathode}} = \frac{I_{\text{anode}} - I_{\text{anode,MMonly}}}{|I_{0,\text{cathode}}|}, \]  
(4.10)
\[ G_{\text{eff,G1T}} = \frac{I_{\text{anode}} - I_{\text{anode,MMonly}}}{I_{0,\text{G1T}}}. \]  
(4.11)

Within the GEM1-MM scans, the stack settings have been adjusted in a way to keep the average gain \( G_{\text{eff}} = \left( G_{\text{eff,cathode}} + G_{\text{eff,G1T}} \right) / 2 \) constantly at 2 000. The uncertainties are given by the systematic deviations of both methods to determine the gain as well as the statistical uncertainties of the pA-meter currents.

MCA spectra have been measured using an \(^{55}\text{Fe}\) source (rate of about 40 kHz). Only the inner ring R1 was taken into account in order to reduce the dimension (the capacity) of the pad plane and thus to improve the signal-to-noise ratio (SNR). The energy resolution has been calculated using the advanced fit model from Sec. 4.3.2 as well as - for comparison - based on a single Gaussian function. The obtained results are listed in Tab. 4.4 for different values of TF2. The uncertainties of the energy resolutions \((\sigma/\mu)\) are based on the outcome of the fit results for the mean \( \mu \pm \Delta \mu \) and the standard deviation \( \sigma \pm \Delta \sigma \) (in case of the advanced fit model: mean and sigma of the \( K_\alpha \) peak) and obtained by an error propagation according to \[ \Delta (\sigma/\mu)^2 = (\Delta \sigma)^2/\mu^2 + (\Delta \mu)^2 \sigma^2/\mu^4. \] The environmental conditions (oxygen/water content and overpressure inside the detector, atmospheric pressure, peak temperature of the day as well as relative humidity) during the operation of the setup are listed in Tab. 4.5. As the gain scales with \( p/T \) it is an important quantity which influences the ion backflow and the energy resolution \cite{29}. The relative fluctuations of \( p/T \) (about 2 %) can be neglected as the environmental conditions and the conditions inside the detector were more or less stable during the period of data acquisition.

Fig. 4.21 shows the measured values of the energy resolution, the ion backflow as well as the gain for

\(^6\) The procedure has been proposed and used for an alternative hybrid setup of the Yale university \cite{111,112} and was adopted to allow for a comparison with the Bonn hybrid detector.
the GEM1-Micromegas scans. Increasing the potential (the gain) of GEM1 produces more electrons $N$ and thus improves the energy resolution according to $(\sigma/\mu) \propto 1/\sqrt{N}$ (see Eq. 2.33). At the same time more ions are produced which immediately enter the drift region, i.e. the ion backflow increases. As expected, the gain stays more or less stable during the GEM1-Micromegas scans. Fig. 4.22 shows the energy resolution as a function of the ion backflow for the hybrid detector. The aspired goals of the upgrade of the ALICE TPC are indicated by the red box. The results of the quadruple GEM stack are indicated as well. Since the advanced fit model was not used in case of the quadruple GEM stack, both models (fitting a single Gaussian function to the tip of the photo peak versus advanced fit model) are compared. The investigated hybrid setup consisting of two GEM foils and a single Micromegas is clearly capable of fulfilling the requirements of the upgrade program. Speaking in terms of energy resolution and ion backflow, the hybrid detector clearly competes with the quadruple S-LP-LP-S solution.

Comparing both methods to determine the energy resolution of the hybrid detector (see Tab. 4.4) indicates that even deviations in a range of approximately $1 - 2\%$ (difference of absolute values) are possible ($1.1 - 1.2\%$ for $E_{\text{TF}2} = 80\,\text{V cm}^{-1}$, $1.1 - 2.0\%$ for $E_{\text{TF}2} = 200\,\text{V cm}^{-1}$ and $1.0 - 1.6\%$ for $E_{\text{TF}2} = 400\,\text{V cm}^{-1}$).

With the decision to upgrade the ALICE TPC a system consisting of four GEM foils (S-LP-LP-S configuration) has been chosen as multiple GEM stacks were more studied and understood at this time.

Comparison to Yale measurements

In addition to the research in Bonn, a hybrid detector was investigated in parallel at Yale university by S. Aiola et al. Like the Bonn setup, it consisted of two $10\,\text{cm} \times 10\,\text{cm}$ standard-pitch GEM foils (produced at CERN) and a bulk Micromegas with a mesh of 400 lines per inch. The geometry of the Yale setup was similar to Bonn, i.e. width of TF1 / TF2 approximately 2 mm / 4 mm and a gap of the Micromegas of about 126 $\mu\text{m}$. With 8 mm, only the width of the drift region was chosen smaller (Bonn setup 26 mm). The definition of the ion backflow differs in [113] according to

$$\text{IB}_{\text{Yale}} = \frac{|I_{\text{cathode}}|}{|I_{\text{anode}}|} - \frac{|I_{0,\text{cathode}}|}{|I_{0,\text{anode}}|}$$

(4.12)

where $I_{0,\text{cathode}}$ and $I_{0,\text{anode}}$ are the currents from the initial ionization in the drift gap and from the ionization in the gap of the Micromegas, i.e. the influence of the primary ionization is neglected. To allow for a comparison between the Yale and the Bonn measurements in the following, the backflow was determined according to the definition of Yale. Like in [113], the energy resolutions are obtained by fitting a single Gaussian distribution to the tip of the photo peak for each $^{58}\text{Fe}$ spectrum. Fig. 4.23 shows a GEM1-Micromegas scan from Yale and Bonn in case of comparable settings and for Ne-CO$_2$ (90-10). Indeed both curves are in a good agreement and merge.

Furthermore some specific settings from Yale have been applied to the Bonn setup in order to crosscheck the detector performances in terms of energy resolution, ion backflow and total gain. The results are listed in Tab. 4.6. Equal settings or similar detector performances are indicated by highlighted numbers. In a first step, exactly the same settings have been applied to the Bonn setup (row 1b / 2b). As the detector performance highly depends on the environmental conditions, deviations of the total stack gain can be expected. Commonly the potentials of all stages are scaled in order to tune the total gain of the detector (row 1c / 2c). Additionally only the potentials of GEM1 and the Micromegas have been scaled while...
4.3 Characterization of the detector

<table>
<thead>
<tr>
<th>TF2 / V cm(^{-1})</th>
<th>(U_{\text{GEM1}} / \text{V})</th>
<th>(U_{\text{MM}} / \text{V})</th>
<th>(\sigma/\mu / %) (Adv. fit)</th>
<th>(\sigma/\mu / %) (Single Gauss)</th>
<th>IB / %</th>
<th>Eff. Gain</th>
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</thead>
<tbody>
<tr>
<td>80</td>
<td>274</td>
<td>305</td>
<td>7.72 ± 0.01</td>
<td>8.80 ± 0.01</td>
<td>2.22 ± 0.01</td>
<td>1992 ± 40</td>
</tr>
<tr>
<td>80</td>
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<td>312</td>
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<td>8.95 ± 0.01</td>
<td>1.94 ± 0.05</td>
<td>2000 ± 40</td>
</tr>
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<td>320</td>
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<td>9.17 ± 0.01</td>
<td>1.65 ± 0.06</td>
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</tr>
<tr>
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<td>328</td>
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<td>9.41 ± 0.01</td>
<td>1.4 ± 0.07</td>
<td>2013 ± 40</td>
</tr>
<tr>
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<td>336</td>
<td>8.62 ± 0.01</td>
<td>9.75 ± 0.01</td>
<td>1.2 ± 0.08</td>
<td>1984 ± 40</td>
</tr>
<tr>
<td>80</td>
<td>228</td>
<td>344</td>
<td>8.92 ± 0.01</td>
<td>10.06 ± 0.01</td>
<td>1.02 ± 0.09</td>
<td>2010 ± 40</td>
</tr>
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<td>354</td>
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<td>10.64 ± 0.01</td>
<td>0.85 ± 0.12</td>
<td>1996 ± 40</td>
</tr>
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<td>361</td>
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<td>10.93 ± 0.01</td>
<td>0.74 ± 0.11</td>
<td>2006 ± 40</td>
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<tr>
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<td>294</td>
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<td>9.06 ± 0.01</td>
<td>1.7 ± 0.01</td>
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</tr>
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<td>303</td>
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<td>9.34 ± 0.01</td>
<td>1.46 ± 0.02</td>
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</tr>
<tr>
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<td>311</td>
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<td>2023 ± 154</td>
</tr>
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<td>319</td>
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<td>9.92 ± 0.01</td>
<td>1.07 ± 0.03</td>
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<td>328</td>
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<td>0.91 ± 0.03</td>
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<td>335</td>
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<td>10.73 ± 0.01</td>
<td>0.8 ± 0.04</td>
<td>2012 ± 153</td>
</tr>
<tr>
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<td>342</td>
<td>9.88 ± 0.01</td>
<td>11.11 ± 0.01</td>
<td>0.71 ± 0.04</td>
<td>2007 ± 152</td>
</tr>
<tr>
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<td>350</td>
<td>10.35 ± 0.02</td>
<td>12.31 ± 0.01</td>
<td>0.62 ± 0.04</td>
<td>2012 ± 153</td>
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<td>295</td>
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<td>1.13 ± 0.01</td>
<td>2002 ± 85</td>
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<td>9.26 ± 0.01</td>
<td>10.39 ± 0.01</td>
<td>0.96 ± 0.03</td>
<td>1997 ± 85</td>
</tr>
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<td>400</td>
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<td>10.81 ± 0.01</td>
<td>0.84 ± 0.05</td>
<td>2021 ± 86</td>
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<td>11.29 ± 0.01</td>
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<td>2005 ± 85</td>
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<td>2028 ± 86</td>
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<td>358</td>
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<td>13.32 ± 0.01</td>
<td>0.53 ± 0.06</td>
<td>2027 ± 86</td>
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</table>

Table 4.4: Results of the GEM1-Micromegas scans with the hybrid detector in Ne-CO\(_2\) (90-10) (settings \(E_{\text{drift}} = 400\) V cm\(^{-1}\), \(TF1 = 3\) kV cm\(^{-1}\) and \(U_{\text{GEM2}} = 250\) V). Advanced fit model as well as single Gaussian function has been used in order to determine the energy resolution.

<table>
<thead>
<tr>
<th>TF2 / V cm(^{-1})</th>
<th>Oxygen / ppm</th>
<th>Water / ppm</th>
<th>(p / \text{mbar})</th>
<th>(p_{\text{atmo}} / \text{hPa})</th>
<th>(T_{\text{max}} / ^\circ\text{C})</th>
<th>rH / %</th>
<th>(p/T / \text{hPa K})</th>
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<tr>
<td>80</td>
<td>34</td>
<td>177</td>
<td>26</td>
<td>1027</td>
<td>20.0</td>
<td>53</td>
<td>3.50</td>
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<td>230</td>
<td>30</td>
<td>1030</td>
<td>21.5</td>
<td>59</td>
<td>3.50</td>
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Table 4.5: Environmental conditions and gas properties during the GEM1-Micromegas scans. Environmental data taken from [114] (station "Köln/Bonn"). Dates of measurements are 4/20/2015 for TF2 = 80 V cm\(^{-1}\), 4/15/2015 for TF2 = 200 V cm\(^{-1}\) and 4/21/2015 for TF2 = 400 V cm\(^{-1}\).
Figure 4.21: Results of the GEM1-Micromegas scans with the hybrid detector in Ne-CO₂ (90-10) (settings $E_{\text{drift}} = 400\,\text{V}\,\text{cm}^{-1}$, TF1 = $3\,\text{kV}\,\text{cm}^{-1}$ and $U_{\text{GEM2}} = 250\,\text{V}$). Energy resolution determined using advanced fit model. Errors of energy resolution multiplied by 20.
4.3 Characterization of the detector

Figure 4.22: Energy resolution versus ion backflow for the hybrid detector compared to the quadruple S-LP-LP-S baseline solution for ALICE (data points from [113]). Errors of the energy resolution multiplied by 20 in case of the Bonn hybrid detector and unknown for quadruple GEM stack.

Figure 4.23: Comparison of Bonn hybrid ($U_{\text{GEM2}} = 250$ V and TF2=80 V cm$^{-1}$, uncertainties of the energy resolution multiplied by 20) to Yale hybrid ($U_{\text{GEM2}} = 230$ V and TF2=75 V cm$^{-1}$, uncertainties unknown, see [111]). Common settings are $E_{\text{drift}} = 400$ V cm$^{-1}$ and TF1=3 kV cm$^{-1}$. 
Chapter 4  Hybrid detector

<table>
<thead>
<tr>
<th></th>
<th>$U_{\text{GEM1}}$ / V</th>
<th>$U_{\text{GEM2}}$ / V</th>
<th>$U_{\text{MM}}$ / V</th>
<th>Total gain</th>
<th>IB$_{\text{Yale}}$ / %</th>
<th>$\sigma/\mu$ / % (Single Gauss)</th>
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<tr>
<td>Yale</td>
<td>1a 195</td>
<td>210</td>
<td>400</td>
<td>2133</td>
<td>0.350</td>
<td>12.25</td>
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<tr>
<td>Bonn</td>
<td>1b 195</td>
<td>210</td>
<td>400</td>
<td>1561 ± 53</td>
<td>0.45 ± 0.02</td>
<td>13.62 ± 0.02</td>
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<td></td>
<td>1c 200</td>
<td>214</td>
<td>405</td>
<td>2151 ± 73</td>
<td>0.367 ± 0.001</td>
<td>13.00 ± 0.02</td>
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<td>1d 202</td>
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<td>407</td>
<td>2142 ± 72</td>
<td>0.369 ± 0.001</td>
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<td>Yale</td>
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<td>2160</td>
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<tr>
<td>Bonn</td>
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<td>185</td>
<td>420</td>
<td>1338 ± 45</td>
<td>0.39 ± 0.04</td>
<td>16.13 ± 0.02</td>
</tr>
<tr>
<td></td>
<td>2c 205</td>
<td>194</td>
<td>421</td>
<td>2138 ± 72</td>
<td>0.53 ± 0.02</td>
<td>14.25 ± 0.02</td>
</tr>
<tr>
<td></td>
<td>2d 206</td>
<td>185</td>
<td>430</td>
<td>2157 ± 73</td>
<td>0.30 ± 0.08</td>
<td>14.44 ± 0.02</td>
</tr>
</tbody>
</table>

Table 4.6: Comparison between Bonn (water content $\approx$ 160 ppm, oxygen content $\approx$ 30 ppm, overpressure $\approx$ 30 mbar) and Yale setup (water content $<$ 200 ppm, oxygen content $<$ 30 ppm) for some selected Yale settings [111]. Highlighted numbers indicate equal settings or similar detector performances.

fixing the potential of GEM2 (row 1d / 2d).

Keeping the potential of GEM2 constant while adjusting the total gain with GEM1 and the Micromegas (1d / 2d) gives best results in the reproduction of the Yale measurements. In fact specific ion backflow values can be reproduced within the limits of the measurements (2d) or turn out to be comparable (1d). However, the obtained energy resolutions deviate by approximately 1% (difference of absolute values) and the obtained Bonn values yield a slightly increased / worse energy resolution. The operation of MPGD and especially an appropriate determination of the energy resolution is a sophisticated topic and turns out to be strongly influenced by the operational conditions. Factors like environmental conditions or geometric manufacturing tolerances of the GEM foils bias the energy resolution and could cause deviations. As the conditions (such as temperature, atmospheric pressure, geometric tolerances) are unknown in case of the Yale measurements, deviations can be expected. Nevertheless both detectors lead to comparable measurements and do not indicate major inconsistent behaviors. Within the limits of the given measurable accuracy and the environmental uncertainties, trends of the detector performances can be reproduced.
A model for charge transfer in GEM foils

5.1 Charge transfer in GEM foils

The ion backflow, the energy resolution as well as the gain of GEM stacks highly depend on the charge transfer processes between the different GEM stages. Thus a good understanding of the transfer processes is mandatory. Commonly these processes can be described using the collection and extraction efficiency for the charge carriers of a GEM. The charge transfer processes can be characterized as follows:

- The electron collection efficiency ($\epsilon_{\text{coll}}$) is defined as the ratio between the number of electrons entering the holes (i.e. number $N_{\text{coll}}$ of electrons which are not blocked by the top electrode of the GEM) and the number of electrons approaching the GEM foil ($N_0$).

$$\epsilon_{\text{coll}} = \frac{N_{\text{coll}}}{N_0}$$ (5.1)

- The electron extraction efficiency ($\epsilon_{\text{extr}}$) is defined as the ratio between the number of electrons extracted from the holes (i.e. number $N_{\text{extr}}$ of electrons reaching the next amplification stage / readout plane) and the total number of electrons available in the holes after the multiplication ($N_{\text{in}}$).

$$\epsilon_{\text{extr}} = \frac{N_{\text{extr}}}{N_{\text{in}}}$$ (5.2)

Generally the average (macroscopic) movement of the charge carriers is defined by the electric and the magnetic field according to the Langevin equation (Eq. 2.20). In this approach no magnetic field has been taken into account and the efficiencies will be modeled by calculating the electric field inside a GEM hole. Fig. 5.1 shows the electric field lines inside a GEM hole for a standard-pitch GEM based on the calculations of the electric fields which will part in the following chapter. The field lines have been calculated for different ratios of the electric fields above / below the GEM ($E_{\text{above}}$ and $E_{\text{below}}$) and inside the GEM hole $E_{\text{GEM}}$. Field lines with common origin and ending positions are indicated by the same color. Fig. 5.1(a) shows field lines originating from the cathode for a high field above the GEM, i.e. some field lines end on the top side of the GEM (magenta). The remaining field lines (yellow) enter and leave the hole at the bottom side of the GEM. For a low field above the GEM, no field lines from the cathode end on the top side of the GEM (Fig. 5.1(b)). In this case only field lines originate from [5.1.3]

1 A detailed discussion about the ratios and the calculations of the electric fields will be part of Sec. 5.2.3.
Chapter 5 A model for charge transfer in GEM foils

the top side of the GEM (Fig. 5.1(c)). While some of the field lines manage to reach the anode (green), some of them may end at the bottom side of the GEM (orange). If the field below the GEM is sufficiently high (Fig. 5.1(d)), field lines will additionally start at the bottom side and end at the anode (blue).

All these common field lines can be combined to several electric fluxes within a GEM hole. Generally the electric flux \( \phi \) through a certain region (or surface \( A \) with orthogonal normal vector \( \vec{n} \)) is defined as the surface integral of the electric field \( \vec{E} \) according to

\[
\phi = \int_A \vec{E} \cdot d\vec{A}
\]

where \( d\vec{A} = \vec{n} \, dA \). The collection as well as the extraction efficiency that have been defined above in terms of number of electrons can alternatively be defined as a function of the electric fluxes inside a GEM. This has been studied by Killenberg et al. [115]. A schematic view of all common field lines from Fig. 5.1 is shown in Fig. 5.2 as electric fluxes. The electron collection efficiency can be defined as the part \( D \) of the total flux \( \phi_{\text{cath}} \) which enters the GEM hole. The extraction efficiency of a GEM is defined as the ratio of the flux through the hole which reaches the anode (given by the sum of flux \( D \) and the ring-shaped flux \( R \), i.e. \( D + R \)) compared to the total flux \( \phi_{\text{hole}} \) through the hole. In [115] Killenberg gets the electric flux by performing a numerical simulation using the finite element program Maxwell 3D. It allows to solve Maxwell’s equations for a three-dimensional problem. The obtained results are thereupon parameterized in order to describe the collection and extraction efficiency. The parametrization is later used to predict the transfer properties of a GEM. An approach that is based on the same methodology can be found by Bencivenni et al. in [116]. Also here a parametrization of the efficiencies has been used to predict measurements of the transfer processes. However, both parameterizations do not fully explain the behavior of three different regions that can be clearly seen in the data. In addition measurements and simulations reveal that the extraction efficiency for electrons has an offset that is not included in the parametrization.

Based on the following electric flux calculations, analytic expressions can be found for the collection and extraction efficiency by modeling a GEM as a two–dimensional charge density distribution. Effects like diffusion and charge carrier losses caused by attachment are not taken into account. The obtained results inherently include an offset for the extraction efficiency and predict the appropriate limits for a GEM.

5.2 Efficiency calculations based on an analytic two-dimensional model

5.2.1 Charge-density model

The electric fluxes and thus the efficiencies are calculated analytically based on a simplified two-dimensional cut through the hexagonal GEM structure. The top and the bottom side of the GEM are modeled by metal electrodes carrying constant electric line charges. A central assumption in this work are the constant charge density distributions which cause nonconstant potentials on the electrodes. The dielectric polyimide layer between the top and the bottom side is ignored, i.e. a differentiation between an inner and an outer hole diameter is not considered. Based on the charge density \( \rho \) for this configuration the electric potential can be obtained by solving Poisson’s equation \( \Delta V = -\rho/\epsilon_0 \). The electric field is thereupon given by \( \vec{E} = -\nabla V \). The electric fluxes can then be calculated according to Eq. 5.3, which
5.2 Efficiency calculations based on an analytic two-dimensional model

Figure 5.1: Different field lines inside a GEM hole for different field ratios. In this two–dimensional calculation of the electric field only half of the GEM hole is shown. Same colors indicate common start and end positions of the field lines and group them. Yellow (region D): Cathode to anode. Magenta (region φC): Cathode to top side of the GEM. Green (region R): Top side of the GEM to anode. Orange (region Z): Top side of the GEM to the bottom side of the GEM. Blue (region φA): Bottom side of the GEM to anode.
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Figure 5.2: Separation of the different electric fluxes in a GEM hole based on [115].

