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Electron swarm parameters of the hydrofluoroolefine HFO1234ze

A. Chachereau\(^1\), M. Rabie\(^1\), C. M. Franck\(^1\)

\(^1\)Power Systems and High Voltage Laboratories, ETH Zurich, Physikstr. 3, 8092 Zurich, Switzerland
E-mail: alisec@ethz.ch

Abstract. In this contribution, the electron swarm parameters of the hydrofluoroolefine HFO1234ze, systematic name 1,3,3,3-Tetrafluoro-1-propene, are experimentally investigated. The analysis of the electron avalanche current measured in a pulsed Townsend experiment yields the effective ionization rate coefficient, the electron drift velocity and the longitudinal electron diffusion coefficient. The subsequent ion current is analyzed as well, to obtain separately the ionization and attachment rate coefficients. Measurements in pure HFO1234ze at different pressures show that the effective ionization rate is strongly influenced by three-body attachment and the three-body attachment rate coefficient is derived.

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Introduction

The hydrofluoroolefine HFO1234ze (CF\(_3\)CH=CHF) is a refrigerant gas which was proposed as a replacement to the hydrofluorocarbon R134a (CF\(_3\)CH\(_2\)F) with a lower global warming potential (GWP) [1]. Recently HFO1234ze also attracted attention as a potential alternative to SF\(_6\) for gaseous high voltage insulation [2, 3, 4]. It has a similar molecular structure as hexafluoropropylene (CF\(_3\)CF=CF\(_2\)), which has a comparable electric strength to SF\(_6\) [5, 6, 7]. It is expected that the electric strength of HFO1234ze is lower than that of 1-C\(_3\)F\(_6\) but still substantial. Moreover, HFO1234ze has an extremely low global warming potential on a hundred years horizon (\(<1\) ), compared to 23’500 for SF\(_6\) [8].

In the present work, the electron swarm parameters of pure HFO1234ze and of several diluted mixtures of HFO1234ze with N\(_2\), CO\(_2\) and Ar are investigated in a pulsed
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Townsend experiment. Part of these measurements were used for making breakdown voltage predictions in HFO1234ze [9]. The results are available on the online database ETHZ [10].

In the first section, the experimental setup and the method for analysing the measured current are briefly described. The obtained electron swarm parameters are presented in section 2. The effective ionization rate coefficient, the electron drift velocity and the longitudinal electron diffusion coefficient are obtained in pure HFO1234ze at different gas pressures. At one of these pressures, the ion currents in pure HFO are recorded and analyzed to obtain separately the ionization and attachment rate coefficients. The swarm parameters are also obtained in several diluted mixtures of HFO1234ze with N₂, CO₂ and Ar at the total pressure of 10 kPa. In the third section of this work, the pressure dependence of effective ionization rate coefficient in pure HFO1234ze is investigated using a kinetic model for three-body electron attachment, and the electron attachment properties of HFO1234ze are discussed in view of the swarm parameters of the diluted mixtures of HFO1234ze with N₂, CO₂ and Ar.

1. Methods

1.1. Swarm experiment

The pulsed Townsend experimental setup used in this work has been described previously [11]. A short laser pulse (1.5 ns FWHM) releases about 10⁷ electrons from a back-illuminated photocathode. The electron swarm drifts in a uniform electric field through a gap between two Rogowski type electrodes. The gap distance is varied in the range of 11 to 21 mm and the applied voltage is up to 40 kV. The measurements are made at room temperature, and in a pressure range of 2 to 45 kPa. The transport parameters depend on the reduced field strength \( E/N \), given in Townsend (1 Td = 10⁻²¹ Vm²). Here, \( E \) is the electrical field strength and \( N \) is the number density of the gas. The displacement current \( I_{\text{exp}} \) of the charged particle swarm drifting in the gap is recorded and evaluated. Sample current measurements are shown in figure 1.