Figure 5.3: Two-dimensional charge-density model of a GEM foil for $N=3$. The inner electrodes of the GEM top / bottom side are indicated as black lines. Outer electrodes indicated as gray lines.

finally leads to the efficiencies for a GEM.

As shown in Fig. [5.3] the GEM geometry is described by the parameters $w$ and $L$ that are related to the pitch by $p = (w + L)/2$. The thickness of the GEM (distance between top and bottom electrode) is described by the parameter $d$. As the transfer processes in GEM foils depend on the external field configuration, a cathode is placed at distance $g_1/2$ and an anode at $-g_2/2$. The line charge densities at the cathode $\mu_1$ and at the top side of the GEM $-\mu_1$ describe the electric collection field $E_1$. Accordingly, the line charge densities $\mu_2$ and $-\mu_2$ are used in order to express the extraction field $E_2$. The electric field within the GEM hole is described by the line charge density $\lambda$. The charge density for a GEM with

---

2 The division by a factor of 2 was historically introduced since the electrodes of the GEM are located at $y = \pm d/2$ and is only of comfortable reason for the calculations.
2N − 1 holes \((N \in \mathbb{N} \text{ and } N \geq 1)\) can be written as

\[
\rho_N = (\lambda - \mu_1) \cdot \delta \left( y - \frac{d}{2} \right) \cdot \left[ \sum_{n=0}^{N-2} f_n(x, L, w) + \Theta_N^+(x, L, w) + \Theta_N^-(x, L, w) \right] \\
+ (\mu_2 - \lambda) \cdot \delta \left( y + \frac{d}{2} \right) \cdot \left[ \sum_{n=0}^{N-2} f_n(x, L, w) + \Theta_N^+(x, L, w) + \Theta_N^-(x, L, w) \right] \\
+ \mu_1 \cdot \delta \left( y - \frac{g_1}{2} \right) - \mu_2 \cdot \delta \left( y + \frac{g_2}{2} \right)
\]

(5.4)

with

\[
f_n(x, L, w) = \Theta_{n,0} \left( x + \frac{nL + (n + 1)w}{2} \right) \cdot \Theta_{n,1} \left( -x - \frac{(n + 1)L + nw}{2} \right) \\
+ \Theta_{n,2} \left( x - \frac{(n + 1)L + nw}{2} \right) \cdot \Theta_{n,3} \left( -x + \frac{nL + (n + 1)w}{2} \right).
\]

(5.5)

\[
\Theta_N^+(x, L, w) = \Theta \left( \pm x - \frac{NL + (N - 1)w}{2} \right).
\]

(5.6)

Within the calculations the Dirac delta function \(\delta\) and the Heaviside step function \(\Theta\) are used to express the positions of the GEM electrodes as well as the anode and the cathode. The first two terms in Eq. (5.4) describe the charge densities at the top and at the bottom side of the GEM: Each outer electrode of the GEM (see Fig. 5.3) is described by a single Heaviside function \(\Theta_N^\pm\) as indicated in Fig. 5.4(a). Each inner electrode is described by the product of two Heaviside functions \(f_n\) as indicated in Fig. 5.4(b). The last two terms describe the charge densities at the cathode and the anode.

With the introduced expressions \((\gamma_1, \gamma_2, \delta_1, \delta_2)\), Eq. (5.4) can be rewritten in a more compact form:

\[
\rho_N = (\gamma_1 + \gamma_2) \left( \sum_{n=0}^{N-2} f_n \right) + (\gamma_1 + \gamma_2) (\Theta_N^+ + \Theta_N^-) + \mu_1 \delta_1 - \mu_2 \delta_2.
\]

(5.7)

### 5.2.2 Calculation of potential and electric fields

To obtain the electric potential \(V\) and finally the electric field for this specific charge distribution, Poisson’s equation \(\Delta V_N = -\rho_N/\epsilon_0\) has to be solved. For an infinite volume in two-dimensional Cartesian coordinates the solution of Poisson’s equation can be constructed with the corresponding Green’s function \(G_{\Delta 2D} = \ln(r)/2\pi\) where \(r = |\vec{r} - \vec{r}'| = \sqrt{(x - x')^2 + (y - y')^2}\) [117):

\[
V_N = -\frac{1}{2\pi\epsilon_0} \int \rho_N(\vec{r}') \ln|\vec{r} - \vec{r}'| \, d\vec{r}'.
\]

(5.8)
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Figure 5.4: Introduced Heaviside functions which describe the electrodes of the GEM (example for \( N = 3 \), hole diameter \( L = 50 \mu m \) and pitch \( p = 140 \mu m \)).

Together with the charge density (Eq. [5.7]) the electric potential can be written as

\[
V_N = - \frac{1}{2\pi\epsilon_0} \sum_{n=0}^{N-2} \int \int_{=h_n^{(1)}(x,y)} (\gamma_1 + \gamma_2) f_n \ln r \, dx' \, dy' \\
- \frac{1}{2\pi\epsilon_0} \int \int_{=G_N^{(1)}(x,y)} [(\gamma_1 + \gamma_2)(\Theta_N + \Theta_N) + \mu_1 \delta_1 - \mu_2 \delta_2] \ln r \, dx' \, dy'.
\]  

(5.9)

Based on the potential \( V_N \), the electric field can be determined by \( \vec{E} = -\nabla V_N \) which finally leads to

\[
E_N^x = - \frac{\partial}{\partial x} V_N = - \frac{1}{2\pi\epsilon_0} \left[ \sum_{n=0}^{N-2} \frac{\partial h_n^{(1)}}{\partial x} + \frac{\partial G_N^{(1)}}{\partial x} \right],
\]

(5.10)

\[
E_N^y = - \frac{\partial}{\partial y} V_N = - \frac{1}{2\pi\epsilon_0} \left[ \sum_{n=0}^{N-2} \frac{\partial h_n^{(1)}}{\partial y} + \frac{\partial G_N^{(1)}}{\partial y} \right].
\]

(5.11)

While the \( G_N^{(1)} \) term describes the central hole of the GEM, the electric field is corrected for additional outer holes by the \( h_n^{(1)} \) terms. Especially for \( N = 1 \) (only one hole) the expressions for the electric field components \( E_x \) and \( E_y \) become simple enough allowing to calculate the average electric field inside the central GEM hole and in the regions above and below the GEM.

According to Eq. [5.12] the average electric field inside the GEM hole can be estimated by an integral of
5.2 Efficiency calculations based on an analytic two-dimensional model

The deviation factor $\kappa_m$ as a function of the order $m$ of the Taylor expansion for $d \approx L$ without an external field ($\mu_1 = \mu_2 = 0$) and only for one central GEM hole ($N = 1$).

\begin{figure}[h]
\centering
\includegraphics[width=0.8\textwidth]{figure5.5.png}
\caption{The deviation factor $\kappa_m$ as a function of the order $m$ of the Taylor expansion for $d \approx L$ without an external field ($\mu_1 = \mu_2 = 0$) and only for one central GEM hole ($N = 1$).}
\end{figure}

the $m$-th Taylor expansion of the absolute electric field over the central hole.

$$\langle E_{\text{GEM}} \rangle_m = \frac{1}{dL} \int_{-\frac{L}{2}}^{\frac{L}{2}} \int_{-\frac{L}{2}}^{\frac{L}{2}} \sum_{m_1=0}^{m} \sum_{m_2=0}^{m} \frac{x^{m_1} y^{m_2}}{m_1! m_2!} \left( E_x^2 + E_y^2 \right) \ dx \ dy$$

Without an external field ($\mu_1 = \mu_2 = 0$) and for an arbitrary order $m$ the average electric field inside the GEM hole ($N = 1$) can be written as

$$\langle E_{\text{GEM}} \rangle_m = \frac{\lambda}{\epsilon_0} \kappa_m .$$

Similar calculations for an infinite parallel plate capacitor lead to an average electric field of $\lambda/\epsilon_0$ (which is equal to the limit of $L \to 0$ for a GEM, cf. Fig. 5.6). Thus the expression $\lambda/\epsilon_0$ in Eq. 5.13 can be identified as the electric field of a parallel plate capacitor $U/d$. The factor $\kappa_m$ describes the deviation from the average GEM field to the field of a parallel plate capacitor. Accordingly the average electric field inside a GEM hole can be written as:

$$\langle E_{\text{GEM}} \rangle_m = E_{\text{GEM}}^\parallel \kappa_m = \frac{U}{d} \kappa_m .$$

Some first order values for $\kappa_m$ are given by (only odd orders contribute)

$$\kappa_1 = 1 - \frac{2}{\pi} \arctan \frac{L}{d} ,$$

$$\kappa_3 = 1 - \frac{2}{\pi} \arctan \frac{L}{d} + \frac{2}{3\pi} \frac{dL^3 + Ld^3}{(L^2 + d^2)^2} .$$

In case of $d \approx L$ the factor $\kappa_m$ becomes independent of the GEM geometry. Numerical values for higher orders of $m$ are given in Fig. 5.5.

With external fields ($\mu_1 \neq 0, \mu_2 \neq 0$ and for $m = 1$) the average electric field inside of the GEM
hole is given by \((N = 1, \kappa_1 \geq 0)\)

\[
\langle E_{\text{GEM}} \rangle_1 = \frac{1}{\epsilon_0} \left[ -\lambda \left( 1 - \frac{2}{\pi} \arctan \frac{L}{d} \right) - \frac{\mu_1}{\pi} \arctan \frac{L}{d} - \frac{\mu_2}{\pi} \arctan \frac{L}{d} \right] \\
= \frac{\lambda}{\epsilon_0} \left( \kappa_1 + \frac{1}{2} \left( 1 - \kappa_1 \right) (\eta_1 + \eta_2) \right). 
\tag{5.17}
\]

In case of \(d \approx L\) (as an approximation for a GEM) Eq. 5.17 turns into

\[
\langle E_{\text{GEM}} \rangle_1 = \frac{1}{2} E_{\text{GEM}}^{\parallel} + \frac{1}{4} E_{\text{above}}^{\parallel} + \frac{1}{4} E_{\text{below}}^{\parallel} 
\tag{5.18}
\]

where the expressions \(\mu_1/\epsilon_0\) and \(\mu_2/\epsilon_0\) have been identified as the (parallel plate) electric fields above and below the GEM. The average electric field inside the GEM hole is dominated by the parallel plate term \(E_{\text{GEM}}^{\parallel}\) (50%) and both external fields couple in with a factor of \(1/4\) (25%). For more realistic values \((d = 50 \mu m\) and \(L = 70 \mu m\)) the deviation factor is given by \(\kappa_1 \approx 0.4\), i.e. \(E_{\text{GEM}}^{\parallel}\) contributes with 40% to the average electric field inside a GEM hole. Both external fields couple in with approximately 30%. The average field above the GEM can be approximated by a Taylor expansion at \(x = 0\) and \(y = (d + g_1)/4\) (central position between cathode and top electrode of the GEM) similar to Eq. 5.12. In this case the integration limits are given by \(-p/2 \leq x \leq p/2\) and \(d/2 \leq y \leq g_1/2\). For \(N = 1\) and the first order \(m = 1\) the average electric field above the GEM is given by

\[
\langle E_{\text{above}} \rangle_1 = \frac{1}{\epsilon_0} \left[ -\lambda D_1(d, L, g_1) - \mu_1 D_2(d, L, g_1) - \mu_2 D_3(d, L, g_1) \right] 
\tag{5.19}
\]
where the following coefficients have been introduced

\[
D_1(d, L, g) = -\frac{1}{\pi} \left[ \arctan \left( \frac{2L}{d-g} \right) + \arctan \left( \frac{2L}{3d+g} \right) \right], \tag{5.20}
\]

\[
D_2(d, L, g) = 1 + \frac{1}{\pi} \arctan \left( \frac{2L}{d-g} \right), \tag{5.21}
\]

\[
D_3(d, L, g) = \frac{1}{\pi} \arctan \left( \frac{2L}{3d+g} \right). \tag{5.22}
\]

Accordingly the average field below the GEM can be obtained by a Taylor expansion at \(x = 0\) and \(y = -(d + g_2)/4\). The integration limits are \(-p/2 \leq x \leq p/2\) and \(-g_2/2 \leq y \leq -d/2\). For \(N = 1\) and \(m = 1\) the field below the GEM is given by

\[
\langle E_{\text{below}} \rangle_1 = \frac{1}{\varepsilon_0} \left[ -\lambda D_1(d, L, g_2) - \mu_1 D_3(d, L, g_2) - \mu_2 D_2(d, L, g_2) \right]. \tag{5.23}
\]

Obviously Eq. 5.24 holds for the coefficients \(D_1, D_2\) and \(D_3\), i.e. all contributions of the charge densities \(\lambda, \mu_1\) and \(\mu_2\) add up to 100%.

\[
D_1 + D_2 + D_3 = 1 \tag{5.24}
\]

Commonly the distance of the cathode / the anode to the GEM is much higher than the thickness or the diameter of the GEM hole, i.e. \(g \gg d\) and \(g \gg L\). For this limit the coefficients \(D_1\) and \(D_3\) vanish and the coefficient \(D_2 \to 1\) dominates. As example for a standard-pitch GEM \((d = 55 \mu m, L = 70 \mu m)\) and \(g_1 = g_2 = 2000 \mu m\) the coefficients are \(D_1 = 0.0023, D_2 = 0.9771\) and \(D_3 = 0.0206\). Thus the average external electric fields can be approximated by

\[
\langle E_{\text{above}} \rangle \approx \frac{\mu_1}{\varepsilon_0} = E_{\text{||}}^{\text{above}}, \tag{5.25}
\]

\[
\langle E_{\text{below}} \rangle \approx \frac{\mu_2}{\varepsilon_0} = E_{\text{||}}^{\text{below}}. \tag{5.26}
\]

Consequently the average electric field above or below a GEM (e.g. transfer field or induction field) can be well approximated as the field of a parallel plate capacitor. The field inside the GEM requires an additional correction factor \(\kappa\) which describes the deviation to a parallel plate capacitor.

Fig. 5.7 shows the electric field inside the central GEM hole for different numbers of total holes and charge density configurations. For low \(N\) an asymmetry of the electric field at the anode and cathode becomes dominant and field lines from neighboring GEM unit cells can enter each other. As example in Fig. 5.7(c) the field lines at the left bottom side of the GEM unit cell enter the neighboring cell. This effect can be compensated for higher values of \(N\) (cf. Fig. 5.7(b)). Indeed this effect biases the efficiencies but disappears for \(N \to \infty\) (as shown in Sec. 5.2.5). A first impression of the charge transfer efficiencies can qualitatively be seen in Fig. 5.7. For a low collection field \(E_1\) (Fig. 5.7(a)) more field lines from the cathode are entering the GEM hole as compared to a high collection field (Fig. 5.7(b)). Here a large part of the field lines ends on the top side of the GEM. Based on this the collection of electrons can be expected to be higher for a low collection field which finally decreases for an increasing collection field. Similarly, the extraction of electrons below the GEM can be expected to be higher for high extraction fields and lower for low extraction fields.
5.2.3 Charge density ratios and electric field ratios

In the following the efficiencies will be expressed in unitless ratios of the charge densities (external charge density / GEM charge density), i.e. \( \eta_1 := \frac{\mu_1}{\lambda} \) and \( \eta_2 := \frac{\mu_2}{\lambda} \). Based on the previous investigations the charge density ratios can be expressed as functions of the average electric fields and parallel plate fields. With Eq. 5.17 and Eq. 5.25 the following ratio can be obtained:

\[
\frac{\langle E_{\text{above}} \rangle}{\langle E_{\text{GEM}} \rangle} = \frac{\eta_1}{\kappa_1 + \frac{1}{2} (1 - \kappa_1) (\eta_1 + \eta_2)}.
\]

(5.27)

The ratio of both external fields is given by

\[
\frac{\langle E_{\text{above}} \rangle}{\langle E_{\text{below}} \rangle} = \frac{\eta_1}{\eta_2}.
\]

(5.28)

By combining Eq. 5.27 and Eq. 5.28 the charge density ratios can be written as

\[
\eta_1 = \frac{\kappa_1 \langle E_{\text{above}} \rangle}{\langle E_{\text{GEM}} \rangle - \frac{1}{2} (1 - \kappa_1) (\langle E_{\text{above}} \rangle + \langle E_{\text{below}} \rangle)},
\]

(5.29)

\[
\eta_2 = \frac{\kappa_1 \langle E_{\text{below}} \rangle}{\langle E_{\text{GEM}} \rangle - \frac{1}{2} (1 - \kappa_1) (\langle E_{\text{above}} \rangle + \langle E_{\text{below}} \rangle)}.
\]

(5.30)

Commonly the field inside the GEM hole is much higher than the applied external fields, i.e. \( \langle E_{\text{GEM}} \rangle \gg \langle E_{\text{above}} \rangle \) and \( \langle E_{\text{GEM}} \rangle \gg \langle E_{\text{below}} \rangle \). In this case the charge density ratios can be written as

\[
\eta_1 \approx \kappa_1 \frac{\langle E_{\text{above}} \rangle}{\langle E_{\text{GEM}} \rangle},
\]

(5.31)

\[
\eta_2 \approx \kappa_1 \frac{\langle E_{\text{below}} \rangle}{\langle E_{\text{GEM}} \rangle}.
\]

(5.32)
By using Eq. 5.17, 5.25 and 5.26 the charge density ratios (Eq. 5.29 and 5.30) can be expressed as functions of the parallel external fields

\[ \eta_1 = \frac{E_\text{above}}{E_{\text{GEM}}} \]  
\[ \eta_2 = \frac{E_\text{below}}{E_{\text{GEM}}} \]  

### 5.2.4 Flux calculations and efficiencies

As shown in Fig. 5.2, the field inside a GEM can be separated into several fluxes [115]. The collection efficiency (Eq. 5.35) can be expressed as the part \( D \) of the total cathode flux \( \phi_{\text{cath}} \) which enters the GEM hole. For common GEM settings all field lines originating from the cathode point to the GEM foil. Depending on the field configuration, some of them might end on the top electrodes of the GEM (flux \( \phi_C \)). These field lines point downwards (i.e. negative y-component) at the top side of the GEM. The flux \( \phi_C \) is adjoined by the flux \( R \). At the top side of the GEM, the field lines from flux \( R \) point upwards (i.e. positive y-component). By finding the point on the top side of the GEM where the electric field in y-direction \( E_y \) changes the sign gives access to calculate \( \phi_C \) and finally the collection efficiency. The flux \( \phi_C \) can be calculated as it corresponds to all field lines which end on the top side of the GEM (point downwards, i.e. negative y-component) and which is adjoined by the flux \( R \) which field lines originate from the GEM (point upwards, i.e. positive y-component).

\[ \epsilon_{\text{coll}} = \frac{D}{\phi_{\text{cath}}} = \frac{\phi_{\text{cath}} - 2\phi_C}{\phi_{\text{cath}}} \]  
\[ \epsilon_{\text{extr}} = \frac{D + 2R}{\phi_{\text{hole}}} = \frac{\phi_{\text{anode}} - 2\phi_A}{\phi_{\text{hole}}} \]  

The extraction efficiency (Eq. 5.36) of a GEM is defined as the flux \( D + 2R \) through the hole \( \phi_{\text{hole}} \) which manages to reach the anode. The flux \( D + 2R \) can be expressed by the flux \( \phi_A \) which again can be found by a flip of the electric field in y-direction at the bottom side of the GEM.

In general the electric flux \( \phi \) through an arbitrary two–dimensional surface which is parallel to the GEM (starting from \( x_a \) to \( x_b \) at \( y_a \)) can be calculated according to Eq. 5.37. For the following calculations the direction of the limit is of importance which is notated by \( y_{a}^{\pm} \) (+ limit from top side, − limit from bottom side).

\[ \phi = \int_A \vec{E} \, d\vec{A} = -\frac{1}{2\pi \epsilon_0} \lim_{y \rightarrow y_a^{\pm}} \left[ \sum_{n=0}^{N-2} \int_{x_a}^{x_b} \frac{\partial h_n^{(1)}}{\partial y} \, dx \right] + \int_{x_a}^{x_b} \frac{\partial G_N^{(1)}}{\partial y} \, dx \]  

### Collection efficiency

In order to calculate the collection efficiency the fluxes \( \phi_{\text{cath}} \) and \( \phi_C \) need to be known. The flux at the cathode \( \phi_{\text{cath}} \) can be calculated according to Eq. 5.37 with \( x_a = -(w + L)/4, x_b = (w + L)/4 \) and \( y = (q_1/2)^{-} \). The calculation of the flux \( \phi_C \) turns out to be more complicated since three different cases need to be considered for \( x_b \).

1. **Plateau region (Fig. 5.9(a))**: For low collection field ratios \( \eta_1 \) all field lines on the top side of the
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Table 5.1: Integration limits of the flux $\phi_C$ for the different regions.

<table>
<thead>
<tr>
<th>Region</th>
<th>$x_a$</th>
<th>$x_b$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\eta_1 &lt; \eta_{1,k1}$</td>
<td>$-(w + L)/4$</td>
<td>$-(w + L)/4$</td>
</tr>
<tr>
<td>$\eta_{1,k1} \leq \eta_1 \leq \eta_{1,k2}$</td>
<td>$-(w + L)/4$</td>
<td>$x_{B,top}(\eta_1, \eta_2)$</td>
</tr>
<tr>
<td>$\eta_{1,k2} &lt; \eta_1$</td>
<td>$-(w + L)/4$</td>
<td>$-L/2$</td>
</tr>
</tbody>
</table>

GEM point upwards, i.e. no field lines from the cathode end on the top side (no sign flip of electric field). With $x_a = x_b = -(w + L)/4$ and $y \rightarrow (d/2)^+$, Eq. [5.37] becomes $\phi_C = 0$. The collection efficiency forms a plateau and is limited by $\epsilon_{cell} = 1$.

2. **Formation of the kink** (Fig. 5.9(b)): With an increasing collection field ratio $\eta_1$, the field lines originating from the cathode start to reach the top side of the GEM. A sign flip of the electric field in $y$-direction starts to develop and spreads over the upper copper layer of the GEM (from $x = -(w + L)/4$ to $x = -L/2$ on the left side of the unit cell and from $x = (w + L)/4$ to $x = L/2$ on the right side of the unit cell). The electric field in $y$-direction at the top side of the GEM can be approximated by a Taylor expansion of Eq. [5.11] at $x_T = -(w + L)/4$: The solution of Eq. [5.38] gives the $x$-position of the sign flip of the electric field in $y$-direction $x_{B,\text{top}} = x_{B,\text{top}}(\eta_1, \eta_2)$ for an arbitrary charge density configuration.

$$\text{Taylor} \left[ E^N_y \left( y \rightarrow \frac{d}{2}, x = x_T \right) \right] (x) = 0 \rightarrow x_{B,\text{top}}(\eta_1, \eta_2)$$

(5.38)

The region of the kink can be obtained by investigating $x_{B,\text{top}}(\eta_1, \eta_2)$. The requirement $x_{B,\text{top}}(\eta_1, \eta_2) = -(w + L)/4$ (sign flip starts) leads to the starting point $\eta_{1,k1}$ of the kink. Accordingly the end point $\eta_{1,k2}$ of the kink is given by the requirement $x_{B,\text{top}}(\eta_1, \eta_2) = -L/2$. Finally the collection efficiency for the kink region ($\eta_{1,k1} \leq \eta_1 \leq \eta_{1,k2}$) can be calculated by Eq. [5.37] with $x_a = -(w + L)/4$ and $x_b = x_{B,\text{top}}$.

3. **High collection field** (Fig. 5.9(c)): Once the flip of the electric field in $y$-direction has completely expanded over the top side of the GEM, no field lines originate from the copper layer. For this case the integration limits of Eq. [5.37] are fixed at $x_a = -(w + L)/4$ and $x_b = -L/2$.

The integration limits for the flux $\phi_C$ are summarized in Tab. 5.1. The development and expansion of the position $x_a$ can be seen in Fig. 5.8 for a GEM with standard pitch. At a ratio of $\eta_{1,k1} \approx 0.14$ the electric field lines which originate from the cathode start to reach the top side of the GEM, i.e. the kink of the collection efficiency begins. At a ratio of $\eta_{1,k2} \approx 0.17$ no field lines start from the top side any more and the sign flip has completely expanded over the GEM.

**Extraction efficiency**

The extraction efficiency can be calculated similar to the collection efficiency. The flux at the anode $\phi_{anode}$ can be obtained by Eq. [5.37] where $x_a = -(w + L)/4$, $x_b = (w + L)/4$ and $y \rightarrow (-d/2)^+$. Electrons which manage to reach the bottom side of the hole are either extracted to the anode/next amplification stage or collected at the bottom electrodes of the GEM. The flux through the hole $\phi_{hole}$ can be calculated at the bottom side of the GEM with $x_a = -L/2$, $x_b = L/2$ and $y \rightarrow (-d/2)^+$.  

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Figure 5.8: Development of the integration limit $x_b$ for a standard-pitch GEM at the top electrode ($p = 140 \mu m$, $L = 50 \mu m$, $d = 60 \mu m$, $g_1 = g_2 = 2000 \mu m$, $N = 1000$ and $\eta_2 = 0.1$).

(a) Plateau region ($\eta_1 = 0.10$).
(b) Kink formation ($\eta_1 = 0.16$).
(c) High collection field ($\eta_1 = 0.20$).

Figure 5.9: Electric field component in y-direction at the top side of the GEM for the central unit cell.
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As for the collection efficiency, the flux \( \phi_A \) at the bottom side of the GEM can be separated in three different regions:

1. **Low extraction field:** For low extraction field ratios \( \eta_2 \) no field lines leave from the bottom side to the next amplification stage / anode. The bottom side is fully occupied by a part of the field lines which originate from the GEM hole. In this case the integration limits for the flux \( \phi_A \) are given by \( x_a = x_b = -(w + L)/4 \) and the contribution of flux \( \phi_A = 0 \) vanishes. Thus the extraction efficiency is given by the ratio \( \epsilon_{\text{extr}} = \frac{\phi_{\text{hole}}}{\phi_{\text{hole}}} \) which leads to an offset in case of \( \eta_2 \to 0 \) as a small amount of field lines still reaches the next amplification stage / anode.

2. **Formation of the kink:** Field lines start to leave from the bottom side of the GEM and end on the anode. Like for the collection efficiency, a sign flip of the electric field in y-direction starts to develop and expands at the bottom side. The position of the flip for the extraction efficiency can be obtained from the solution of the following Taylor expansion at \( x_T = -(w + L)/4 \):

\[
\text{Taylor} \left[ E_y^N \left( y \to \left( -\frac{d}{2} \right), x = x_T \right) \right] (x) = 0 \to x_{B, \text{bot}}(\eta_1, \eta_2).
\] (5.39)

Like for the collection efficiency the region of the kink \( (\eta_{2,k1} \leq \eta_2 \leq \eta_{2,k2}) \) can be obtained from the requirement \( x_{B, \text{bot}}(\eta_1, \eta_2) = -(w + L)/4 \) and \( x_{B, \text{bot}}(\eta_1, \eta_2) = -L/2 \).

3. **Plateau region:** No field lines end on the bottom side of the GEM. In this case the integration limits of Eq. (5.37) are fixed at \( x_a = -(w + L)/4 \) and \( x_b = -L/2 \).

The integration limits for the extraction efficiency are similar to the limits of the collection efficiency (cf. Tab. 5.1) with \( x_{B, \text{bot}}(\eta_1, \eta_2) \) instead of \( x_{B, \text{top}}(\eta_1, \eta_2) \). Especially for the symmetric case \( (x_{B, \text{top}}(\eta_1, \eta_2) = \eta) \), \( x_{B, \text{bot}}(\eta_1 = \eta, \eta_2) \) and \( g_1 = g_2 \) the curves of the integration limits \( x_b \) turn out to be equal.

### 5.2.5 Collection and extraction efficiency

Fig. 5.10 and 5.11 show the resulting collection and extraction efficiency for different GEM pitches. The dashed part of the efficiencies describes the expansion of the kink. The position of the kink is highly influenced by the pitch of the GEM: The larger the pitch of the GEM, the earlier starts the kink. If the collection efficiency is desired to be high for a large range of collection ratios \( \eta_1 \), a smaller GEM pitch should be considered. For a fixed ratio \( \eta_2 \), the extraction efficiency can be increased by choosing a GEM with a larger pitch. The range of the kink region depends also on the pitch as the size of the copper layer changes, e.g. the sign flip has to spread over a shorter distance for smaller GEM pitches.

Based on the definitions of the efficiencies (Eq. 5.35 and Eq. 5.36), the definition of the electric flux (Eq. 5.37) and with the discussed integration limits for \( x_a \) and \( x_b \) (cf. Sec. 5.2.4), the following relations can be derived for the collection and extraction efficiency for an arbitrary GEM geometry \( (d, w, L, g_1, g_2, N) \):

\[
\epsilon_{\text{extr}} = \frac{2\pi (c_7 + c_8 \eta_1 + c_9 \eta_2)}{c_4 + c_5 \eta_1 + c_6 \eta_2}, \quad (5.40)
\]

\[
\epsilon_{\text{coll}} = \frac{2\pi (c_7 + c_8 \eta_1 + c_9 \eta_2)}{c_1 + c_3 \eta_1 + c_2 \eta_2}. \quad (5.41)
\]

The efficiencies as given by Eq. (5.40) and Eq. (5.41) are the results of a top-sided collection and a bottom-sided extraction. Additionally, a collection / extraction efficiency can be defined for the bottom
5.2 Efficiency calculations based on an analytic two-dimensional model

Figure 5.10: Collection efficiency for different GEM pitches \((d = 60\, \mu\text{m}, L = 50\, \mu\text{m}, g_1 = g_2 = 2000\, \mu\text{m}, N = 1000\) and \(\eta_2 = 0.1\). The dashed part of the efficiencies describes the expansion of the kink.

Figure 5.11: Extraction efficiency for different GEM pitches \((d = 60\, \mu\text{m}, L = 50\, \mu\text{m}, g_1 = g_2 = 2000\, \mu\text{m}, N = 1000\) and \(\eta_1 = 0.1\). The dashed part of the efficiencies describes the expansion of the kink.
N = 10
N = 50
N = 100
N = 1000

η
col
0 0.05 0.1 0.15 0.2 0.25 0.3 0.35 0.4 0.45 0.5 1.2
0.2 0.4 0.6 0.8 1.0 1.2

Figure 5.12: Collection efficiency for different numbers of GEM holes N (standard-pitch GEM with p = 140 µm, d = 60 µm, L = 50 µm, g₁ = g₂ = 2000 µm and η₂ = 0.1).

/ top side as well. With the definition of the electric fluxes from [115], the following relations can be obtained for the top-sided extraction and the bottom-sided collection efficiencies:

\[ \eta_{\text{extr, top}} = \frac{D}{\phi_{\text{hole}}} = \frac{2\pi(c_7 + c_8 \eta_1 + c_9 \eta_2)}{c_4 + c_6 \eta_1 + c_5 \eta_2}, \quad (5.42) \]

\[ \eta_{\text{coll, bot}} = \frac{D + 2R}{\phi_{\text{anode}}} = \frac{2\pi(c_7 + c_8 \eta_1 + c_9 \eta_2)}{c_1 + c_2 \eta_1 + c_3 \eta_2}. \quad (5.43) \]

The efficiencies are not independent of each other and share common coefficients \( c_i \) (\( i = 1..9 \)) where \( \tilde{c}_i = c_i(g_2 \leftrightarrow g_1) \) (flipped \( g_1 \) and \( g_2 \)).

Fig. 5.12 and 5.13 show the collection and the extraction efficiency as a function of the number of GEM holes \( N \) for a standard-pitch GEM. For lower number of GEM holes field lines from neighboring cells enter the central GEM unit cell at the anode. As a consequence the flux \( \phi_{\text{anode}} \) is overestimated and extraction efficiencies larger than 1 can be observed. This effect decreases with an increasing number of GEM holes and the extraction efficiency is limited by 1. In Sec. 5.2.5 it will be shown that \( \eta_{\text{extr}} \rightarrow 1 \) for \( N \rightarrow \infty \). Since the changes for the efficiencies are small for \( N = 50 \) to \( N = 100 \), a good approximation is already given by \( N = 100 \). Indeed almost all parameters converge for \( N = 100 \) (cf. Fig. 5.15).

**Coefficients of the efficiencies**

The coefficients \( c_i \) and \( \tilde{c}_i \) are parameters which depend on the GEM geometry, i.e. they are functions of the pitch \( p \), the thickness \( d \), the hole diameter \( L \), the distance to the previous amplification stage or cathode \( g_1/2 \), the distance to the next amplification stage or anode \( g_2/2 \) and the number of GEM holes \( N \). Additionally the coefficients \( c_7, c_8 \) and \( c_9 \) turn out to be functions of \( \eta_1 \) and \( \eta_2 \). The same holds for \( \tilde{c}_7, \tilde{c}_8 \) and \( \tilde{c}_9 \). Due to geometric symmetries, the coefficients of the collection efficiency are the coefficients of the extraction efficiency for flipped \( g_1 \) and \( g_2 \), i.e. \( \tilde{c}_i(g_1, g_2) = c_i(g_2, g_1) \) (for \( i = 7, 8, 9 \)). Fig. 5.14 shows the coefficients \( \tilde{c}_7, \tilde{c}_8 \) and \( \tilde{c}_9 \) for a standard-pitch GEM. The coefficients turn out to be constant outside of
5.2 Efficiency calculations based on an analytic two-dimensional model

The kink region (as the integration limits are fixed) and only change within the region of the kink. As the coefficients are not independent of each other the following relations hold for any value of \( N \geq 1 \):

\[
\begin{align*}
    c_1 + c_2 + c_3 &= 2\pi p \\
    c_4 + c_5 + c_6 &= 2\pi L \\
    c_7 + c_8 + c_9 &= \begin{cases} 
        p & \text{for } \eta_2 < \eta_{2,k_1}(\eta_1) \\
        L & \text{for } \eta_2 > \eta_{2,k_2}(\eta_1) 
    \end{cases} \\
    \tilde{c}_7 + \tilde{c}_8 + \tilde{c}_9 &= \begin{cases} 
        p & \text{for } \eta_1 < \eta_{1,k_1}(\eta_2) \\
        L & \text{for } \eta_1 > \eta_{1,k_2}(\eta_2) 
    \end{cases} \\
    c_4 + c_5 &= \pi L
\end{align*}
\]
Hence the sum of all coefficients is constant outside of the kink region, i.e.:

\[ \sum_{i} c_i + c_7 + c_8 + c_9 = \begin{cases} 2\pi(p + L) + p & \text{for } \eta_2 < \eta_{2,k1}(\eta_1) \\ 2\pi(p + L) + L & \text{for } \eta_2 > \eta_{2,k2}(\eta_1) \end{cases} \quad (5.45) \]

\[ \sum_{i} \bar{c}_i + \bar{c}_7 + \bar{c}_8 + \bar{c}_9 = \begin{cases} 2\pi(p + L) + p & \text{for } \eta_1 < \eta_{1,k1}(\eta_2) \\ 2\pi(p + L) + L & \text{for } \eta_1 > \eta_{1,k2}(\eta_2) \end{cases} \]

Fig. [5.15] shows the convergence of the coefficients \(c_1\) to \(c_9\) for an increasing number of GEM holes \(N\). In case of \(N \to \infty\) the following limits and relations can be found:

\[ c_1 \to 0 \]
\[ c_2 \to \pi L \]
\[ c_3 \to \pi(2p - L) \]
\[ c_6 \to \pi L \]

\[ \bar{c}_7(\eta_1 < \eta_{1,k1}) \to 0 \]
\[ \bar{c}_8(\eta_1 < \eta_{1,k1}) \to L/2 \]

\[ \bar{c}_7 + \bar{c}_8 \to \begin{cases} L/2 & \text{for } \eta_1 < \eta_{1,k1}(\eta_2) \\ L/2 & \text{for } \eta_1 > \eta_{1,k2}(\eta_2) \end{cases} \]

\[ \bar{c}_9 \to \begin{cases} p - L/2 & \text{for } \eta_1 < \eta_{1,k1}(\eta_2) \\ L/2 & \text{for } \eta_1 > \eta_{1,k2}(\eta_2) \end{cases} \]

\[ c_7(\eta_2 < \eta_{2,k1}) \to 0 \]
\[ c_8(\eta_2 < \eta_{2,k1}) \to L/2 \]

\[ c_7 + c_8 \to \begin{cases} L/2 & \text{for } \eta_2 < \eta_{2,k1}(\eta_2) \\ L/2 & \text{for } \eta_2 > \eta_{2,k2}(\eta_2) \end{cases} \]

\[ c_9 \to \begin{cases} p - L/2 & \text{for } \eta_2 < \eta_{2,k1}(\eta_2) \\ L/2 & \text{for } \eta_2 > \eta_{2,k2}(\eta_2) \end{cases} \]

**Limits and offsets**

Taking the relations and limits for \(N \to \infty\) into account, the following limits can be obtained for the collection and extraction efficiency where the two–dimensional optical transparency \(\tau := L/p\) has been introduced:

\[ \epsilon_{\text{extr}}(\eta_2 \to \infty, N \to \infty) = \frac{2\pi c_9}{c_6} = 1, \quad (5.46) \]

\[ \epsilon_{\text{coll}}(\eta_1 \to \infty, N \to \infty) = \frac{2\pi \bar{c}_9}{c_3} = \frac{L}{2p - L} = \frac{\tau}{2 - \tau}. \quad (5.47) \]
5.3 Three–dimensional approach for a single GEM hole

The average electric field will be calculated for a GEM hole in 3D (see Fig. 5.16) in order to estimate the influence of the dimensionality. Due to the complexity of the three–dimensional hexagonal structure of a GEM foil, this will only be done for a single GEM hole. The results can be compared to the two–dimensional average field calculations and the deviation factor \( \kappa \) (for \( N = 1 \) and \( \mu_1 = \mu_2 = 0 \), see Sec. 5.2.2). Both holes have a radius of \( R \) and the distance between the top and the bottom electrode is given by \( d \). The upper electrode carries a surface charge density of \( (+\lambda) \) and the lower electrode is described by \( (-\lambda) \). In cylindrical coordinates \((r, \phi, z)\) the charge density configuration is given by Eq. 5.49.

\[
\rho = \lambda \Theta (r - R) \left[ \delta \left( z - \frac{d}{2} \right) - \delta \left( z + \frac{d}{2} \right) \right] ~ (5.49)
\]

For high collection ratios \( \eta_1 \) and \( N \to \infty \) the collection efficiency is limited by the optical transparency \( \tau \). With \( 0 \leq \tau \leq 1 \) the limit of the collection efficiency is given by \( 0 \leq \tau/(2 - \tau) \leq 1 \). As a GEM with no holes \( (\tau = 0) \) can not collect any electrons the collection efficiency vanishes in this case. In the limit of \( \tau = 1 \) all electrons are collected since there are no top or bottom electrodes.

The extraction efficiency \( \epsilon_{\text{extr}} \) has an offset \( \epsilon_{\text{off}} \) for \( \eta_2 \to 0 \). In case of \( N \to \infty \) and for \( \eta_1 < 1 \) the offset is given by

\[
\epsilon_{\text{off}} = \epsilon_{\text{extr}}(\eta_2 \to 0) = \frac{\eta_1}{1 - \frac{\pi}{2L} (1 - \eta_1)} . ~ (5.48)
\]

As expected the extraction efficiency is limited by \( \epsilon_{\text{extr}} = 1 \) for \( N \to \infty \) and large extraction ratios \( \eta_2 \), i.e. the previously mentioned asymmetry of the field lines at the anode disappears and each unit cell gets indistinguishable from each other. In this case all field lines are extracted from the GEM hole and end on the anode. All electrons are extracted.