1.2. Electron and Ion swarm analysis

The current is measured on two different timescales, namely during the electron transit time and during the ion transit time, see figure 2. The electron-dominated current of the electron swarm released by the laser pulse is followed by a current that is dominated by the ions that were produced during the electron transit. Ions are about two to three orders of magnitude slower than electrons, the ion current amplitude is consequently lower and the ion transit longer. The details of the electron and ion swarm analysis used in this work can be found in the appendix. The electron current analysis yields the effective ionization rate \( \nu_{\text{eff}} \), the electron drift velocity \( v_e \) and the longitudinal electron diffusion coefficient \( D_L \). The total ionization rate \( \nu_i \) and the total attachment rate \( \nu_a \) such that \( \nu_{\text{eff}} = \nu_i - \nu_a \), as well as the drift velocities for anions \( v_n \) and cations \( v_p \), are
obtained from the subsequent ion current analysis. This approach considers one drift velocity for the anions $v_n$ and one for the cations $v_p$, whereas the possible presence of additional ion species, reaction between species or clusters formation is not considered. In the case of several anions, as in the present work, $v_n$ must be considered as an averaged drift velocity over all anion species.
Figure 2: Current versus time in HFO1234ze at the pressure \( p = 3 \text{kPa} \) and for an electrode gap distance \( d = 15 \text{mm} \). (a) Electron current and (b) ion current at \( E/N = 180 \text{Td} \). (c) Electron current and (d) ion current in HFO1234ze at \( E/N = 210 \text{Td} \). The thin lines are the measured currents whereas the thick lines correspond to the models of electron and ion swarms described in the appendix 3.4 and 3.5, in particular \( I_p \) is the positive ion current, \( I_n \) is the negative ion current and \( I_{\text{ion}} \) is the total ion current. The times \( T_1, T_2, T_3 \) and \( T_e \) are defined in the appendix 3.6.

2. Results

2.1. HFO1234ze at different pressures

The effective ionization rate coefficient \( k_{\text{eff}} = \nu_{\text{eff}}/N \) in HFO1234ze has been obtained at pressures ranging from 3 to 45 kPa. The values of \( k_{\text{eff}} \) are shown in figure 3 as a function of \( E/N \). A strong pressure dependence of \( k_{\text{eff}} \) is observed over the whole \( E/N \) range. In particular, the density reduced critical electric field \( (E/N)_{\text{crit}} \), for which \( k_{\text{eff}} \) equals zero, increases strongly with increasing gas pressure.

The electron drift velocity \( v_e \) and the density normalized longitudinal electron diffusion coefficient \( ND_{L} \) in HFO1234ze were obtained at pressures between 3 to 45 kPa.
Figure 3: Effective ionization rate coefficient in HFO1234ze at different gas pressures as a function of $E/N$.

and found to be independent of the gas pressure. Figure 4 shows the average and the standard deviation of $v_e$ and $N D_L$ over the different pressures, as a function of $E/N$. The same quantities are shown for $N_2$, CO$_2$ and Ar for comparison.
Figure 4: (a) Electron drift velocity $v_e$ and (b) density normalized longitudinal electron diffusion coefficient $ND_L$ in HFO1234ze, N$_2$, CO$_2$ and Ar as a function of $E/N$.

2.2. Separating ionization and attachment rate coefficients

At the pressure of 3 kPa, the ion currents in HFO1234ze were measured in addition to the electron currents. In contrast to the electron currents, the ion currents were obtained only for $E/N \geq 130$ Td, since for lower $E/N$ they were too small for being measured with the present instruments. Using the method described in the appendix 3.4, the total ionization and attachment rate coefficients $k_i = \nu_i/N$ and $k_a = \nu_a/N$ are obtained, as well as the average cation and anion drift velocities $v_p$ and $v_n$. The rate coefficients $k_i$ and $k_a$ are shown in figure 5 (a) and the ion drift velocities are shown in figure 5 (b). The total attachment rate coefficient $k_a$ increases slightly over the $E/N$ range 130 to 220 Td. The total ionization rate coefficient $k_i$ starts increasing at about 150 Td, below this value too few positive ions are present to obtain their drift velocity. The crossing point between $k_a$ and $k_i$ corresponds to the density reduced critical electric field $(E/N)_{\text{crit}} \simeq 198$ Td at the pressure of 3 kPa.
Figure 5: (a) Total attachment ($k_a$), total ionization ($k_i$), and effective ionization ($k_{eff}$) rate coefficients and (b) drift velocity of anions ($v_n$) and cations ($v_p$) in HFO1234ze at 3 kPa as a function of $E/N$.