For high collection ratios \( \eta_1 \) and \( N \to \infty \) the collection efficiency is limited by the optical transparency \( \tau \). With \( 0 \leq \tau \leq 1 \) the limit of the collection efficiency is given by \( 0 \leq \tau/(2 - \tau) \leq 1 \). As a GEM with no holes \( (\tau = 0) \) can not collect any electrons the collection efficiency vanishes in this case. In the limit of \( \tau = 1 \) all electrons are collected since there are no top or bottom electrodes.
As in Sec. 5.2.2, the potential can be obtained by solving Poission's equation. For a three-dimensional space the potential is given by

\[ V = \frac{1}{4\pi \varepsilon_0} \int \frac{\rho(r')}{|r - r'|} \, dr'. \]  

(5.50)

Obviously the investigated charge configuration (see Eq. 5.49) has no angular dependence on \( \varphi \). In this case the Green's function can be developed according to Eq. 5.51 [117] where the modified Bessel functions \( I_\nu(x) \) and \( K_\nu(x) \) have been introduced. Here \( r_< \) describes the smaller value of \( r \) and \( r' \), and \( r_\rangle \) describes the larger value of \( r \) and \( r' \).

\[ \frac{1}{|r - r'|} = \frac{2}{\pi} \int_0^\infty \cos[k(z - z')] \, I_0(kr_<) \, K_0(kr_\rangle) \, dk \]  

(5.51)

Thus the electric potential can be rewritten as

\[ V = \frac{\lambda}{\pi \varepsilon_0} \int \gamma(z, k) \left[ K_0(kr) \cdot \tilde{I}_0(r, R, k) + I_0(kr) \cdot \tilde{K}_0(r, R, k) \right] \, dk \]  

(5.52)

where the following relations have been introduced:

\[ \tilde{I}_0(r, R, k) = \int_0^r r' I_0(kr') \Theta(r' - R) \, dr' , \]  

(5.53)

\[ \tilde{K}_0(r, R, k) = \int_r^\infty r' K_0(kr') \Theta(r' - R) \, dr' , \]  

(5.54)

\[ \gamma(z, k) = \cos \left[ k \left( z - \frac{d}{2} \right) \right] - \cos \left[ k \left( z + \frac{d}{2} \right) \right] . \]  

(5.55)
5.4 Comparison to simulations

Depending on \( r \) and \( R \) the integrals \( \tilde{I}_0(r, R, k) \) and \( \tilde{K}_0(r, R, k) \) are given by

\[
\tilde{I}_0(r, R, k) = \begin{cases} 
0 & \text{if } r < R \\
\frac{1}{k} (r \cdot I_1(kr) - R \cdot I_1(kR)) & \text{if } r \geq R
\end{cases}
\]

and

\[
\tilde{K}_0(r, R, k) = \begin{cases} 
\frac{k}{k} \cdot K_1(kR) & \text{if } r < R \\
\frac{1}{k} \cdot K_1(kr) & \text{if } r \geq R
\end{cases}
\]

which leads to the following potential

\[
V = \frac{\lambda}{\epsilon_0} \left( \frac{k}{\pi} \int \frac{I_0(kr) K_1(kR)}{k} dk \right)
\]

For the average electric field inside the hole only the case \( r < R \) has to be investigated. In cylindrical coordinates the electric field components are given by

\[
E_r^\leq = -\frac{1}{r} \frac{\partial V}{\partial r} = -\frac{R \lambda}{\pi \epsilon_0} \int \frac{r I_1(kr) K_1(kR)}{k} dk ,
\]

\[
E_z^\leq = -\frac{1}{\pi} \int \frac{d\gamma}{d\zeta} \int \frac{I_0(kr) K_1(kR)}{k} dk ,
\]

\[
E_{\phi}^\leq = -\frac{1}{r} \frac{\partial V}{\partial \phi} = 0 .
\]

The average electric field inside the hole with radius \( R \) and height \( d \) is given by

\[
\langle E \rangle_{3D} = \frac{1}{V} \int_V E \ dV = \frac{2}{dR^2} \int_0^R \int_{-d/2}^{d/2} dr \int d\zeta \left[ \left( E_r^\leq \right)^2 + \left( E_z^\leq \right)^2 \right]^{1/2} .
\]

Like for the two–dimensional case a unitless deviation factor \( \kappa_{3D} \) can be introduced which only depends on the geometry \( R \) and \( d \):

\[
\kappa_{3D} = \frac{2}{dR} \int_0^R dr \int_{-d/2}^{d/2} d\zeta \left[ \left( \int \gamma I_1(kr) K_1(kR) \ d\gamma \right)^2 + \left( \int \frac{1}{k} \frac{d\gamma}{d\zeta} I_0(kr) K_1(kR) \ d\gamma \right)^2 \right]^{1/2} .
\]

The average electric field can be written as

\[
\langle E \rangle_{3D} = \frac{\lambda}{\epsilon_0} \kappa_{3D}(R, d) .
\]

For a GEM with \( R = 25 \mu m \) and \( d = 55 \mu m \) the three–dimensional deviation factor is given by \( \kappa_{3D} = 0.7408 \). Experimental values were found in a range of 0.73 to 0.825 [118]. The two–dimensional deviation factor is given by \( \kappa_{2D} = 0.5140 \) (see Fig. 5.5).

5.4 Comparison to simulations

In the following section the efficiencies from the charge density calculations will be compared to simulations. A detailed description of the simulations can be found in [42]. For a given GEM configuration
the electric fields were numerically calculated within ANSYS. The obtained field maps were thereupon used in Garfield++ in order to simulate the charge transfer efficiencies. The simulations were made for a Ne-CO$_2$-N$_2$ (90-10-5) gas mixture and a GEM voltage of 300 V. With a thickness of $d = 50 \mu m$ the electric field inside the GEM is given by $E_{GEM} = 60 \text{kV cm}^{-1}$. For the collection efficiency, the collection field was varied from $E_{\text{above}} = 0 - 15 \text{kV cm}^{-1}$ while the field below the GEM was fixed at $E_{\text{below}} = 0 \text{ V cm}^{-1}$ to allow for a comparison with measurements. Here the field below the GEM was set to 0 V cm$^{-1}$ in order to measure the electrons at the bottom side of the GEM. The extraction field was scaled from $E_{\text{below}} = 0 - 15 \text{kV cm}^{-1}$ for the extraction efficiency while the field above the GEM was kept constant at $E_{\text{above}} = 2000 \text{ V cm}^{-1}$. To simulate the efficiencies a fixed amount of electrons $N_{\text{primary}}$ is homogeneously placed above the top side of a GEM unit cell. The trajectory of each electron is thereupon simulated and the end points are counted depending on the final position: Electrons ending on the top side of the GEM are counted as $N_{\text{top}}$, electrons ending on the bottom side as $N_{\text{bottom}}$. Electrons which manage to reach 300 $\mu m$ below the GEM are regarded as extracted and counted as $N_{\text{transfer}}$. The efficiencies are then given by Eq. 5.65 and 5.66.

$$
\epsilon_{\text{coll}} = \frac{N_{\text{primary}} - N_{\text{top}}}{N_{\text{primary}}} \quad (5.65)
$$

$$
\epsilon_{\text{extr}} = \frac{N_{\text{transfer}}}{N_{\text{transfer}} + N_{\text{bottom}}} \quad (5.66)
$$

### 5.4.1 Influence of diffusion

As the Garfield++ simulations allow to disable diffusion the influence of this effect can be studied and compared to the electric flux calculations (which do not include diffusion as well). The simulated efficiency curves are shown in Fig. 5.17 for a standard-pitch GEM with and without diffusion. Without diffusion the initial rise of the extraction efficiency is much higher than with diffusion, i.e. the extraction efficiency reaches the maximum immediately after the position of the kink. Without diffusion the collection efficiency slightly overestimates the collection efficiency with diffusion after the kink. This changes for higher field ratios. The position of the kink is not affected by diffusion. The results from the calculations are shown in Fig. 5.18. Aside from the different scalings of the $\eta$–axes the calculated efficiencies are comparable to the simulated efficiencies without diffusion. The curves, the offset of the extraction efficiency as well as the limits are in a good agreement to the simulations. The influence of the diffusion will be compensated by a correction factor as it will be discussed in the following sections.

### 5.4.2 Influence of constant charge density distributions

To estimate the impact of the simplification based on constant charge density distributions ($\lambda$, $\mu_1$ and $\mu_2$), the geometry of this model has been implemented in ANSYS giving the possibility to numerically calculate the required charge distributions which lead to constant potentials. Fig. 5.19 shows the charge distribution at the cathode for different distances $g_1$ in case of a standard-pitch GEM with $N = 2$. The potentials on the electrodes have always been chosen such that $E_{\text{above}} = 400 \text{ V cm}^{-1}$, $E_{\text{below}} = 400 \text{ V cm}^{-1}$ and $U_{\text{GEM}} = 300 \text{ V}$. The thickness of the region between the top side of the GEM and the cathode is given by $(g_1 - d)/2$ and varies from 70 $\mu m$ (for $g_1 = 200 \mu m$) to 570 $\mu m$ (for $g_1 = 1200 \mu m$). For common detector setups this region (drift or transfer region) is in the order of 1000 $\mu m$ and higher. In this case the charge distribution at the cathode can be well assumed to be constant (cf. Fig. 5.19).
5.4 Comparison to simulations

Figure 5.17: Simulated efficiencies for a standard-pitch GEM showing the influence of diffusion ($E_{\text{GEM}} = 54.5 \text{ kV cm}^{-1}$, for collection eff. $E_{\text{below}} = 0 \text{ V cm}^{-1}$, for extraction eff. $E_{\text{above}} = 2000 \text{ V cm}^{-1}$, electric field range for scan $E = 0 - 15 \text{ kV cm}^{-1}$).

Figure 5.18: Calculated efficiencies for a standard-pitch GEM ($p = 140 \mu\text{m}$, $d = 60 \mu\text{m}$, $L = 50 \mu\text{m}$, $g_1 = g_2 = 2000 \mu\text{m}$, $N = 10k$, $E_{\text{GEM}} = 54.5 \text{ kV cm}^{-1}$, for collection eff. $E_{\text{below}} = 0 \text{ V cm}^{-1}$, for extraction eff. $E_{\text{above}} = 2000 \text{ V cm}^{-1}$, electric field range for scan $E = 0 - 54.4 \text{ kV cm}^{-1}$).
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Figure 5.19: Charge distribution at the cathode for constant potentials and different thicknesses of the drift region (2D ANSYS calculation for $p = 140\,\mu\text{m}$, $d = 60\,\mu\text{m}$, $L = 70\,\mu\text{m}$ and $N = 2$).

Figure 5.20: Charge distribution at the top electrodes for constant potentials and different thicknesses of the drift region (2D ANSYS calculation for $p = 140\,\mu\text{m}$, $d = 60\,\mu\text{m}$, $L = 70\,\mu\text{m}$ and $N = 2$).

The same observation holds for the anode but with flipped charge signs.

However the situation turns out to be different for the top and the bottom electrodes of the GEM. In this case the charges tend to accumulate at the rim of the electrodes, leading to nonconstant charge distributions. The influence of $g_1$ can be neglected (cf. Fig. 5.20). The same behavior can be observed in case of three–dimensional simulations. Fig. 5.21 shows the top electrode of the GEM unit cell as it was used for the efficiency simulations (see [42] for details). A projection of the charges along the dashed line (between the centers of two GEM holes) leads to a nonconstant charge distribution as well (cf. Fig. 5.22). Consequently the assumption of constant charge density distributions will affect the potentials on the GEM electrodes and thus the electric fields and field ratios $\eta_1$ and $\eta_2$. Taking the influence of the dimensionality into account (cf. Sec. 5.3) a deviation between the simulations and the calculations can be expected in terms of the electric field ratio dependencies (scaling of $\eta$–axes, cf. Fig. 5.17 and 5.18).
5.4 Comparison to simulations

Figure 5.21: Charge distribution at the top side of the GEM unit cell (3D ANSYS calculation for a standard-pitch GEM).

Figure 5.22: Projection of the charge distribution between two GEM holes (3D ANSYS calculation for a standard-pitch GEM).
5.4.3 Correction of the equations

To correct the scalings of the electric field ratios and in order to include diffusion in the calculations the following unitless fit parameters \( s_1, s_2 \) and \( s_3 \) are being introduced. The field ratios \( \eta_1 \) and \( \eta_2 \) are scaled by \( s_1 \) and \( s_2 \). The third fit parameter \( s_3 \) is used to quench the curve of the extraction efficiency. The following fits are used:

\[
\begin{align*}
\epsilon_{\text{coll}}(\eta_1, \eta_2) & \rightarrow \epsilon_{\text{coll}} \left( \eta_1, \eta_2 = \frac{E_{\text{below}}}{E_{\text{GEM}}} \right), \\
\epsilon_{\text{extr}}(\eta_1, \eta_2) & \rightarrow \epsilon_{\text{extr}} \left( \eta_1 = \frac{E_{\text{above}}}{E_{\text{GEM}}}, s_2 \eta_2 \right), \\
c_4 & \rightarrow s_3^{\frac{3.5}{3}} c_4, \\
c_6 & \rightarrow \frac{1}{s_3} c_6.
\end{align*}
\]

To obtain a relation between the tuning parameters and the pitch of the GEM multiple geometries have been simulated. The resulting tuning parameters \( s_1, s_2 \) and \( s_3 \) are shown in Fig. 5.23 as a function of the pitch \( p \). Both electric field scaling factors \( s_1 \) and \( s_2 \) are comparable for smaller GEM pitches and diverge with an increasing pitch. The influence of the diffusion turns out to be unaffected by the GEM pitch (as it is related to the gas properties) and leads to a constant value of \( s_3 = 1.297 \pm 0.002 \) (see Fig. 5.24). By definition the parameter \( s_3 \) is given by \( s_3 = 1 \) in case of no diffusion. Thus only two tuning parameters \( (s_1 \text{ and } s_2) \) depend on the GEM pitch \( p \) (given in \( \mu \text{m} \)). Both parameters can be described by quadratic functions leading to

\[
\begin{align*}
s_1(p) & = (-1.81 \pm 0.09) + (0.048 \pm 0.001) \mu \text{m}^{-1} \cdot p + (9.8 \pm 0.3) \times 10^{-5} \mu \text{m}^{-2} \cdot p^2, \\
s_2(p) & = (-3.80 \pm 0.44) + (0.094 \pm 0.005) \mu \text{m}^{-1} \cdot p - (15.6 \pm 1.3) \times 10^{-5} \mu \text{m}^{-2} \cdot p^2.
\end{align*}
\]

Together with the tuning parameters, the efficiencies can be calculated for any electric field configuration and for different GEM pitches. Finally, Fig. 5.25 and 5.26 show the calculated as well as simulated efficiencies for a standard, a medium and a large-pitch GEM. The propagated error bands are caused by the uncertainties of the fitted tuning parameters.
5.4 Comparison to simulations

Figure 5.23: Tuning parameters \( s_1, s_2 \) and \( s_3 \) for different GEM pitches (with \( d = 50 \mu m, L = 60 \mu m, g_1 = g_2 = 2110 \mu m \) and \( N = 200 \)).

Figure 5.24: Detail view of tuning parameter \( s_3 \) for different GEM pitches.
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Figure 5.25: Calculated (lines) and simulated (dots) collection efficiency for different GEM pitches (with $d = 50\,\mu\text{m}$, $L = 60\,\mu\text{m}$, $g_1 = g_2 = 2\,110\,\mu\text{m}$, $N = 200$, $E_{\text{below}}^{\parallel} = 0\,\text{V cm}^{-1}$ and $E_{\text{GEM}}^{\parallel} = 60\,\text{kV cm}^{-1}$).

Figure 5.26: Calculated (lines) and simulated (dots) extraction efficiency for different GEM pitches (with $d = 50\,\mu\text{m}$, $L = 60\,\mu\text{m}$, $g_1 = g_2 = 2\,110\,\mu\text{m}$, $N = 200$, $E_{\text{above}}^{\parallel} = 2\,000\,\text{V cm}^{-1}$ and $E_{\text{GEM}}^{\parallel} = 60\,\text{kV cm}^{-1}$).
Fundamental properties of Micropattern Gaseous Detectors such as the energy resolution or the ion backflow depend highly on the charge transfer processes between the amplification stages. Based on the previous calculations of the efficiencies, expressions will be derived for the energy resolution and for the ion backflow in the following. The obtained model calculations will be compared to the measurements of the hybrid 2GEM-MM detector of Bonn (see Sec. 6.3) and Yale (see Sec. 6.4) as well as to the quadruple S-LP-LP-S stack of the ALICE TPC upgrade (see Sec. 6.5).

6.1 Energy-resolution model

The signal fluctuation and thus the energy resolution $\sigma/\mu$ of a single GEM foil can be written as

$$\left(\frac{\sigma}{\mu}\right) = \sqrt{F + f N}$$

and depends on the Fano factor $F$, the single gain fluctuation $f$ and the number of collected electrons $N$ (cf. Sec. 2.7). The contribution of electronic noise is neglected. Several statistical processes determine the energy resolution:

- The single electron gain can be described by a Polya distribution as introduced in Sec. 2.6.3. Within dedicated Garfield++ and ANSYS simulations [42], the single electron gain has been simulated for different GEM potentials and gas mixtures. The single electron gains can be seen in Fig. 6.1 in case of a standard-pitch GEM in Ne-CO$_2$ (90-10) and for different GEM voltages. Polya distributions have been fitted to describe the data (cf. Sec. 2.6.3). In the limit of a low GEM potential, the Polya distribution approaches an exponential function (cf. Fig. 6.1(b)). The fitted Polya distributions allow to extract $\theta$ and finally to obtain the single gain fluctuation $f = 1/(\theta + 1)$. The resulting values for $f$ are shown in Fig. 6.2 as a function of $U_{GEM}$. The following empirical fit function has been introduced in order to describe the available data points

$$f(U_{GEM}) = f_0 + \exp[(U_0 - U_{GEM}) s_U]$$

where the limit for high GEM potentials is given by $f_0$. The increase of the single gain fluctuation at lower GEM potentials is located at the potential $U_0$. A further parameter $s_U$ has been introduced to allow for a scaling of the x-axis. A fit of $f(U_{GEM})$ to the extracted single gain fluctuations gives $f_0 = 0.5720 \pm 0.0015$, $U_0 = (147 \pm 3)$ V and $s_U = (0.051 \pm 0.005)$ V$^{-1}$. The curve $f(U_{GEM})$ is
used to calculate the single gain fluctuation as a function of the GEM voltage.

- The amount $N^-$ of collected electrons is given by a binomial distribution. The collection (as well as extraction) probability for a single electron is hereby given by the transfer efficiencies as introduced in Sec. 5.4.3. With $N_0$ as the total amount of primary ionization electrons, the number of collected electrons can be written as $N^- = N_0 \epsilon_{\text{coll}}$ and the variance is given by $\sigma^2 = N_0 \epsilon_{\text{coll}} (1 - \epsilon_{\text{coll}})$.

- The Fano factor $F$ characterizes the fluctuations of the primary ionizations $N_0$ for a fixed radiation energy (cf. Sec. 2.6.1).

The energy resolution of a stack consisting of $J$ GEM stages will be calculated according to

$$\left( \frac{\sigma}{\mu} \right)_J = \left[ \frac{F + f_1}{N_1} + \sum_{j=2}^{J} \frac{f_j}{N_j} \right]^{1/2} $$

(6.3)

where $J \geq 1$. Photon conversion and ionization dominates "above" the first GEM stage, i.e. in the drift region. Due to this the fluctuations of the primary charges (given by the Fano factor $F$) is only included in the term of the first GEM stage. The number of collected electrons $N_j$ at a certain stage is given by the individual transfer efficiencies and absolute gains $G_i$ of the previous GEM foils and can be written as

$$N_j = N_0 \epsilon_{e,1} (1 - C_1 d_1) \cdot \prod_{i=2}^{j} G_{i-1} \epsilon_{e,i-1} \epsilon_{c,i} (1 - C_i d_i)$$

(6.4)

_conversion might also appear between the GEM stages which is neglected since the amplitudes of the corresponding signals are smaller as the gain of the first amplification stage is missing.
Figure 6.2: Single gain fluctuation $f$ for a standard-pitch GEM ($p = 140 \mu \text{m}$) in Ne-CO$_2$ (90-10). Data points extracted from fits of the Polya distribution (Eq. 2.31) to the simulated single electron gains [42]. Finally the fit function $f(U_{\text{GEM}})$ has been fitted to the data points. Obtained fit values are $f_0 = 0.5720 \pm 0.0015$, $U_0 = (147 \pm 3) \text{ V}$ and $s_U = (0.051 \pm 0.005) \text{ V}^{-1}$.

where the electron collection efficiency for a specific GEM is denoted as $\epsilon_{\text{c},j}$ and the electron extraction efficiency as $\epsilon_{\text{e},j}$. As the drifting electrons can be lost due to attachment within the drift region or between the GEM stages, the attachment coefficient $C_j$ (cf. Sec. 2.5) has been introduced. The width of each gap is given by $d_j$. Fig. [6.3] shows the configuration of a triple GEM stack and the calculated electron numbers $N_{-j}$ at each stage.

The total effective gain of a GEM stack can be calculated based on Eq. [6.4] if the additional transfer to the anode (or the readout) is taken into account. Following relation holds

$$G_{\text{eff,stack}} = \frac{1}{N_0} \left[ N_j G_j \epsilon_{\text{e},j} (1 - C_{j+1} d_{j+1}) \right]$$

$$= (1 - C_j d_j) \cdot \prod_{j=1}^J G_{\text{eff},j} \left(1 - C_{j+1} d_{j+1} \right)$$

(6.5)

where the effective gain of a single GEM stage has been introduced as $G_{\text{eff},j} = \epsilon_{\text{e},j} G_j \epsilon_{\text{c},j}$. In case of a negligible attachment ($C_j \to 0$), Eq. [6.5] simplifies to

$$G_{\text{eff,stack}} = \prod_{j=1}^J G_{\text{eff},j}$$

(6.6)
as introduced in Sec. 2.8.1, i.e. the total effective gain of a GEM stack is the product of the single effective gains of the individual stages.

### 6.2 Ion-backflow model

#### 6.2.1 Model calculations

A common definition of the ion backflow (IB) is given in terms of the currents at the anode \( I_{\text{anode}} \) (caused by electrons) and at the cathode \( I_{\text{cathode}} \) (caused by the back drifting ions). Following relation is often used in measurements to determine the ion backflow:

\[
\text{IB} = \left| \frac{I_{\text{cathode}}}{I_{\text{anode}}} \right|. \tag{6.7}
\]

The currents can be written as \( I_{\text{cathode}} = \dot{Q}_{\text{cathode}} = (dN_{\text{cathode}}^+/dt) q_i \) and \( I_{\text{anode}} = \dot{Q}_{\text{anode}} = (dN_{\text{anode}}^-/dt) q_e \) where \( q_i \) and \( q_e \) are the charges of the electrons as well as ions. The amount of positively charged ions at the cathode is given by \( N_{\text{cathode}}^+ \). The amount of electrons at the anode is given by \( N_{\text{anode}}^- \). Assuming that the charges \( q_i \) and \( q_e \) are of opposite sign (\( q_i = -q_e \)) for a single ionization process, the ion backflow - as defined in Eq. 6.7 - can be rewritten as

\[
\text{IB} = \frac{dN_{\text{cathode}}^+}{dN_{\text{anode}}^-}. \tag{6.8}
\]

Finding the functional dependence of \( N_{\text{cathode}}^+ (N_{\text{anode}}^-) \) gives access to obtain the ion backflow of a GEM stack. Following assumptions are taken into account for the ion-backflow model:

- Electrons and ions are always created pairwise within a single ionization process. The amount of created avalanche ions \( N_j^{+} \) at a certain stage \( j \) is thus given by

\[
N_j^{+} = N_j^- G_j - N_j^- (G_j - 1) \tag{6.9}
\]
6.2 Ion-backflow model

Figure 6.4: Different contributions to the ion backflow in case of a triple GEM stack ($J = 3$): Avalanche ions in green, drift ions in blue.

and can be expressed by the number of electrons before ($N_{j}^{−}$) and after the GEM amplification ($N_{j}^{−}G_{j}$) using Eq. 6.4.