2.3. Diluted mixtures of HFO1234ze in $N_2$, $CO_2$ and $Ar$

Figures 6, 7 and 8 show the swarm parameters of diluted mixtures of HFO1234ze in $N_2$, $CO_2$ and $Ar$ respectively, compared to measurements in the pure buffer gases $N_2$, $CO_2$ and $Ar$. In the HFO1234ze/$N_2$ and HFO1234ze/$CO_2$ mixtures, only slight changes in the swarm parameters are observed compared to pure $N_2$ and $CO_2$. In the HFO1234ze/$Ar$ mixtures, a strong change of the swarm parameters is observed compared to pure $Ar$. There is a strong increase in ionization in the HFO1234ze/$Ar$ mixtures compared to pure $Ar$, but only weak electron attachment is observed in the HFO1234ze/$Ar$ mixtures. The drift velocity in the HFO1234ze/$Ar$ mixtures differs from that in pure $Ar$, and it increases with increasing percentage of HFO1234ze in the investigated mixtures. For the density reduced electric fields $E/N \geq 14$ Td in the mixture with 0.2% HFO1234ze, and for $E/N \geq 20$ Td in the mixture with 0.9% HFO1234ze, an after-current is observed after the electron transit which compromises the derivation of the electron drift velocity and the diffusion coefficient.
Figure 6: (a) Effective ionization rate coefficient, (b) electron drift velocity and (c) density normalized longitudinal electron diffusion coefficient in pure N$_2$ and in the mixtures of 0.2% and 2% HFO1234ze in N$_2$ as a function of $E/N$. 

(a) $k_{\text{eff}}$ ($10^{-18}$ m$^3$ s$^{-1}$)

(b) $v_e$ ($10^4$ m s$^{-1}$)

(c) $N D_L$ ($10^{24}$ m$^{-1}$ s$^{-1}$)
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Figure 7: (a) Effective ionization rate coefficient, (b) electron drift velocity and (c) density normalized longitudinal electron diffusion coefficient in pure CO$_2$ and in the mixtures of 0.5% and 1.1% HFO1234ze in CO$_2$ as a function of $E/N$. 
Figure 8: (a) Effective ionization rate coefficient, (b) electron drift velocity and (c) density normalized longitudinal electron diffusion coefficient in pure Ar and in the mixtures of 0.2%, 0.9% and 2.8% HFO1234ze in Ar as a function of $E/N$. 
3. Discussion

3.1. Three-body electron attachment to HFO1234ze

The effective ionization rate coefficient $k_{\text{eff}}$ in pure HFO1234ze is clearly decreasing with increasing gas pressure. This pressure dependence has been observed as well in 1-C$_3$F$_6$ [6] and other fluorocarbon gases [12, 13] and is typically due to the occurrence of three-body electron attachment [14]. In this section, we consider the simple model for three-body attachment described in [16] and apply it to the present measurements.

The elementary attachment processes considered are the following

$$C_3H_2F_4 + e^{-} \xrightarrow{k_{\text{da}}} A^{-} + B \text{ (dissociative attachment),} \quad (1)$$

$$C_3H_2F_4 + e^{-} \xrightarrow{k_{\text{at}}} (C_3H_2F_4^-)^* \text{ (parent ion attachment),} \quad (2)$$

$$(C_3H_2F_4^-)^* + C_3H_2F_4 \xrightarrow{k_{\text{stab}}} C_3H_2F_4^- + C_3H_2F_4 \text{ (collisional stabilization),} \quad (3)$$

$$(C_3H_2F_4^-)^* + C_3H_2F_4 \xrightarrow{k_{\text{det}}} e^- + C_3H_2F_4 + C_3H_2F_4 \text{ (collisional detachment).} \quad (4)$$

where $k_{\text{da}}$ is the rate coefficient of dissociative attachment, $k_{\text{at}}$ is the rate coefficient of $(C_3H_2F_4^-)^*$ formation; $\tau^{-1}$ is the autodetachment rate from $(C_3H_2F_4^-)^*$; $k_{\text{stab}}$ and $k_{\text{det}}$ are the rate coefficients of collisional stabilization of $(C_3H_2F_4^-)^*$ and of collisional detachment from $(C_3H_2F_4^-)^*$. Additionally, HFO1234ze is ionized with a total ionization rate coefficient $k_i$ which includes dissociative and non-dissociative ionization.