- Avalanche ions are mostly created in at the bottom side of the GEM holes (see detailed simulations in [42]). The **avalanche ion collection efficiency** at a GEM stage $j$ is denoted as $\epsilon_{a,c,j}$ The **avalanche ion extraction efficiency** is given by $\epsilon_{a,e,j}$.

- Once avalanche ions have been created, they leave the initial GEM hole and propagate through the GEM stack as **drift ions**, i.e. the outgoing avalanche ions of stage $j$ are the incoming drift ions at stage $j − 1$ (plus drift ions that might have passed stage $j$). The **drift ion collection efficiency** at a GEM stage $j$ is introduced as $\epsilon_{d,c,j}$ and the **drift ion extraction efficiency** as $\epsilon_{d,e,j}$.

Eq. 6.4 illustrates the different efficiencies for avalanche as well as drift ions in case of a triple GEM stack. The number of ions at the cathode $N_{cathode}^{+}$ can be calculated using the different efficiencies and leads for a stack with $J$ GEM foils ($J \geq 1$) to

$$N_{cathode}^{+} = N_{0} \left(1 + \epsilon_{j} \right) \quad (6.10)$$

where $\epsilon_{j}$ is given by

$$\epsilon_{j} = \sum_{j=1}^{J} \epsilon_{c,j}^{−} \left( G_{j} − 1 \right) \epsilon_{c,j}^{\alpha} \epsilon_{c,j}^{\epsilon} \left( \prod_{k=1}^{j-1} G_{eff,k}^{d+} \epsilon_{c,k}^{d+} \epsilon_{c,k}^{d+} \right). \quad (6.11)$$

The ion collection (extraction) efficiencies are functions of the field below (above) the GEM as ions drift in the opposite direction than electrons.
The amount of electrons at the anode $N_{\text{anode}}^-$ is given by the total effective gain of the stack according to Eq. (6.6) and leads to

$$N_{\text{anode}}^- = N_0 G_{\text{eff,stack}} = N_0 \prod_{j=1}^{J} G_j \cdot$$  \hspace{1cm} (6.12)

Accordingly Eq. (6.10) can be written as

$$N_{\text{cathode}}^+ (N_{\text{anode}}^-) = \frac{N_{\text{anode}}^-}{G_{\text{eff,stack}}} (1 + \epsilon_j) \cdot$$  \hspace{1cm} (6.13)

Together with Eq. (6.8) the ion backflow of a multiple GEM stack is given by

$$\text{IB} = \frac{1 + \epsilon_j}{G_{\text{eff,stack}}} \cdot$$  \hspace{1cm} (6.14)

Indeed this is the generally known definition of the ion backflow as introduced in Sec. 2.8.1. By comparing the equations, the term $\epsilon_j$ can be identified as the number of back drifting ions per incoming electron: A closed analytic expression has been found for a multiple GEM stack in terms of the individual gains and electron as well as ion transfer efficiencies which allows a detailed study of the ion backflow.

Each GEM stage contributes as a single term in $\epsilon_j$ (see Eq. (6.11)). Each term describes multiplicatively the propagation of an incoming electron to a specific GEM stage (effective gains and electron efficiencies), the ion creation (absolute gain) and finally the back drifting amount of ions (avalanche / drift ion efficiencies). In case of a triple GEM stack $\epsilon_3$ is given as

$$\epsilon_3 = \frac{e_{c,1}^- (G_1 - 1) e_{c,1}^+ e_{c,1}^+}{e_{c,1}^+ e_{c,1}^+} + G_{\text{eff,1}} e_{c,2}^- (G_2 - 1) e_{c,2}^+ e_{c,2}^+ e_{c,1}^+ e_{c,1}^+ + G_{\text{eff,1}} G_{\text{eff,2}} e_{c,3}^- (G_3 - 1) e_{c,3}^+ e_{c,3}^- e_{c,2}^+ e_{c,2}^+ e_{c,1}^+ e_{c,1}^+ \cdot$$

Fig. 6.5 shows a graphical interpretation of the contributing terms for $\epsilon_3$.
6.2 Ion-backflow model

6.2.2 Simulations of ion efficiencies

The ion-backflow model strongly relies on the avalanche as well as drift ion efficiencies as important input parameters. A treatment of the ion efficiencies turns out to be more sophisticated than for electrons and will be discussed in the following. Important aspects are:

- Avalanche ions are created in the vicinity of the GEM hole. Due to this most of the avalanche ions are collected $\epsilon_{a,i}^+ \approx 1$. The extraction $\epsilon_{a,j}^+$ of the avalanche ions scales with the electric field above the GEM (Fig. 6.7).

- The collection $\epsilon_{d,j}^+$ of the drift ions scales with the electric field below the GEM. Once collected, drift ions are almost completely extracted, i.e. $\epsilon_{d,j}^+ \approx 1$ (Fig. 6.7).

- Important parameters which influence the ion transport are alignment properties such as rotation or displacement of two successive GEM foils. These quantities influence the average hole overlap and determine the geometric blocking of drift ions. Additionally the "flavors" of the GEMs (standard, medium or large pitch) play an important role and must be taken into account.

For the simulations, the initial drift ions have been homogeneously distributed in a fixed area below the investigated unit cell which has been displaced with respect to the calculated average hole overlap $\bar{A}$ of two full-sized $10 \text{ cm} \times 10 \text{ cm}$ GEM foils (see [42] for details). The blue area in Fig. 6.6 corresponds to half of the average hole overlap. Ions are initially distributed within the dashed semicircle. The GEM unit cell is thereupon copied in a repetitive way in order to reproduce a larger area. Fig. 6.7 shows the electron as well as ion efficiencies for a standard-pitch GEM (followed by a second standard-pitch GEM, rotated by $90^\circ$) in Ne-$\text{CO}_2$ (90-10). The shown electron efficiencies are described using the model calculations as discussed in Sec. 5.4.3.

Recent investigations indicate that the spatial distribution of the extracted avalanche ions (which are the incoming drift ions) plays an important role in the determination of the drift ion efficiencies. The picture of a homogeneously and fixed starting distribution of the initial drift ions - having a displacement with respect to the average hole overlap - appears to be inappropriate to describe the drift ion efficiencies. The area and the spatial extension of the initial distribution has to scale with the electric field as well. Fig. 6.8 shows the simulated distribution of the extracted avalanche ions as they arrive at the GEM.
above for different fields $E_{\text{above}}$ (where they will be collected as drift ions). For low extraction fields $E_{\text{above}} \approx 200 \text{ V cm}^{-1}$, the distribution is rather smeared across the GEM unit cell. With an increased extraction field, a spatial structure of the extracted avalanche ions becomes clearly visible. As these avalanche ions are the collected drift ions, the spatial distribution must be taken into account. An analytic approach to calculate the drift ion collection efficiency will be discussed in the following where the spatial distribution of the extracted avalanche ions will be considered.
6.2 Ion-backflow model

(a) \(E_{\text{above}} = 200 \text{ V cm}^{-1}\)  
(b) \(E_{\text{above}} = 1000 \text{ V cm}^{-1}\)  
(c) \(E_{\text{above}} = 2500 \text{ V cm}^{-1}\)  
(d) \(E_{\text{above}} = 4000 \text{ V cm}^{-1}\)

Figure 6.8: Simulated extraction points of the avalanche ions in Ar-CO\(_2\) (90-10) as a function of the field \(E_{\text{above}}\) (standard-pitch GEM \(p = 140 \mu\text{m}, U_{\text{GEM}} = 300 \text{ V and } E_{\text{below}} = 0 \text{ V cm}^{-1}\)). The GEM unit cell (origin of avalanche ions) is indicated by the rectangular box [42].

6.2.3 Calculations of the drift ion collection efficiency

In case of the electron collection efficiencies (cf. Sec. 5), a homogeneous distribution of the initial electrons has been assumed which extends to the whole anode (collection from below) of the GEM unit cell. This assumption is based on the strong widening of the electron distribution while drifting.

However, the initial distributions of the drift ions are strongly correlated to the geometric alignment of two successive GEM foils (displacement, rotation) and the electrostatic field configuration in between. Accordingly, the drift ion collection efficiency has to be calculated for ions which emerge from specific intervals \([x_a, x_b]\) on the anode. These intervals will later be correlated to the distribution of the initial drift ions. The corresponding flux of this interval \(\phi_{\text{anode},i}'\) can thereupon be divided up into a flux \(\tilde{D}'_i\) which enters the GEM hole (collected drift ions) and a flux \(\phi_{\text{A},i}'\) which ends on the lower GEM electrodes (blocked drift ions) (see Fig. 6.9). The generalized drift ion collection efficiency \(e_{\text{gc},i}'(x_a, x_b)\) of GEM...
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Figure 6.9: Electric fluxes at the anode inside in order to define the drift ion collection efficiency.

\[ D_i = D_i / 2 \]

\[ \tilde{D}_i = \frac{D'_i}{D'_i + \phi'_{A,i}} \]  \hspace{1cm} (6.15)

As the point of the transition from flux \( \tilde{D}'_i \) to \( \phi'_{A,i} \) on the anode is not known, both fluxes can not be calculated immediately. With \( \tilde{D}_i = 1/2 D_i \) and \( \tilde{D}_i = \tilde{D}'_i + \tilde{D}''_i \) and \( \phi'_{anode,i} = D_i + 2 \phi_{A,i} \), the flux \( \tilde{D}'_i \) can be expressed as

\[ \tilde{D}'_i = \frac{1}{2} \phi'_{anode,i} - \phi_{A,i} - \tilde{D}''_i \]  \hspace{1cm} (6.16)

where the fluxes \( \phi'_{anode,i} \) and \( \phi_{A,i} \) are known from Sec. 5. The flux \( \tilde{D}''_i \) can be obtained like the flux \( \phi'_{A,i} \) just by integrating the y-component of the electric field on the interval \([0, x_a]\) at the anode. The flux \( \phi'_{A,i} \) can be written as

\[ \phi'_{A,i} = \phi_{A,i} - \phi''_{A,i} \]  \hspace{1cm} (6.17)

The missing flux \( \phi''_{A,i} \) is again given by the y-integration of the electric field at the anode on the interval \([x_b, p/2]\). Finally, the generalized drift ion collection efficiency can be rewritten as

\[ \epsilon_{gc,i}^{d+}(x_a, x_b) = \frac{\phi'_{anode,i} - 2\phi_{A,i} - 2\tilde{D}''_i}{\phi'_{anode,i} - 2\phi''_{A,i} - 2\tilde{D}''_i} \]  \hspace{1cm} (6.18)

Fig. 6.10 shows two GEM unit cells of the same pitch on top of each other, i.e. \( p_{i+1} = p_i \). The "cathode" of the lower GEM is the "anode" of the upper GEM. The extraction field of the lower GEM is chosen equally to the collection field of the upper GEM. In addition, both GEM unit cells are displaced by a shift of \( x_{geom} \). In a first step, the distribution of the extracted ions of the lower GEM will be

\footnote{In case of \( \phi_{anode,i} \), the electric field was integrated on the interval \([-p/2, p/2]\), i.e. the whole anode of the GEM unit cell.}
investigated, which will then be used as the input distribution for the drift ion collection efficiency of the upper GEM. Apparently, the diameter \( x_{D,i+1} \) of the extracted avalanche ion distribution scales with the electron collection efficiency of the lower GEM according to

\[
\epsilon_{c,i+1} = \frac{D_{i+1}}{\phi_{\text{cathode},i+1}} \frac{x_{D,i+1}}{p_{i+1}} = \frac{x_{D,i+1}}{x_{D,i+1} + 2\bar{x}_{D,i+1}}
\] (6.19)

and with the pitch \( p_{i+1} \), the diameter can be written as

\[
x_{D,i+1} = p_{i+1} \bar{\epsilon}_{c,i+1}.
\] (6.20)

The diameter of the extracted avalanche ion distribution defines the initial distribution of the drift ions and thus the integration limits for the generalized drift ion collection efficiencies. In the frame of the upper GEM, the points \( x_1, x_2 \) and \( x_3 \) are of interest. The interval \([x_1, x_2]\) defines the generalized drift ion collection efficiency for ions which emerge “from the left” part of flux \( D_{i+1} \), referred to as \( D_{L,i+1} \). Accordingly, ions emerging from the “right part” (interval \([x_2, x_3]\)) are referred as \( D_{R,i+1} \). Due to symmetric reasons, the flux \( D_{R,i+1} \) contributes on the left side of the GEM unit cell as well. As the electrostatic calculations are mirror-symmetric inside the unit cell, the generalized drift ion collection efficiency on the interval \([x_2, x_3]\) is completely equivalent to those on the interval \([x_2 - \Delta x, x_2]\) with \( \Delta x = x_3 - x_2 \). In the reference frame of the upper GEM, the points are given as:

\[
x_1 = x_{\text{geom}} + \frac{1}{2}(p_{i+1} - p_i) - \frac{1}{2} p_{i+1} \bar{\epsilon}_{c,i+1}, \] (6.21)

\[
x_2 = \frac{1}{2} p_i, \] (6.22)

\[
x_3 = x_{\text{geom}} + \frac{1}{2}(p_{i+1} - p_i) + \frac{1}{2} p_{i+1} \bar{\epsilon}_{c,i+1}. \] (6.23)

Depending on the displacement and for some electrostatic configurations it might happen that \( x_3 < x_2 \). In this case only a single term on the interval \([x_1, x_3]\) contributes to the drift ion collection efficiency. The full drift ion collection efficiency \( \epsilon_{c,i}^{\text{d+}} \) is given by the probability to be part of the "left" flux \( D_{L,i+1} \)
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(given by \((x_2 - x_1)/x_{D,i+1}\)) or to be part of the "right" flux \(D_{R,i+1}\) (given by \((x_3 - x_2)/x_{D,i+1}\)) and the corresponding generalized drift ion collection efficiencies \(f_{i,j}^{d+}\):

\[
\epsilon_{i,j}^{d+} = \begin{cases} 
\epsilon_{i,j}^{d+}(x_1, x_2) + \epsilon_{i,j}^{d+}(x_2 - \Delta x, x_3) & \text{for } x_3 \geq x_2, \\
\epsilon_{i,j}^{d+}(x_1, x_3) & \text{for } x_3 < x_2.
\end{cases}
\]

(6.24)

Fig. 6.11 shows various calculated drift ion collection efficiencies for two standard-pitch GEM foils as a function of different displacements \(x_{\text{geom}}\). Generally, four distinct regions can be observed (cf. Fig. 6.11(a)). As for the electron collection efficiency, a plateau (region I) can be found for low collection fields since no field lines end on the GEM electrodes of the first GEM, i.e. all drift ions are collected. With an increasing collection field, the diameter \(x_{D,i+1}\) shrinks which has a focusing effect on the ion distribution. This leads to the first kink, followed by a steep drop (region II). The extend of this first drop is defined by the "right" flux \(D_{R,i+1}\). Once all ions from the "right" flux are blocked, a second kink appears, followed by a flat drop (region III). This region is completely determined by the "left" flux \(D_{L,i+1}\). For a sufficient high collection field, all field lines from flux \(D_{i+1}\) end on the electrodes of the first GEM stage (region IV), i.e. the drift ions are completely blocked.

The contribution of each region changes with different displacements as shown in Fig. 6.11(b). In case of \(x_{\text{geom}} = p/2 = 70 \mu\text{m}\), both standard-pitch GEM foils are shifted symmetrically, i.e. the fluxes \(D_{L,i+1}\) and \(D_{R,i+1}\) and the contributions of the "left" as well as of the "right" ions turn out to be equal. Consequently, only a single kink can be observed, followed by a steep drop. In case of a vanishing displacement (\(x_{\text{geom}} = 0 \mu\text{m}\)), no drift ion suppression can be observed since no GEM electrodes allow to geometrically block the drifting ions.

The two-dimensional calculations already indicate that the treatment of the drift ions becomes more complicated than the treatment of electrons. By taking a dynamic initial drift ion distribution into account (which scales with the electrostatic field configuration), multiple kinks and drift ion contributions can be found: A strong suppression of the drift ion collection efficiency can be observed which is highly correlated to the displacement of two successively followed GEM foils.

The description of the drift ions becomes even more complicated in case of two different GEM pitches. In this case the picture as given in Fig. 6.10 becomes incomplete since - depending on the GEM pitches and the displacements - multiple overlaps of different sizes and kinds might occur. This requires a unique calculation of the drift ion collection efficiency for each possible overlap as a function of both GEM pitches and a fixed displacement. A general drift ion collection efficiency can thereupon be obtained as a superposition of all possible collection efficiency, weighted by the fractional occurrence probability of each possible overlap. Like for the investigated configuration of equal GEM pitches which leads to two contributions ("left" and "right" ions), multiple contributions can be expected. Indeed a treatment turns out to be highly complicated and will be neglected as - in case of the hybrid detector - only two standard-pitch GEM foils need to be taken into account. In case of the S-LP-LP-S configuration, only standard-standard configurations will be considered to calculate the drift ion collection efficiencies. Nevertheless the required scaling factors to tune the equations are still functions of the different GEM pitches. The simplification of equal GEM pitches will be compensated by regarding the displacement \(x_{\text{geom}}\) as a free tuning parameter in terms of an effective displacement which "averages" over all possible
6.3 Comparison to Bonn hybrid detector

The obtained energy resolution, ion backflow and gain measurements of the Bonn hybrid detector (see Sec. 4) will be compared to the model calculations in the following. A C++ class\(^4\) has been developed which allows to define GEM stacks in terms of the geometry (e.g. GEM types, distances) and the electrostatic configuration (e.g. GEM potentials, transfer fields). The energy resolution, the ion backflow as well as the gain can be immediately calculated based on the introduced model calculations. This gives access to a flexible, detailed and time-saving way to study GEM stacks. The contributions of the individual GEM stages to the stack properties can be studied and the working points (at which ion or electron efficiencies do the single GEM stages operate) can be indicated. A part of this analysis tool has been implemented in the ALICE O²\(^5\) framework which will be used for online as well as offline data analysis with the ongoing upgrade of ALICE (see Sec. 6.6).

A simple example on how to implement the hybrid detector is given in the appendix (see. App. A.1): Once a new instance of a detector has been created for an incoming photon energy of \(E_\gamma = 5.9\) keV in Ne-CO\(_2\) (90-10), a constant attachment factor is globally assumed\(^6\) for the whole stack. The stack is build up successively starting from the drift region (\(E_1 = 400\) V cm\(^{-1}\) and \(d_1 = 2.6\) cm), the first GEM (standard pitch, \(U_{GEM1} = 270\) V) and finally ends with a third GEM (standard pitch, \(U_{GEM3} = 370\) V).

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\(^4\) https://github.com/ritzratz/GEMModel

\(^5\) http://alice-o2.web.cern.ch/

\(^6\) In this simplified implementation the attachment factor is assumed to be independent of the electric field configuration.
Chapter 6 Model calculations

The last stage should contribute as a Micromegas which is realized by tuning the properties of the last "GEM". The gain curve is overwritten by the measured curve of the Micromegas (see Sec. 4.3.1). As a Micromegas is almost opaque for ions and transparent for electrons (see Sec. 2.8.2) the efficiencies are set to $\epsilon_{c,3}^+ = 0.99$, $\epsilon_{c,3}^- = 0.01$ for avalanche ions (there are not drift ions) and $\epsilon_{c,3} = 0.99$, $\epsilon_{c,3} = 1.00$ for electrons.

The calculated energy resolution as well as ion backflow values can be seen in Fig. 6.12 for different TF2 scans. The total effective gain of the stack is shown in Fig. 6.13. Generally, the measurements are in a good agreement to the model calculations. Only the calculated gain curve for $E_{TF2} = 400$ V cm$^{-1}$ shows a linear trend which can not be seen in case of the measured data points: For lower GEM1 potential, the effective gain is overestimated. In case of higher GEM1 potentials, the effective gain is underestimated. This trend will be discussed in detail in case of the Yale hybrid detector (see Sec. 6.4).

The error bands of the ion backflow calculations are based on an error propagation of Eq. 6.14. The errors of the electron efficiencies are given by the accuracy of the fits, i.e. by the tuning factors $s_1$, $s_2$ and $s_3$. The simulated avalanche ion efficiencies (cf. Fig. 6.7) are fitted by higher polynomial functions. Accordingly, the uncertainties of the polynomial coefficients determine the error of the avalanche ion efficiencies. As the drift ion collection efficiencies are calculated using the tuning factors $s_1$ for two successive GEM foils (cf. Sec. 6.2.2), these factors determine the errors. The drift ion extraction efficiency is assumed to be constant $\epsilon_{c,d}^+ = 1$ for the calculations and no error is assumed ($\Delta\epsilon_{c,d}^+ = 0$). The errors of the gains are given by the uncertainties of the fitted exponential functions to the simulated gain curves.

The propagation of $n$ electrons between two amplification stages can be described by a binomial distribution where the probability $p$ is given by the transfer efficiencies and the attachment coefficients. At each stage, $np$ electrons can be expected (cf. Sec. 6.1) and the variance is given by $(\Delta N_1)^2 = np(1-p)$. In the following no errors are assumed for the attachment coefficient as well as for the Fano factor. As the error of the single gain fluctuation turns out to be asymmetric ($\Delta f_{Hi}$ and $\Delta f_{Lo}$, see Fig. 6.2), the following equations are used in order to estimate the lower / upper limit of the energy resolution

$$\Delta \left( \frac{\sigma}{\mu} \right)_{f_{Hi}} = \frac{F + f_1 + \Delta f_{1,Hi}}{N_1 - \Delta N_1} + \frac{f_2 + \Delta f_{2,Hi}}{N_2 - \Delta N_2} + \ldots ,$$

$$\Delta \left( \frac{\sigma}{\mu} \right)_{f_{Lo}} = \frac{F + f_1 - \Delta f_{1,Lo}}{N_1 + \Delta N_1} + \frac{f_2 - \Delta f_{2,Lo}}{N_2 + \Delta N_2} + \ldots .$$

The model calculations allow a detailed study of the dynamics within a GEM stack and to analyze the influence of each stage to the whole setup. Fig. 6.14(a) shows the individual GEM/MM contributions to the total effective gain of the detector for each data point in case of $E_{TF2} = 400$ V cm$^{-1}$: As the total effective gain was kept approximately at 2 000 during the GEM1-MM scans, the calculated gain can be expected to be constant too (see black bars). The effective gain contribution of the first GEM increases with an increasing GEM1 potential (red bars), while the effective gain of the Micromegas decreases (green bars). Most of the gain is contributed by the Micromegas.

The contributions of the single terms for $\epsilon_3$ are shown in Fig. 6.14(b) for each data point within the scans (low GEM1 potentials to high GEM1 potentials). With an increasing GEM1 potential, more
6.3 Comparison to Bonn hybrid detector

<table>
<thead>
<tr>
<th>$E_{TF2}$ / V cm$^{-1}$</th>
<th>Gain scaling</th>
<th>Geometric displacement $x_{geom}$ / µm</th>
<th>Attachment $C$ / cm$^{-1}$</th>
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</thead>
<tbody>
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</tr>
<tr>
<td>400</td>
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<td>52</td>
<td>0.040</td>
</tr>
</tbody>
</table>

Table 6.1: Bonn hybrid detector. Required attachment coefficients, gain scaling factors and geometric displacements (GEM1-Micromegas scan, different TF2). Settings are $E_{Drift} = 400$ V cm$^{-1}$, $E_{TF1} = 3$ kV cm$^{-1}$ and $U_{GEM2} = 250$ V in Ne-CO$_2$ (90-10).

Electrons and thus ions are created within the first GEM stage. As expected the ion contribution of the first GEM increases as well (red bars). However most ions emerge from the second term in $e_3$ (blue bars) which describes the electron collection at the second GEM ($G_{eff,1} e_{c,2}$), followed by the ion creation at the second GEM ($G_2 - 1$) and the ion propagation through the first GEM ($e_{c,2}^{d+} e_{c,1}^{d+} e_{c,1}^{e}$) back to the drift volume. As the second GEM potential and the transfer field between GEM1 and GEM2 was constant during the GEM1-MM scan, only the effective gain of the first GEM ($G_{eff,1}$) and the drift ion transfer at the first GEM ($e_{c,1}^{d+} e_{c,1}^{e}$) changes. Fig. 6.14(c) shows both contributions for the second term in $e_3$ in detail. The increase of the second term is obviously dominated by the effective gain of the first GEM stage. The ion backflow within the GEM1-MM scans is dominated by the number of collected electrons at the second GEM and thus the number of created ions at the second stage. The (drift) transfer probability for these ions back to the drift field (through GEM1) increases slightly but contributes much less compared to the increased amount of ions available at GEM2. The influence of the ions created at the first GEM as well as at the Micromegas is of minor importance.

Nevertheless a scaling of the single GEM gains and an adjustment of the displacement coefficient is required to reproduce the results of the measurements. The factors and coefficients are listed in Tab. 6.1. Generally, an additional gain of a few percent is required as the simulated values seem to underestimate the (real) measured gain curves. Yet not understood, this behavior has already been observed and appears to be an open issue of Garfield++ simulations [119]. The required GEM displacement coefficients $x_{geom}$ and attachment factors turn out to be comparable throughout the TF2 scans. With a hole diameter of $L = 70$ µm and $x_{geom} \approx 50 - 60$ µm for two standard-pitch GEMs, the average hole overlap $\bar{A} = 1 - x_{geom}/L$ is given by approximately 15 – 25 %. Calculations for the hexagonal and three-dimensional scenario of large-sized GEM areas lead to an average hole overlap which corresponds to a shift of about 50 µm of two GEM holes (with respect to the centers, see [42]) which is in a good agreement to the lower limits of the observed displacements.

Fig. 6.15 shows the ion as well as the electron efficiencies of the first GEM in case of $E_{TF2} = 400$ V cm$^{-1}$. The operational working points / conditions are indicated on the curves for $U_{GEM1} = 163$ V. The arrows indicate the directions and ranges in which the efficiencies evolve for increasing GEM1 potentials. With an increasing GEM1 potential (or decreasing ratio $E_{Extern}/E_{GEM}$), less electrons are extracted from GEM1 and the electron transfer probability decreases (product of blue curves, Fig. 6.15). Although less electrons are transferred, an improvement of the energy resolution can still be observed since the increased gain of GEM1 compensates the drop of the transfer probability, i.e. the effective gain increases (cf. Fig. 6.14(a)). The avalanche ion transfer of the first GEM decreases for an increasing GEM1 potential (black curves). Again the exponential growth of the gain in GEM1 compensates this drop which leads to the observed increase of the contribution to the ion backflow (see Fig. 6.14(b)). With an increasing GEM1 potential,
Figure 6.12: Bonn hybrid detector. Energy resolution and ion backflow compared to models (GEM1-Micromegas scan, different TF2). Settings: $E_{\text{Drift}} = 400 \text{ V cm}^{-1}$, $E_{\text{TF1}} = 3 \text{ kV cm}^{-1}$ and $U_{\text{GEM2}} = 250 \text{ V}$ in Ne-CO$_2$ (90-10). The bands indicate the uncertainties of the model calculations.
6.3 Comparison to Bonn hybrid detector

Figure 6.13: Bonn hybrid detector. Total effective gain compared to model calculations (GEM1-Micromegas scan, different TF2). Settings are $E_{\text{Drift}} = 400 \text{ V cm}^{-1}$, $E_{\text{TF1}} = 3 \text{ kV cm}^{-1}$ and $U_{\text{GEM2}} = 250 \text{ V}$ in Ne-CO$_2$ (90-10). The bands indicate the uncertainties of the model calculations.
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Figure 6.14: Bonn hybrid detector. Contributions of the single amplification stages to the total effective gain and the ion backflow for each data point (GEM1-Micromegas scan, $E_{TF2} = 400$ V cm$^{-1}$). Settings are $E_{Drift} = 400$ V cm$^{-1}$, $E_{TF1} = 3$ kV cm$^{-1}$ and $U_{GEM2} = 250$ V in Ne-CO$_2$ (90-10).
more drift ions are capable of passing through GEM1 (red curves). However this is not the dominating contribution to the ion backflow as explained before. The final results are summarized in Fig. 6.16 and show the energy resolution as a function of the ion backflow for all TF2 scans.
6.4 Comparison to Yale hybrid detector

The GEM1-Micromegas scans of the Yale hybrid detector (see [111] for details) will be compared to the model calculations. Instead of changing TF2 during all scans (Bonn scans), the potential of the second GEM has been altered (Yale scans). The used scaling factors and coefficients for the model calculations are listed in table 6.2 and turn out to be stable throughout all scans. Only minor corrections are required. With a constant drift field TF2 of 75 V cm$^{-1}$ and in Ne-CO$_2$ (90-10), the Yale setup has been operated under comparable conditions like the Bonn setup with 80 V cm$^{-1}$. The obtained displacement coefficients of the Yale setup ($x_{\text{geom}} = 59 - 60 \mu$m) are in a good agreement to the coefficients of the Bonn setup ($x_{\text{geom}} = 52 - 60 \mu$m). The increased attachment coefficients might result from different environmental conditions or different contents of gaseous impurities like oxygen or water (water content Yale / Bonn setup < 200 ppm / ≈ 160 ppm, cf. Tab. 6.5). In case of the Bonn hybrid detector, the measured gain curve of the Micromegas has been used for the model calculations. Accordingly only two GEM stages are affected by the underestimated gain prediction of the Garfield++ simulations (cf. Sec. 6.3) which explains the increased gain scaling factors in case of the Yale hybrid.

As the Yale energy resolutions have been obtained by fitting a single Gaussian distribution to the photo peak, divergences in the order of a percent (difference of absolute values) can be expected (see Sec. 4.3.3). Indeed this is shown in Fig. 6.19. In terms of the energy resolution, the raw data of the Yale hybrid deviates by an offset in the order of a percent compared to the model calculations. A subtraction of about 1.2 % (maximum deviation observed with the Bonn hybrid detector in case of $E_{\text{TF2}} = 80$ V cm$^{-1}$, cf. Sec. 4.3.3) shifts the data and leads to comparable results. This again indicates that fitting a single Gaussian strongly overestimates the (real) energy resolution (difference of absolute values). The measured Yale data points are in a good coincidence with the calculated energy resolutions and ion backflows if a shift of the energy resolution is assumed. The shift is justified by the influence of fitting a single Gaussian distribution to the photo peak of $^{55}$Fe. A detailed view of the energy resolution and the ion backflow can
6.4 Comparison to Yale hybrid detector

<table>
<thead>
<tr>
<th>$U_{\text{GEM2}}$ / V</th>
<th>Gain scaling</th>
<th>Geometric displacement $x_{\text{geom}}$ / µm</th>
<th>Attachment $C$ / cm$^{-1}$</th>
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</thead>
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<tr>
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<td>210</td>
<td>1.25</td>
<td>59</td>
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<td>230</td>
<td>1.24</td>
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Table 6.2: Yale hybrid detector. Required attachment coefficients, gain scaling factors and geometric displacements (GEM1-Micromegas scan, different GEM2 potentials [111]). Settings are $E_{\text{Drift}} = 400$ V cm$^{-1}$, $E_{\text{TF1}} = 3$ kV cm$^{-1}$ and $E_{\text{TF2}} = 75$ V cm$^{-1}$ in Ne-CO$_2$ (90-10).

Table 6.3: Obtained coefficients for $f(U_2)$ from linear fits. The uncertainties for the Yale hybrid detector are in the order of $O(10^{-9})$ for $D/\alpha_1$ and $O(10^{-12})$ for $\alpha_2/\alpha_1$ and thus not denoted.

The calculated gain curves are shown in Fig. 6.22. Following a linear trend, they overestimate the measured gain in case of low GEM1 potentials and underestimated the measured gain for higher GEM1 potentials. Assuming that the gains of two individual stages are given by $G_1(U_1, \alpha_1) = g_1 \cdot \exp(\alpha_1 U_1 + m_1)$ and $G_2(U_2, \alpha_2) = g_2 \cdot \exp(\alpha_2 U_2 + m_2)$, the total gain can be written as the product $G(U_1, U_2, \alpha_1, \alpha_2) = G_1(U_1, \alpha_1) \cdot G_2(U_2, \alpha_2)$. Within the GEM1-MM scans, the potentials $U_1$ and $U_2$ have been manually adjusted in a way to keep the total gain constantly at 2 000. Let the functional dependence $U_1(U_2) = f(U_2, \alpha_1, \alpha_2)$ to be chosen such that the total gain

$$G(U_1, U_2, \alpha_1, \alpha_2) = g_1 g_2 \cdot \exp[a_1 f(U_2, \alpha_1, \alpha_2) + m_1 + a_2 U_2 + m_2] = C$$

is constant $C$ for all possible potentials $U_2$. It immediately follows that the dependence between both potentials is linearly given by

$$f(U_2, \alpha_1, \alpha_2) = \frac{D}{\alpha_1} - \frac{\alpha_2}{\alpha_1} U_2$$

where $D = \ln[C/(g_1 g_2)] - m_1 - m_2$. Indeed this can be seen for all GEM1-MM scans with the Bonn as well as with the Yale hybrid detectors (Fig. 6.17). A linear regression has been fitted to the data and the obtained fit values are listed in Tab. 6.3. Throughout all measurements, the ratio $\alpha_2/\alpha_1$ is found to be similar and close to 1.

1Transfer efficiencies and attachment are neglected in this simple approach.
The model calculations rely on simulated gain curves which - as already introduced - require scaling factors as they underestimate the (real) measured gains. Additionally, the simulated gain curves are based on ideal GEM geometries. Due to tolerances within manufacturing processes, deviations can be expected (e.g. inner and outer hole diameters differ from the design values) which lead to gain fluctuations. Consequently, the measured dependence of $f(U_2, \alpha_1, \alpha_2)$ differs from the simulated gain curves. Fig. 6.18 shows the ideal (solid red) curve of $G(U_1 = f(U_2, \alpha_1, \alpha_2), U_2, \alpha_1, \alpha_2)$ where the slopes $\alpha_1$ and $\alpha_2$ of the "measured" and fixed dependence $f(U_2, \alpha_1, \alpha_2)$ are equal to the simulated slopes. As expected the gain stays constant. Small deviations of the simulated slopes ($\alpha_1 \rightarrow \delta_1 \alpha_1$ and $\alpha_2 \rightarrow \delta_2 \alpha_2$) will now cause deviations of the total gain according to $G(U_1 = f(U_2, \alpha_1, \alpha_2), U_2, \delta_1 \alpha_1, \delta_2 \alpha_2)$ (solid blue and solid black curves). Apparently already minor changes in the order of less a percent have drastic consequences on the predicted gain curves and lead to the linear deviations as observed for the model calculations. Deviations of the constant values $m_1$ and $m_2$ lead to the observed shifts of the gain curves (dashed black curve).

The gain of the hybrid detector is dominated by the Micromegas. As the gain curve of the Micromegas is known for the Bonn hybrid detector, the observed differences are much smaller compared to the Yale hybrid detector. The gain discrepancies can be traced back to small deviations of the simulated gain curves.

Figure 6.17: Potential of GEM1 as a function of the Micromegas potential for the GEM1-MM scans of the Bonn as well as the Yale hybrid detectors. Linear regressions have been fitted to the data points.
6.4 Comparison to Yale hybrid detector

Figure 6.18: Influence of small deviations of the slopes $\alpha_1$ and $\alpha_2$ as well as constant values $m_1$ and $m_2$ to the calculated gain. Apparently already minor changes have drastic consequences and lead to strong deviations of the gain curves (for the calculations: ratio $\alpha_1/\alpha_2 = 1$, $\alpha_1 = \alpha_2 = 1$, $g_1 = g_2 = 1$, $m_1 = m_2 = 1$ and $C = 2000$).

Figure 6.19: Yale hybrid detector. Energy resolution versus ion backflow (model calculations and measurements for GEM1-Micromegas scans, different GEM2 potentials). Data points from [111], errors unknown. Settings are $E_{\text{Drift}} = 400 \text{ V cm}^{-1}$, $E_{\text{TF1}} = 3 \text{ kV cm}^{-1}$ and $E_{\text{TF2}} = 75 \text{ V cm}^{-1}$ in Ne-CO$_2$ (90-10).
Figure 6.20: Yale hybrid detector. Energy resolution compared to model calculations (GEM1-Micromegas scans, different GEM2 potentials). Data points from [111], errors unknown. An offset of 1.2 % has been subtracted from the energy resolutions in order to correct for the influence of fitting a Gaussian distribution to the photo peak of $^{55}$Fe. Settings are $E_{\text{Drift}} = 400 \text{ V cm}^{-1}$, $E_{\text{TF1}} = 3 \text{ kV cm}^{-1}$ and $E_{\text{TF2}} = 75 \text{ V cm}^{-1}$ in Ne-CO$_2$ (90-10). The bands indicate the uncertainties of the model calculations.
6.4 Comparison to Yale hybrid detector

Figure 6.21: Yale hybrid detector. Ion backflow compared to model calculations (GEM1-Micromegas scans, different GEM2 potentials). Data points from [111], errors unknown. Settings are $E_{\text{Drift}} = 400 \text{ V cm}^{-1}$, $E_{\text{TF1}} = 3 \text{ kV cm}^{-1}$ and $E_{\text{TF2}} = 75 \text{ V cm}^{-1}$ in Ne-CO$_2$ (90-10). The bands indicate the uncertainties of the model calculations.
Figure 6.22: Yale hybrid detector. Total effective gain compared to model calculations (GEM1-Micromegas scans, different GEM2 potentials). Data points from [111], errors unknown. Settings are $E_{\text{Drift}} = 400 \text{ V cm}^{-1}$, $E_{\text{TF1}} = 3 \text{ kV cm}^{-1}$ and $E_{\text{TF2}} = 75 \text{ V cm}^{-1}$ in Ne-CO$_2$ (90-10). The bands indicate the uncertainties of the model calculations.
6.5 Comparison to quadruple GEM stack

The S-LP-LP-S quadruple GEM stack of the ALICE TPC upgrade program (measurements from [113]) will be compared to the model calculations. During each scan, the potential of GEM1 was changed and the total effective gain was kept constant by tuning the potentials of GEM3 and GEM4 (with fixed ratio \( U_{\text{GEM3}} / U_{\text{GEM4}} = 0.8 \)). The obtained energy resolution, ion backflow and gain calculations are shown in Fig. 6.24, Fig. 6.25 and Fig. 6.26 in case of different \( U_{\text{GEM2}} \) potentials in Ne-CO\(_2\)N\(_2\) (90-10-5). Like for the Yale hybrid detector, an offset has been subtracted from the energy resolution data in order to correct for the influence of fitting a single Gaussian distribution to the photo peak of \(^{55}\)Fe. The final plot showing the energy resolution as a function of the ion backflow is shown in Fig. 6.23.

The required coefficients as well as scaling factors are listed in table 6.4. A summary of all required scaling factors, coefficients and the water / oxygen contents (Bonn hybrid, Yale hybrid, ALICE S-LP-LP-S) is given in table 6.5. Again stable and consistent factors can be found throughout all scans with the quadruple stack and only minor changes are observed. For Ne-CO\(_2\) (90-10), the attachment coefficient increases with the amount of impurities (comparing Yale and Bonn setups). A drop can be seen in case of Ne-CO\(_2\)-N\(_2\) (90-10-5) which is in accordance with the simulated attachment coefficients (cf. Sec. 2.5) for different gas compositions. The required gain scaling factors for Ne-CO\(_2\)-N\(_2\) (90-10-5) turn out to be higher compared to Ne-CO\(_2\) (90-10). However corrections up to a factor of 2 have been reported [119]. The calculated gain curves show linear deviations which have already been observed in case of the hybrid detectors. The origin of these discrepancies has been discussed in detail for the Yale hybrid detector (cf. Sec. 6.4).

In case of the S-LP-LP-GEM stack, only standard-standard GEM pitch configurations have been considered in order to calculate the drift ion collection efficiencies (cf. Sec. 6.4). Yet the scaling factors are still functions of the GEM pitches. The resulting effective geometric displacement \( x_{\text{geom}} = 59 – 60 \mu \text{m} \) for the whole quadruple GEM stack turns out to be comparable to the Yale as well as to the Bonn hybrid detector. Nevertheless a worse description can be observed in case of the ion backflow (Fig. 6.25). This discrepancy might result from the simplifications of the drift ion collection efficiencies which is expected to increases with the number of amplification stages (four GEM stages compared to three amplification stages in case of the hybrid detectors). Generally a good agreement can be observed in case of the energy resolution calculations.

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<th>( C / \text{cm}^{-1} )</th>
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<tr>
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Table 6.4: ALICE S-LP-LP-S quadruple GEM stack. Required attachment coefficients, gain scaling factors and geometric displacements (GEM1-GEM3/4 scans for fixed ratio \( U_{\text{GEM3}} / U_{\text{GEM4}} = 0.8 \), different GEM2 potentials [113]). Settings are \( E_{\text{Drift}} = 400 \text{ V cm}^{-1} \), \( E_{\text{TF1}} = 4 \text{ kV cm}^{-1} \), \( E_{\text{TF2}} = 4 \text{ kV cm}^{-1} \), \( E_{\text{TF3}} = 0.1 \text{ kV cm}^{-1} \) and \( E_{\text{TF4}} = 4 \text{ kV cm}^{-1} \) in Ne-CO\(_2\)N\(_2\) (90-10-5).
Chapter 6 Model calculations

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<th>Setup</th>
<th>Gas</th>
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<th>( C / \text{cm}^{-1} )</th>
<th>( \text{O}_2 / \text{ppm} )</th>
<th>( \text{H}_2\text{O} / \text{ppm} )</th>
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<td>\text{Ne-CO}_2</td>
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<tr>
<td>Yale hybrid</td>
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<td>0.083 – 0.095</td>
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<td>&lt; 200</td>
</tr>
<tr>
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<td>59 – 60</td>
<td>0.06</td>
<td>≈ 33</td>
<td>≈ 250</td>
</tr>
</tbody>
</table>

Table 6.5: Summary of all used attachment coefficients, gain scaling factors and geometric displacements for the hybrid detectors (Yale and Bonn) as well as the ALICE S-LP-LP-S quadruple GEM stack. Oxygen and water content values for S-LP-LP-S / Yale from [113]/[111].