This simple kinetic model is sufficient to reproduce the present experimental findings. Introducing additional kinetic processes involving for instance ion clusters cannot be justified based on these findings alone, but it is clear that such processes could occur. The problem of non-unicity of the kinetic model derived from swarm data is well known [15]. Complementary investigations of HFO1234ze based for instance on ion mass spectrometry would be required to establish with certitude the attachment mechanism to HFO1234ze as was done for 1-C$_3$F$_6$ [7].

Through the processes (2) and (4), the density of electrons is coupled with that of $(C_3H_2F_4^-)^*$. After a few $\tau$, the growth of the electron number is exponential, with the rate [16, 17, 18]

$$\nu_{\text{eff}}(N) = (k_i - k_{\text{da}})N - \frac{k_{\text{at}}k_{\text{stab}}N^2}{\tau^{-1} + (k_{\text{det}} + k_{\text{stab}})N}, \quad (5)$$

If we note

$$N_{\text{sat}} = ((k_{\text{stab}} + k_{\text{det}})\tau)^{-1}, \quad (6)$$

$$k_{\text{quad}} = k_{\text{at}}k_{\text{stab}}\tau, \quad (7)$$

equation (5) becomes

$$\nu_{\text{eff}}(N) = (k_i - k_{\text{da}})N - \frac{k_{\text{quad}}N^2}{1 + N/N_{\text{sat}}}, \quad (8)$$

Two limiting cases of equation (8) can be identified by comparing the gas density $N$ to the quantity $N_{\text{sat}}$ [16].
(i) When $N \ll N_{\text{sat}}$, equation (8) simplifies as
$$\nu_{\text{eff}}(N) = (k_i - k_{\text{da}})N - k_{\text{quad}}N^2,$$
and the three-body attachment rate increases quadratically with the gas density.

(ii) When $N \gg N_{\text{sat}}$, equation (8) simplifies as
$$\nu_{\text{eff}}(N) = (k_i - k_{\text{da}})N - k_{\text{quad}}N_{\text{sat}}N,$$
and the three-body attachment rate increases linearly with the gas density.

The quantity $N_{\text{sat}}$ can be seen as a "saturation" density for three-body attachment. Indeed, when the gas density greatly exceeds $N_{\text{sat}}$, the autodetachment lifetime $\tau$ is much larger than the mean time for the unstable anion to collide with a third body ($\tau \gg ((k_{\text{stab}} + k_{\text{det}})N)^{-1}$), therefore, the unstable anion is systematically collisionally stabilized/detached whereas the autodetachment becomes irrelevant. In this case, three-body attachment appears outwardly as a two-body process, with the apparent two-body rate coefficient $k_{\text{quad}}N_{\text{sat}}$.

The present measurements of the effective ionization rate in HFO1234ze are clearly obeying equation (9), as shown for some sample $E/N$ values in figure 9. Thus, they correspond to the case (i) where $N \ll N_{\text{sat}}$. The quantities $(k_i - k_{\text{da}})$ and $k_{\text{quad}}$ are obtained as the coefficients from the linear regression of $\nu_{\text{eff}}(N)$ on $N$ and $N^2$. The quantity $N_{\text{sat}}$ cannot be obtained from the present measurements as it does not appear in equation (9). Examples of the regression of equation (9) for sample $E/N$ values are shown in figure 9. The rate coefficients $(k_i - k_{\text{da}})$ and $k_{\text{quad}}$ are shown in figure 10 (a) and (b). The three-body attachment rate $k_{\text{quad}}$ reaches a maximum at 120 Td, then slowly decreases with increasing $E/N$.

The analysis developed here is independent from the ion current analysis that was presented in section 1.2. For comparison, the total attachment rate $k_a$ that was obtained with the ion current analysis at the pressure $p = 3$ kPa would correspond here to
$$k_a(N_1) = k_{\text{da}} + k_{\text{quad}}N_1,$$
(11)
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Figure 10: (a) Two-body rate coefficient \((k_i - k_{da})\) and (b) three-body attachment rate coefficient \(k_{quad}\) in HFO1234ze as a function of \(E/N\). The points are determined using the regression (9), the lines are fits used in figure 11.

where \(N_1\) is the gas number density at the pressure \(p = 3\) kPa and at room temperature.

Using the quantities \((k_i - k_{da})\) and \(k_{quad}\), the rate coefficient \(k_{eff}\) can be calculated at different gas pressures with equation (9). These calculations are plotted in figure 11 at the same pressures where measurements in pure HFO1234ze were performed.

Figure 11: Effective ionization rate coefficient in HFO1234ze at different gas pressures as a function of \(E/N\). The markers are measurement points that were presented in figure 3, whereas the lines are calculated from \((k_i - k_{da})\) and \(k_{quad}\), in particular line for \(p \rightarrow 0\) corresponds to the rate coefficient for two-body processes \((k_i - k_{da})\).
Since \( N_{\text{sat}} \) could not be obtained from the present measurements, it is uncertain up to which pressure the three-body attachment rate will continue increasing quadratically with the gas density following equation (9). At higher pressures, three-body attachment will eventually reach "saturation" and \( \nu_{\text{eff}} \) will depend linearly on the gas pressure according to equation (10). Therefore equation (9) should not be used to extrapolate the present data towards higher gas pressures than where measured. However, the present considerations give the limit of \( k_{\text{eff}} \) towards low pressures, where three-body attachment is negligible. This limit corresponds to the term \( (k_i - k_{\text{da}}) \).

3.2. Diluted mixtures of HFO1234ze in \( N_2 \) and \( CO_2 \)

The addition of small percentages of HFO1234ze in \( N_2 \) or \( CO_2 \) barely affects the effective ionization rate coefficient shown in figures 6 and 7, whereas pure HFO1234ze proved strongly electron attaching, with a density reduced critical electric field \( \geq 188 \text{Td} \) as shown in figure 3. For strongly attaching gases, it is rather common to observe a strong change in \( k_{\text{eff}} \) relative to the pure buffer gas even in such diluted mixtures [18]. The absence of this change proves that, in the investigated \( E/N \) range, the attachment cross section of HFO1234ze has little overlap with the electron energy distribution functions in \( N_2 \) and \( CO_2 \). Thus, attachment to HFO1234ze would occur at relatively high electron energies. This is usually the case for dissociative electron attachment, but is less common for parent ion attachment.

3.3. Diluted mixture of HFO1234ze in Ar

The addition of a small percentage of HFO1234ze in Ar causes a strong change of the electron velocity at low \( E/N \) (see figure 8). At low \( E/N \), the drift velocity in the HFO1234ze/Ar mixtures is decreasing with increasing \( E/N \), and is higher than the drift velocity in either Ar or HFO1234ze. These phenomena, often referred to as negative differential mobility or negative differential conductivity, have been studied extensively [19, 21, 20] and are considered common for diluted mixtures of molecular gases in Ar.

The effective ionization rate coefficient is strongly affected by a small addition of HFO1234ze in Ar, see figure 8. For the most diluted mixture, which is 0.2% HFO1234ze in Ar, a strong increase of ionization relative to pure Ar is observed. The increase of ionization could be due to Penning ionization of HFO1234ze by metastable excited Ar states. Penning ionization could in addition account for the after-current observed in some of the electron transients in the mixtures of 0.2% HFO1234ze in Ar and 0.9% HFO1234ze in Ar in the higher \( E/N \) range. For the mixtures with 0.9% and 2.8% HFO1234ze, some electron attachment is observed which partially compensates the increase in ionization.
Conclusion