Figure 6.23: ALICE S-LP-LP-S quadruple GEM stack. Energy resolution versus ion backflow (model calculations and measurements for GEM1-GEM3/4 scans with fixed ratio \( U_{\text{GEM1}}/U_{\text{GEM4}} = 0.8 \), different GEM2 potentials). Data points from [113], errors unknown. An offset of 1.0 % has been subtracted from the energy resolutions in order to correct for the influence of fitting a Gaussian distribution to the photo peak of \( ^{55}\text{Fe} \). Settings are 

\[
E_{\text{Drift}} = 400 \text{ V cm}^{-1},
E_{\text{TF1}} = 4 \text{ kV cm}^{-1},
E_{\text{TF2}} = 4 \text{ kV cm}^{-1},
E_{\text{TF3}} = 0.1 \text{ kV cm}^{-1}
\]

and

\[
E_{\text{TF4}} = 4 \text{ kV cm}^{-1}
\]

in Ne-CO\(_2\)N\(_2\) (90-10-5).
6.5 Comparison to quadruple GEM stack

Figure 6.24: ALICE S-LP-LP-S quadruple GEM stack. Energy resolution compared to model calculations (GEM1-GEM3/4 scans for fixed ratio $U_{\text{GEM3}}/U_{\text{GEM4}} = 0.8$, different GEM2 potentials). Data points from [113], errors unknown. An offset of 1.0 % has been subtracted from the energy resolutions in order to correct for the influence of fitting a Gaussian distribution to the photo peak of $^{55}$Fe. Settings are $E_{\text{Drift}} = 400 \text{ V cm}^{-1}$, $E_{\text{TF1}} = 3 \text{ kV cm}^{-1}$ and $E_{\text{TF2}} = 75 \text{ V cm}^{-1}$ in Ne-CO$_2$N$_2$ (90-10-5). The bands indicate the uncertainties of the model calculations.
Figure 6.25: ALICE S-LP-LP-S quadruple GEM stack. Ion backflow compared to model calculations (GEM1-GEM3/4 scans for fixed ratio $U_{GEM3}/U_{GEM4} = 0.8$, different GEM2 potentials). Data points from [113], errors unknown. Settings are $E_{Drift} = 400 \text{ V cm}^{-1}$, $E_{TF1} = 3 \text{ kV cm}^{-1}$ and $E_{TF2} = 75 \text{ V cm}^{-1}$ in Ne-CO$_2$N$_2$ (90-10-5). The bands indicate the uncertainties of the model calculations.
Figure 6.26: ALICE S-LP-LP-S quadruple GEM stack. Total effective gain compared to model calculations (GEM1-GEM3/4 scans for fixed ratio $U_{GEM1}/U_{GEM4} = 0.8$, different GEM2 potentials). Data points from [113], errors unknown. Settings are $E_{\text{Drift}} = 400 \text{ V cm}^{-1}$, $E_{\text{TF1}} = 3 \text{ kV cm}^{-1}$ and $E_{\text{TF2}} = 75 \text{ V cm}^{-1}$ in Ne-CO$_2$N$_2$ (90-10-5). The bands indicate the uncertainties of the model calculations.
6.6 Implementation in the ALICE O² framework

Due to the upgrade of ALICE and the increased data rates during the continuous readout, a new computational concept is currently in development which will be used for the online as well as for the offline (Online - Offline, O²) data reconstruction and analysis during Run 3 and Run 4 of the LHC. A major aspect of the upgrade is the immediate transfer of all detector data to the O² computing farm. The O² system performs detector calibration and data reconstruction online, e.g. cluster finders reduce the amount of data on-the-fly by replacing the raw data. Only reconstructed data will be processed and stored permanently. Raw data will be discarded. Up to now the reconstruction was done fully offline. Detailed information about the upgrade concept of the ALICE O² computing system can be found in the dedicated Technical Design Report [120]. The ALICE O² software repository[^1] is developed from scratch and uses state of the art C++ and modern coding concepts. It includes the relevant and detector specific source code for the reconstruction, the calibration and the simulation of the ALICE experiment.

The models of the electron charge transfers and the energy resolutions have been added to the TPC implementation and allow to dynamically calculate the efficiencies as a function of the electrostatic field configuration and the applied GEM potentials. Prior to this these values have been hard-corded. The gains of the individual stages as well as the energy resolution of the stack can now be immediately calculated. The implemented energy resolution curves are adjusted on the outcome of the comparison of the S-LP-LP-S stack to the model calculations (cf. Sec. 6.5). The implementation in O² is based on a simplified version of the C++ class presented in Sec. 6.3. An example source code which allows to reproduce the energy resolution and the gain of the ALICE baseline configuration (cf. Sec. 6.5) with the ALICE O² framework is given in the appendix A.2.

[^1]: https://github.com/AliceO2Group/AliceO2
CHAPTER 7

Summary

In the context of the ongoing upgrade of the Large Hadron Collider at CERN, the interaction rate of ALICE will be increased to 50 kHz for Pb-Pb collisions. As a consequence, a continuous readout of the Time Projection Chamber will be required which was previously limited to rates in the order of 300 Hz due to a gated readout system based on Multiwire Proportional Chambers. To keep the amount of space-charge distortions at a manageable size, the ion backflow of the charge amplification system has to be significantly reduced. At the same time an excellent detector performance and stability of the system has to be maintained. In Ne-CO$_2$-N$_2$ (90-10-5) and at a total gain of 2 000, the energy resolution must not exceed 12% for $^{55}$Fe which translates to a $dE/dx$ resolution of about 5.5% and 7% in pp and central Pb-Pb collisions. The ion backflow should be lower than 1% in order to be able to completely correct all effects from space-charge distortions. A solution with four GEM foils (S-LP-LP-S configuration) has been adopted as baseline solution for the upgraded chambers, as the concept of GEM detector was more studied and understood at the time of the decision.

7.1 Experimental setup

As an alternative approach to the ALICE baseline solution for the upgrade of the Time Projection Chamber, a hybrid GEM-Micromegas detector consisting of one Micromegas and two GEM foils has been assembled and investigated in terms of the energy resolution, the ion backflow and the gain. Micromegas offer an intrinsic suppression of the ion backflow in the order of a few percent at relatively high gains. As a consequence less GEM stages are required to block the back drifting ions. The GEM foils are commonly used as preamplification stages in order to reduce the needed gain of the Micromegas and to keep the discharge probability at a reasonable level.

The investigated setup consisted of two standard-pitch GEM foils (pitch of 140 $\mu$m) with an active area of 10 cm $\times$ 10 cm. A bulk Micromegas with a gap of 128 $\mu$m and a mesh with approximately 160 lines / cm has been operated as last stage. The whole setup has been assembled within a gas-tight vessel with a volume of about 6 liters. Premixed gas bottles were used in order to flush the setup with Ne-CO$_2$ (90-10) and flows of about 2 – 6 L h$^{-1}$. A Kapton window on the lid and on the side of the vessel allowed to externally irradiate the drift volume with an $^{55}$Fe source and an X-ray tube (Mini-X from Amptek). The pressure, the water as well as the oxygen content was monitored using a Rapidox device.

A summation card has been designed in order to sum up the 128 pads at the segmented readout plane. The card can be directly connected to the Micromegas and is placed inside the vessel. Different configurations
can be selected, allowing to read out only the innermost pads up to the full pad plane.

Within the scope of this work, a slow control has been developed in LabView which gives full access to control and to operate Micro-Pattern Gaseous Detectors (MPGD) such as GEM stacks or the hybrid detector. The operation can be fully automated by user-defined script files which can be loaded and executed within the slow control. This allows to run dedicated measurements over long time scales, e.g. ramp to specific voltage settings, start or stop current measurements, record and save MCA spectra and more. The slow control is capable of communicating with the Bonn PAMeters as well as with the Zagreb PAMeters which are used within the QA program of the ALICE TPC upgrade. The slow control has become the new standard for MPGD operation in the laboratories.

7.2 Background / Energy resolution studies

A common way to determine the energy resolution (in terms of $\sigma/\mu$) is to fit a single Gaussian distribution to the photo peak of $^{55}$Fe spectra. In fact, the photo peak consists of two energy lines ($K_\alpha$ and $K_\beta$) which is neglected in this case as they cannot be resolved with the achievable energy resolution given by gaseous detectors. The influence of the suppressed $K_\beta$ line was studied by looking at the possible Gamma and Auger electron emissions. Comparing both methods indicates that fitting a single Gaussian distribution overestimates the energy resolutions, i.e. the $K_\beta$ peak should be taken into account. Additionally a small excitation of copper was studied as copper layers are irradiated as well (like copper layers of the GEM foils or the cathode): Copper excitations are expected with a probability of a few percent and lead to minor changes of the energy resolutions, i.e. the influence of copper excitations can be neglected.

A dedicated study of the recorded MCA spectra for three different conditions of irradiation ($^{55}$Fe collimated / not collimated from top window, collimated from side window) indicates that the background is composed of two different contributions: Sharing causes (with approximately 68% of background contribution) a lower energetic tail on the left side of the photo peak and appears if charges from photo conversions are partially lost. This effects dominates in case of uncollimated or displaced sources (with respect to the readout pads) as more photons convert in the vicinity of the investigated pads. The second background contribution (approximately 32%) appears as an intrinsic broad exponential component of the detector and an estimation of the background rate yields that cosmics can be excluded. Both background contributions are correlated by different conditions of irradiation, i.e. a subtraction of two recorded background spectra (under different conditions) results in the third background spectrum.

Based on the background studies and the investigations of the $K_\beta$ line, a fit model has been derived in order to describe the full MCA spectra. This model takes sharing (modeled by the product of an error function and a linear function) as well as the second broad exponential background contribution into account. Both the $K_\alpha$ as well as the $K_\beta$ lines are described by Gaussian functions which are coupled by their ratios of intensities and amplitudes. A further exponential function is used in order to describe the lower energetic electronic noise given by the readout electronics. Comparing the advanced fit model to the method of fitting a single Gaussian distribution to the tip of the photo peak yields that the energy resolutions differ in a range between 1 – 2% (difference of absolute values, energy resolutions obtained by a single Gaussian function are overestimated). Consequently a proper description of the background and the inclusion of the $K_\beta$ line should be always considered.
7.3 Measurements and comparison to Yale hybrid / quadruple GEM stack

The hybrid detector has been investigated within GEM1-Micromegas scans in Ne-CO$_2$ (90-10) and for different transfer fields between the second GEM and the Micromegas. The potential of the first GEM stage has been scanned while adjusting the Micromegas potential in order to achieve a constant gain of 2 000. The measurements of the ion backflow and the energy resolution clearly show that the hybrid detector fulfills the requirements of the ALICE TPC upgrade (energy resolution $\sigma/\mu \leq 12\%$ for $^{55}$Fe and ion backflow $\leq 1\%$ at a gain of 2 000). By using the advanced fit model for the energy resolution, a value of about $\sigma/\mu \approx 10\%$ and an ion backflow of $0.7\%$ can be achieved. To allow for a comparison with the ALICE S-LP-LP-S baseline solution, a single Gaussian distribution has been fitted to the photo peak as well. In this case the hybrid detector clearly competes with the quadruple GEM stack in terms of ion backflow and energy resolution. A comparison to an alternative hybrid setup of the Yale University allows to reproduce a specific GEM1-Micromegas scan. A crosscheck of some specific settings allows to reproduce the gain as well as the ion backflow. However divergences of about $0.5\%$ to $1.0\%$ (difference of absolute values) can be observed for the energy resolution. Nevertheless both detectors lead to comparable measurements and do not indicate major inconsistent behaviors. Within the limits of the given measurable accuracy and the environmental uncertainties, trends of the detector performances can be reproduced.

7.4 Model calculations

Characteristic quantities of MPGD (like energy resolution, ion backflow and gain) highly depend on the charge transfer processes between the individual amplification stages. The charge transfer between multiple GEM stages can be described by the collection as well as the extraction efficiencies. The collection efficiency describes the probability to collect the charge carriers into the holes of the GEM foil. Accordingly the extraction efficiency describes the probability of extraction the charges to the next amplification stage or the readout plane. Within this work, an analytic model has been derived in order to describe the transfer efficiencies for an arbitrary GEM geometry and electrostatic configuration. The model is based on a two-dimensional and simplified cut of the hexagonal GEM structure. The GEM electrodes as well as the anode and the cathode are modeled by constant charge distributions. No polyimide layer is considered. The geometry of the GEM is defined in terms of the pitch, the thickness and the hole diameter (no differentiation between an inner and an outer diameter). Based on the charge distribution, the electric field can be obtained by solving Poisson’s equation.

The electric field calculations inside a GEM hole show that the average electric field can be expressed as the field of a parallel plate capacitor where an additional deviation factor is required. The deviation factor $\kappa$ can be expressed in terms of the GEM geometry and lower orders are derived. Three-dimensional calculations for a standard-pitch GEM lead to a value of $\kappa_{3D} = 0.7408$ which is in a good agreement to observed measurements (0.73 to 0.825 [118]). The average field above or below a GEM (e.g. induction field, transfer field) can be well approximated as the field of a parallel plate capacitor. In this case the deviation factor can be neglected.

Electric flux calculations lead to closed analytic expressions of the efficiencies as functions of the charge density ratios. The obtained equations inherently include the appropriate limits and offsets that can be seen in simulations so far. Both the collection as well as the extraction efficiency turn out to be
correlated by common geometric coefficients. In addition correlations between the coefficients can be derived. Most coefficients already show a sufficient convergence in case of lower number of GEM holes for \( N > 100 \). Analytic limits are derived in case of \( N \to \infty \) which allow to express the limitations and offsets of the efficiencies in terms of the optical transparency and the geometry of the GEM. Generally three characteristic regions of the efficiency curves can be observed which can be traced back to three different electric field line scenarios.

As the obtained efficiencies are solely based on two-dimensional electric field calculations, a certain deviation to simulations can be expected as no polyimide layers and no gas properties such as diffusion have been taken into account. Furthermore this model is based on the assumption of constant charge density distributions. Numerical calculations of three-dimensional GEM structures in ANSYS allow to obtain the charge distributions on the electrodes if constant potentials and electric fields are assumed. The assumption of a constant charge density distribution on the anode and the cathode turns out to be applicable for distances exceeding 1 000 \( \mu \)m which is usually the case. However non-constant distributions can be observed on the GEM electrodes. In fact, this turns out to be a strong simplification of the model calculations. By introducing three tuning parameters, an appropriate transformation of the geometric parameters becomes possible, allowing to correct for the electrostatic simplifications and to include diffusion. Both tuning factors to correct for the electrostatic simplifications scale with the pitch of the GEM. The diffusion factor turns out to be independent of the pitch and changes with the gas composition. Finally the efficiencies can be calculated for an arbitrary electric field configuration and GEM pitch.

The derived transfer efficiencies can be used in combination with the gains of the amplification stages to calculate the effective gain of GEM stacks. A model has been introduced to calculate the energy resolution \( \sigma/\mu \) in which each GEM stage contributes with the corresponding single gain fluctuation and the number of collected electrons. Furthermore, a closed analytic expression has been found for the ion backflow as a function of the individual gains and electron as well as ion transfer efficiencies. The number of back drifting ions per incoming electron \( \epsilon \) can be traced back to the charge transfers and gains of the individual stages. An important aspect is the differentiation between drift and avalanche ion efficiencies: Avalanche ions (i.e. ions created within an avalanche processes inside of a GEM hole) show different transfer properties as drift ions which enter the GEM (i.e. ions which emerged from previous amplification processes / GEMs). Drift (avalanche) ions are almost completely extracted (collected). The drift ion collection as well as the avalanche ion extraction scale with the according external electric fields. An important aspect is the initial distribution of the ions which enter a GEM as drift ions. These distributions highly influence the drift ion collection efficiency. Within the two-dimensional charge transfers calculations, two GEM foils of equal pitches have been placed on top of each other, allowing to calculate the drift ion collection efficiencies. Depending on the displacement of both GEM foils and the electrostatic configuration, strong suppressions of the drift ion collection efficiencies can be observed; Mostly dominated by the geometric blocking of two GEM foils.

The derived model calculations give access to a flexible, time-saving and in-depth study of the internal processes and dynamics of GEM stacks. In this context a C++ class has been developed which allows to define GEM stacks in terms of the geometry (e.g. GEM types, distances) and the electrostatic configuration (e.g. GEM potentials, transfer fields). The energy resolution, the ion backflow as well as the gain can be immediately calculated. The contributions of the individual GEM stages to the stack properties can be studied in detail and the working points (at which ion or electron efficiencies do the single GEM stages operate) can be indicated.
The GEM1-Micromegas scans of the Bonn and the Yale hybrid detectors show a good agreement to the model calculations in terms of the ion backflow and the energy resolution. In case of the Yale setup, a constant offset has been subtracted for the energy resolution. The offset is justified by comparing both methods (advanced fit model versus simple Gaussian distribution) to determine the energy resolution in case of the Bonn hybrid detector. Gain scaling factors are still required as the simulated gain curves are known to underestimate the (real) measured gains. The obtained scaling factors are consistent within each measurement and between both detector setups. The calculated gains show a linear trend which can not be observed in the data. This discrepancies can be traced back to small deviations between the simulated gain curves and the measured voltage dependencies within the GEM1-Micromegas scans. Indeed already minor deviations of the slopes (less than a percent) can lead to the observed deviations. The discrepancies are stronger in case of the Yale hybrid detector, since the dominating gain curve of the Micromegas is known in case of the Bonn setup. The geometric displacement between both standard-pitch GEM foils turns out to be consistent within the measurements and between both detector setups. With approximately 50 – 60 µm, the displacement is comparable to three-dimensional calculations of the hexagonal structure for large-sized GEM foils [42]. The energy resolutions of the ALICE S-LP-LP-S configuration can be well described by the model calculations. As the drift ion collection efficiencies are only calculated for standard-standard GEM pitch configurations (tuning factors are still functions of the GEM types), smaller discrepancies can be found in case of the ion backflow. Like for the hybrid detectors, linear deviations can be observed for the gain curves.

The model calculations of the electron efficiencies, the energy resolution and the gain have been implemented in the ALICE O² framework which will be used by ALICE at CERN for Run 3 and Run 4. The implementations allow to calculate immediately these properties as functions of the electrostatic configuration (electric fields, GEM potentials) as well as the geometric setup (GEM types, distances).
Outlook

The hybrid detector has been investigated in terms of the energy resolution, the ion backflow as well as the gain and competes with the S-LP-LP-S baseline solution for the upgrade of the ALICE Time Projection Chamber. A further important aspect is the discharge probability and the rate stability. Although studies of a 2GEM+MM setup at the Super Proton Synchrotron (SPS) and the Proton Synchrotron (PS) at CERN yield a reduced discharge stability compared to the quadruple GEM stack, an experimental investigation has not been carried out in case of the Bonn hybrid detector.

The origin of the broad exponential background component as it can be seen in the recorded MCA spectra is not known yet. Copper excitations and cosmics can be excluded. The background might be caused by intrinsic electronic noise of the hybrid detector itself as it disappears once the setup has been disconnected from the readout chain and the data acquisition.

The constant charge density distribution of the GEM electrodes is a strong assumption of the two-dimensional efficiency calculations. ANSYS calculations showed that the charges accumulate at the rims of the GEM holes. Instead of assuming a single step function to describe the electrodes between the GEM holes, multiple step functions with different charge amplitudes can be considered. The ratios of the amplitudes are thereupon modeled according to the outcome of the ANSYS simulations. Within iterating steps, a more realistic charge density distribution of higher orders might be approached which approximates the real charge distribution. A further promising ansatz to describe the charge distribution in a continuous way is given in [121] where the surface charge has been derived in case of a conducting circular disk with a fixed electric potential. The introduced radial behavior of the charge distribution $\sigma(r) \propto \left(R^2 - r^2\right)^{-1/2}$ (where $R$ is the radius of the disc, i.e. the diameter corresponds to the distance $(w - L)/2$ between two GEM holes in the model calculations) shows the observed charge accumulations of the ANSYS calculations and leads to constant potentials on the disc electrodes.

The ion-backflow model relies strongly on the simulated ion efficiencies. Two-dimensional calculations of the drift ion collection efficiencies (for two GEMs with equal pitches) already indicate that a proper and full description turns out to be complex which is still an open issue. Dedicated studies and particularly simulations are still needed in order to fully understand the drift ion efficiencies and to extend the model calculation class with more realistic efficiency curves.

The calculated values of the energy-resolution model are strongly biased by the curves of the single gain fluctuations. Up to now an empirical function has been implemented which is motivated by the
outcome of simulations. However a more detailed investigation of the single gain fluctuations might lead to more optimized functional expressions. Additionally scalings of the gains are still required as the simulations underestimate the real gains. This issue is already known for Garfield++, yet not understood. Up to now only a constant attachment factor is globally assumed for the whole GEM stack which is manually specified. In fact, the attachment factor depends on the gas, the amount of impurities (water and oxygen content) and the applied electric field. An advanced implementation of the attachment requires a detailed simulation of the whole parameter space in order to obtain suitable lookup tables. Alternatively, an instance of Garfield++ could be included in the C++ class of the model calculations, allowing to calculate the attachment factors on-the-fly.

Up to now only the gain as well as the energy-resolution model have been implemented in the O² framework of the ALICE experiment. As the treatment of the drift ion efficiencies is still an open issue, no ion-backflow model has been implemented yet. The energy-resolution model allows to calculate $\sigma/\mu$ for the $^{55}$Fe photo peak. A translation back to the $dE/dx$ energy resolution for Pb-Pb and pp collisions is still an open issue.

Although still adjustments are required for the models (e.g. gain scaling, GEM displacement or attachment), the calculations open the door to a huge field of detailed investigations of MPGD and trends / relative changes can be reproduced. The influence of each stage can be quickly studied. The efficiencies and the contributions of the single stages to the whole detector can be examined. How is the gain composited? How and at which stages do the electrons or ions contribute to the ion backflow or the energy resolution? The model calculations offer great opportunities for in-depth studies of the processes within multiple amplification stages, making the dynamics inside of Micropattern Gaseous Detectors much more accessible and transparent.
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**Appendix**

**A.1 Source code: Bonn Hybrid Detector**

Example C++ source code to calculate the energy resolution, the ion backflow and the total effective gain for the hybrid 2GEM–MM detector (in case of $E_{TF2} = 400 \text{ V cm}^{-1}$). Additional functions are used in order to study the stack in details.

```cpp
#include "GEMStack.h"

void HybridDetector ()
{
    // Define stack and gas properties
    GEMStack * Detector = new GEMStack();
    Detector->SetPhotonEnergy(5900);
    Detector->SetGas(GAS_NECO2);
    Detector->SetAttachmentFactor(0.04);
    Detector->SetAttachment(1);

    // Define stack geometry and potentials
    Detector->AddVolume(2.6, 400.0);
    int GEM1 = Detector->AddGEM(GEM_ST, 270.0);
    Detector->AddVolume(0.2, 3000.0);
    int GEM2 = Detector->AddGEM(GEM_ST, 350.0);
    Detector->AddVolume(0.4, 400.0);
    int MM = Detector->AddGEM(GEM_ST, 370.0);
    Detector->AddVolume(0.0, 0.0);

    // Overwrite properties of last "GEM" to those of a MM
    Detector->SetGEMElectronCollection(MM, 0.99);
    Detector->SetGEMElectronExtraction(MM, 1.0);
    Detector->SetGEMAvalancheIonCollection(MM, 0.99);
    Detector->SetGEMAvalancheIonExtraction(MM, 0.01);
    Detector->SetGEMAbsGain(MM, 0.02195, -2.116);
    Detector->SetGEMSINGLEGainFluctuation(MM, 0.8);

    // Set models for calculations
    Detector->SetSingleGainFluctuationModel(MODEL_EXP_NOLIMIT);
    Detector->SetDriftIonModel(MODEL_DRIFT_ION_CALC);
}
```
// Tune total gain and drift ion collection efficiency
Detector->SetAbsGainScalingFactor(1.0);
Detector->SetDriftIonGEMsDisplacement(52.0);

// Get stack properties
double EnergyResolution = Detector->GetEnergyResolution();
double IonBackflow = Detector->GetIonBackflow();
double EffGain = Detector->GetTotalEffGain();

// Effective or absolute gain of a single stage
double EffGainGEM1 = Detector->GetGEMEffGain(GEM1);
double AbsGainGEM1 = Detector->GetGEMAbsGain(GEM1);

// Epsilon contribution of second GEM
double EpsilonGEM2 = Detector->GetGEMEpsilonContribution(GEM2);

// A simple way to plot the efficiencies of the first GEM
TCanvas *c = new TCanvas("c","c",0,0,1024,768);
Detector->DrawGEMEfficiencyCurves(GEM1, c, 1, 1, 1);
}

A.2 Source code: S-LP-LP-S configuration with the ALICE O² framework

using namespace o2::TPC;

#include <vector>

void CalcStackSettings()
{
ModelGEM Model;

std::array<int, 4> mGeometry;
std::array<float, 5> mDistance;
std::array<float, 4> mPotential;
std::array<float, 5> mElectricField;

mGeometry[...];
mDistance[...];
mPotential[...];
mElectricField[...];

Model.setAttachment(0.06);
Model.setAbsGainScalingFactor(1.53);
Model.setStackProperties(mGeometry, mDistance, mPotential, mElectricField);

std::cout << Model.getStackEnergyResolution() * 100.0 << std::endl;
std::cout << Model.getStackEffectiveGain() << std::endl;
}
A.3 Documentation of the slow control

A.3.1 Connection panel

The slow control was developed using LabView 2014 (Version 14.0 32-bit). Since the used binaries for the SNMP communication with the MPOD modules are only available in 32-bit architecture, the version of LabView should be chosen to be of 32-bit too. Running a 64-bit architecture of LabView will cause problems with the MPOD communication. All relevant information to start the communication with the MPOD modules and the Bonn pA-meters are specified in the connection panel (see Fig. A.1):

- **IP address**: Specifies the IP address of the MPOD controller unit.

- **Path to SNMP binaries**: Folder where the binaries (32-bit architecture) for the SNMP communication with the MPOD crate are located. Commonly this folder should be located in the directory of the slow control.

- **Wiener MIB file path**: Folder where the required MIB file is located in order to communicate with a WIENER MPOD crate. Commonly this folder should be located in the directory of the slow control.

- **UDL File for pSQL**: The communication with the Bonn pA-meters is done via a pSQL database. This file includes all relevant information to connect to the database (such as the name of the database, login information, password). A new UDL file (for a new database) can be created within LabView by choosing **Tools → Create Data Link → Microsoft OLE DB Provider for SQL server** and following the instructions.

- **Working directory**: Folder where all measurements will be saved.

- **Connect**: Connect to MPOD crate. In case of an established connection the LED will turn to green and connection information will be listed in the log panel.

![Figure A.1: Connection panel of slow control.](image)

A.3.2 MPOD: High voltage status panel

This panel indicates the status of operation for the connected channels of the MPOD modules. MPOD channels can be edited or added by clicking on the channel field (Ch.) and entering the appropriate channel number before connecting to the MPOD crate (see A.3.1). Following channel information are available:

- **Ch.**: Number of MPOD channel.

- **Stat**: Indicates if channel is either on (light green) or off (dark green).

- **Set U/V**: The voltage which has been set for the channel in Volt.

- **Read U/V**: The actual voltage as it is read by the MPOD controller unit in Volt.

- **I/nA**: The actual current as it is read by the MPOD controller unit in nA.
Appendix A  Appendix

- **Time**: The (estimated) remaining time to reach the set voltage in case channel is ramping. Note: The time is only calculated using the read voltage, the set voltage and the specified ramping speed, i.e. if the read voltage strongly differs from the set voltage (as example due to a short circuit), a remaining time might stay.

- **Rmp**: Indicates if channel is in ramping mode (light green).

- **Trip/mA**: Set trip current for the channel in mA.

- **Spd**: Set ramping speed for the channel in V/s.

- **Trip**: Red LED indicates trip in module or channel (depending on specified trip behavior, see A.3.3).

![Figure A.2: Information about the MPOD high voltage channels.](image)

**A.3.3 MPOD: Trip panel**

Specifies the behavior of the MPOD channels / modules in case of trips, i.e. if the currents exceed a user-defined threshold (see Fig. [A.3]). Settings to specify the trip behavior are:

- **Kill enable** (red LED): Hardware behavior can not be modified. The trip currents are set for the specified channels using the set button.

- **Kill disable** (LED off): Hardware behavior can be modified. The trip behavior (do nothing, ramp down channel, shut off channel, shut off module) can be specified for the channels using the set button. In order to prevent immediate actions, a trip time (16 – 4 000 ms) can be defined. The overcurrent has to last longer than this time before actions are taken.

Use clear events in order to reset the error states in the MPOD high voltage status panel (see A.3.2).

Note: The trip currents can also be set for all channels using "*" as input.

Important: Not all modules support these settings. Make sure to check this before blindly relying on the trip behavior.

**A.3.4 MPOD: Log panel**

If Log Voltages and Currents is ticked, the currents as well as the voltages of the MPOD channels will be logged to the specified file. The log file will be located in the working directory (see A.3.1).
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Figure A.3: Information about the MPOD trip behavior.

Figure A.4: Log currents and voltages as they are read out by the MPOD controller.

A.3.5 pA-meter: Zagreb panel

Indicates the currents as they are measured by the pA-meters "PicoLogic PA125" from Zagreb. Channels can be modified / added by clicking on the channel field (Ch) before connecting to the MPOD crate (see A.3.1). The unit is defined by display format given in the general pA-meters panel (see A.3.7).

The communication with the pA-meters is done using the National Instruments Distributed System Manager (NI-DSM, see Fig. A.6): The pA-meter software provided by PicoLogic stores the measured values to the shared variables Channel1 to Channel24 which are thereupon read out by the slow control. Before setting up a slow control on a new machine make sure these shared variables exist. Otherwise they can be created using the Distributed System Manager.

A.3.6 pA-meter: Bonn panel

The Bonn pA-meters are read out by the software PAMOS. This software allows to store the measured data in a pSQL database. The slow control is capable of connecting to this database (see A.3.1) for details) and to poll the stored information. The station numbers of the pA-meters can be edited / added by modifying the source fields in the Bonn pA-meter panel before connecting to the MPOD crate (see A.3.1). Following information can be obtained:

- **Current I**: Currents as they are recorded by the pA-meters. The unit is defined by display format given in the general pA-meters panel (see A.3.7).
Appendix A Appendix

Figure A.6: Shared variables listed in the NI Distributed System Manager in order to communicate with the pA-meter “PicoLogic PA125”.

- **Error current dl**: Error of the currents as they are recorded by the pA-meters. The unit is defined by display format given in the general pA-meters panel (see A.3.7).
- **M**: The mode in which the pA-meter is being operated.
- **B**: The battery status of the operated pA-meter.
- **Cflag**: The continue flag of a pA-meter. As long as the flag is set (light green) the pA-meter continues to record and store data. Once deactivated, the readout of the station will stop.

Figure A.7: Currents as they are read out by the pA-meters from Bonn.

A.3.7 pA-meter: General panel

This panel manages the communication with the Bonn as well as the Zagreb pA-meters and records data to the specified working directory (see A.3.1) if wanted. The options are:

- **Display format**: Sets the displayed units in the data panels of the pA-meters.
- **Use pA-meter (Zagreb)**: If ticked the defined channels of the Zagreb pA-meter will be updated.
- **Use pA-meter (Bonn)**: If ticked the defined channels of the Bonn pA-meters will be updated.
• **Station:** Allows to set the CFlag (continue flag) of a specific station number.

• **Save currents:** If selected the currents of all channels from the specific source (dropdown menu) will be written to the file (located in working directory, see \[A.3.1\]). In case of the Zagreb pA-meter also the MP0D voltages and currents will be logged for the specified channel next to the dropdown menu (here channel 401).

![Figure A.8: Currents as they are read out by the pA-meters from Bonn.](image)

**A.3.8 Ramping, stack configuration and channel mapping**

The ramping panel (see Fig. [A.9]) allows to ramp individual high voltage channels with a user-defined ramp speed. The potential can be set by the ramp voltage field for a single, multiple (separated by comma, e.g. "401, 402, 405") or all (indicated by "+") channels. Unaffected by the channels field, all high voltage channels can be switched off using the all off button which ramps down the channels (the ramp speed is not overwritten by this procedure). Some HV modules technically require a negative ramp voltage during their operation which can be set by the Neg. Ramp Speed checkbox.

In case of multiple amplification stages (Micro-Pattern Gaseous Detectors (MPGD) like GEM foils, Micromegas,...) the electrostatic and geometric properties of the stack can be defined using the Stack Settings cluster. The checkbox 4 GEMs allows to switch between a triple and a quadruple GEM stack configuration. The layout of the GUI will change accordingly. In addition a Last Strip and a Skirt electrode can be added to the detector settings. The potentials are thereupon calculated according to the defined stack settings once Ramp it! is pressed. A scaling factor linearly scales the potentials for tuning and adjusting. The mapping of the channels can be specified using the Channel Mapping cluster.

![Figure A.9: Ramping panel in order to set the potential of high voltage channels.](image)

**A.3.9 Multichannel Analyzer (MCA)**

The communication with the "Pocket MCA8000D" is based on the API / LabView SDK provided by Amptek. Fig. [A.11] shows the MCA panel as it is used to communicate with the MCA:

• **COM Port:** Specifies the COM port where the MCA is plugged in (check Microsoft Windows Device Manager if not known).
• **Threshold:** Sets the channel where the MCA spectrum should be cut.

• **Acquisition time:** Sets the acquisition time for data taking. The MCA would stop recording data once this time elapsed.

• **ADC Channels:** Selects how many ADC channels should be used.

• **Control buttons:** Allows to start or to stop a running measurement. The MCA data buffer can be cleared using the reset button.

• **Get Data:** Reads and plots the current recorded MCA spectrum.

• **File and Save:** Allows to save the recorded MCA spectrum to an ASCII file in the current working directory (see [A.3.1]) once Get Data has been used.

### A.3.10 X-Ray source

The slow control is capable of operating X-ray sources from Amptek (Mini-X) using the provided API / LabView SDK given by the manufacturer. Fig. [A.12] shows the control panel. Make sure the X-ray tube is plugged in correctly (USB) and that the Windows driver is installed. Press connect to start the communication. Once a connection has been established the Connected LED will light up, allowing to set the emission current \( I \) field (5-200 µA) and the high voltage \( U \) field (10-40 kV). The grayed out fields
indicate the actual measured value in case the Mini-X is in operation. As soon as the interlock of the X-ray source has been set, the Interlock LED will light up and the X-ray can be switched on (and off) by the HV ON checkbox. The red LED X-ray on will light up.

Figure A.12: Panel to communicate with Amptek Miniature X-Ray source (Mini-X).

A.3.11 Automation of jobs / automation scripting

A big advantage of the slow control is the ability to perform automated measurements. These jobs (e.g. ramping, starting the Mini-X or simply idling for a certain time) are executed successively from top (job 0) to bottom. Fig. A.13 shows the panel to program and execute the automated procedures: Currently job 0 (command "SetDir") is about to be executed once the automation is started by the Start automation checkbox, followed by the next job (command "RampV") and so on. The actual job is always indicated by the green LED and the Current job field. Additional information of the ongoing job are always given in the Job Status Info and the log output. The automation can be interrupted by deactivating the Start automation checkbox. The current job will continue till it is done and the automation stops.

A complete list of the available commands and example scripts are given in the following pages. Some commands require additional parameters (fields Parameter 1, Parameter 2 and Parameter 3), e.g. time, voltages or channels. The automation can be programmed using the graphical user interface or by loading automation scripts with the Load button.
List of commands

The following list includes all available jobs (command, parameter 1, parameter 2, parameter 3) and a description for the automation (Slow Control v5.4). Note: Command and parameters are separated by tabulator within the script files.

- **RampV, VOLTAGE, SPEED, CHANNELS**

  Ramp channel/channels as defined in CHANNELS to specific VOLTAGE (in Volt) with ramp SPEED (in Volt/s).

- **4 GEM setup:**
  RampS, LAST_SKIRT_NGEM_SPEED_SCALING, DRIFT_TF1_TF2_TF3_CF_GEM1_GEM2_GEM3_GEM4, DRIFTGEM1_LASTGEM1_SKIRTGEM1_GEM1_GEM2_GEM2_GEM3_GEM3PADS

  3 GEM setup:
  RampS, LAST_SKIRT_NGEM_SPEED_SCALING, DRIFT_TF1_TF2_CF_GEM1_GEM2_GEM3, DRIFTGEM1_LASTGEM1_SKIRTGEM1_GEM1_GEM2_GEM2_GEM3_GEM3PADS

  Ramp to specific stack settings. LAST (parameter 1) defines if last strip should be used (0 no, 1 yes). SKIRT (parameter 1) defines if skirt should be used (0 no, 1 yes). NGEM defines number of GEMS (3 or 4), SPEED the ramping speed (Volt/s), SCALING defines the scaling factor (0.0 – 1.0). Parameter 2 and 3 depend on NGEM. Parameter 2 defines all fields (in Volt/cm) and all GEM voltages (in Volt). Parameter 3 defines all distances (in mm).

- **RampDone**

  Wait till all channels have ramped (ramp status LED is off). Note: Before using RampDone it is recommended to let the Slow Control idle for some seconds (e.g. 10 s) since it takes some time to update the ramp status of the channels after changing to new settings.

- **Idle, TIME**

  Idle and do nothing for specified TIME (in seconds).
• **StartC, FILENAME, PAMETER**

Starts a currents measurement and saves the data in the file FILENAME (within the working directory). PAMETER defines which pA-meter should be used (i.e. "Zagreb" or "Bonn"). The output format of the saved currents is defined by "Display Format" in the GUI. Note: This command does not activate "Use pA-Meter (...)" option in the GUI. It just defines the path, filename, sets the source and starts the measurement.

• **StopC**

Stops an ongoing current measurement.

• **Stop**

End of the automation script.

• **SetCFlag, STATIONNO, FLAG**

Set the continue FLAG (0 or 1) of a Bonn pA-meter with station number STATIONNO.

• **SetDir, DIR**

Set the working directory to DIR.

Note: The path should not end with "\", e.g. "C: \ working_directory" is correct.

• **GoTo, JOBNO**

Jump to a specific job with number JOBNO (counting starts with 0 for the first job).

• **MiniXOn**

Switches Mini-X on (requires a connected device and a closed interlock, otherwise this command will be skipped).

• **MiniXOff**

Switches Mini-X off (requires a connected device and a closed interlock, otherwise this command will be skipped).

• **MiniXSetUI, VOLTAGE, CURRENT**

Set the high VOLTAGE (value between 10 – 40 kV) and the emission CURRENT (value between 5 – 200 µA). This requires a connected device and a closed interlock, otherwise this command will be skipped. This command can only be executed when the HV is switched off.

• **MPODLogStart, FILENAME**

Start logging of MPOD voltages and currents to file specified by FILENAME in current working directory.
Appendix A  Appendix

- **MPODLogStop**
  Stop logging of MPOD voltages and currents.

- **MCASetChannels, CHANNELS**
  Set the amount of MCA CHANNELS (must be 256, 512, 1024, 2048, 4096, 8192, 16384).

- **MCASetTime, TIME**
  Set the data acquisition TIME (in seconds).

- **MCASetThreshold, CHANNEL**
  Set the threshold (CHANNEL must be >= 1).

- **MCAStrart**
  Start data taking with the MCA.

- **MCASstop**
  Stop data taking with the MCA.

- **MCAReset**
  Clear recorded MCA spectrum and reset timer.

- **MCAGetData**
  Receive recorded data from MCA (depending on number of channels this might take some time).

- **MCASave, FILENAME**
  Save recorded and received MCA spectrum to FILENAME in current working directory (MCAGetData must be called before MCASave can save any data). If no FILENAME is specified, an output file with default name "MMDDYYYY_HHMMSS.txt" will be created (starting time of the measurement).

- **ShutterIn**
  Moves the shutter counter-clockwise to shield the beam of the Mini-X.

- **ShutterOut**
  Moves the shutter clockwise to let the beam of the Mini-X pass through.

**Example: Working with Bonn pA-meters**

The following script ramps channel 201 and 202 with 5 V s^{-1} to 200 V. To give the ramping status LEDs some time to update it waits for 5 seconds. Finally it waits till all channels have finished ramping. As the
A.3 Documentation of the slow control

currents might need some time to stabilize, the automation idles for some additional 30 seconds. The currents are recorded using the Bonn pA-Meters and written to file "Example_Measurement.txt" in the working directory "C:\my_working_directory". After 180 seconds the script stops the measurement. All channels will be ramped down to 0 V with 10 V s\(^{-1}\). Finally the pA-Meter with station number 7 will be deactivated and the script stops.

<table>
<thead>
<tr>
<th>Command</th>
<th>Value 1</th>
<th>Value 2</th>
<th>Value 3</th>
</tr>
</thead>
<tbody>
<tr>
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<td>200</td>
<td>5</td>
<td>201,202</td>
</tr>
<tr>
<td>Idle</td>
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<td></td>
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<td>RampDone</td>
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</tr>
<tr>
<td>Idle</td>
<td>30</td>
<td></td>
<td></td>
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<td>SetDir</td>
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<td>StartC</td>
<td>Example_Measurement.txt Bonn</td>
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<td></td>
</tr>
<tr>
<td>Idle</td>
<td>180</td>
<td></td>
<td></td>
</tr>
<tr>
<td>StopC</td>
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<td></td>
<td></td>
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<tr>
<td>RampV</td>
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<td>10</td>
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<td>SetCFlag</td>
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<td>Stop</td>
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</table>

**Example: Quality assurance for ALICE**

The script has been used for the high voltage measurements in the framework of the QA program of the ALICE TPC upgrade in Bonn.

<table>
<thead>
<tr>
<th>Command</th>
<th>Value 1</th>
<th>Value 2</th>
<th>Value 3</th>
</tr>
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