The electron swarm parameters of HFO1234ze were determined experimentally for the first time. The strong pressure dependence of the effective ionization rate coefficient was investigated assuming a three-body attachment mechanism to HFO1234ze. This model described well the observed pressure dependence of the effective ionization rate, and the three-body attachment rate was obtained. The two-body processes of ionization and attachment represent the limit of the effective ionization rate coefficient in HFO1234ze towards low pressures. The attachment to HFO1234ze at higher pressures would require further investigation, but the strong electron attachment observed already at relatively low pressures make HFO1234ze a promising gas for electric insulation at higher pressures. Since HFO1234ze would liquefy at the typical pressures used in high voltage gaseous insulation equipments, HFO1234ze mixtures with suitable buffer gases should be investigated as well.

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Appendix

3.4. Model of electron swarms

In our experiment, a voltage $U$ is applied between two Rogowski type electrodes separated by a gap distance $d$, resulting in a homogenous electric field $E$ in the gap. At time $t = 0$, a short laser pulse releases $n_0$ photo-electrons from a photocathode into the gas of number density $N$. The electron swarm drifts, after a very short non-equilibrium phase, constantly in the direction of the anode. The measured current is a consequence of the drift of all charged particles (electrons and ions) in the gap. For this situation, equations for the current from electrons or ions have been obtained previously [22, 23, 5]. The electron swarm model used here is based on a Gaussian spatial distribution of the swarm electrons and the corresponding electron current can be expressed for $t \geq 0$ as [11]

$$I_e(t) = \frac{I_0}{2} \exp \left( \nu_{\text{eff}} t \right) \left( 1 - \text{erf} \left( \frac{t - T_e}{\sqrt{2} \tau_D} \right) \right), \quad (12)$$

$$I_0 = \frac{n_0 q_0}{T_e}, \quad (13)$$

where $q_0$ is the electron charge, $I_0$ is the electron current at time $t = 0$, $T_e$ is the drift time and $\nu_{\text{eff}}$ is the effective ionization rate. The canonical error function “erf” accounts for the absorption of the swarm at the anode. The characteristic time for longitudinal diffusion $\tau_D$ is related to the longitudinal diffusion coefficient $D_L$ via $2D_L = \nu_e^2 \tau_D$, where $\nu_e$ is the electron drift velocity. The initial distribution of the released electrons is
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assumed to be a Dirac delta. In case of strong electron detachment, cathodic feedback or photo-ionization by secondary photons, equation (12) cannot be used to derive precise values of $v_e$ and $\tau_D$ from our measured currents. However, it can be still applied to derive $\nu_{\text{eff}}$, as will be described in section 3.6.

3.5. Model of ion swarms

The total ion current is the sum of the currents arising from the drift of anions and cations. In this section, we assume that there is only one species of cations and one species of anions. Furthermore, we assume that the effective rate in equation (12) is given by the ionization minus the attachment rate $\nu_{\text{eff}} = \nu_i - \nu_a$.

3.5.1. During electron transit

Ions are treated as immobile during the electron transit, since electron drift velocities typically exceed ion drift velocities by up to three orders of magnitude. The charge densities for electrons $\rho_e(x,t)$, cations $\rho_p(x,t)$ and anions $\rho_n(x,t)$ obey

$$\frac{d}{dt}\rho_p(x,t) = \nu_i \rho_e(x,t),$$

(14)

$$\frac{d}{dt}\rho_n(x,t) = \nu_a \rho_e(x,t).$$

(15)

The currents associated with electron, cations and anions are

$$I_{e,p,n}(t) = \frac{v_{e,p,n}}{d} \int_0^d \rho_{e,p,n}(x,t) dx$$

(16)

where $v_{e,p,n}$ are the electron, cation and anion drift velocities. Thus, the spatial integration of equations (14) and (15) over the electrode gap yields

$$\frac{d}{dt}I_p(t) = \nu_i \frac{v_p}{v_e} I_e(t),$$

(17)

$$\frac{d}{dt}I_n(t) = \nu_a \frac{v_n}{v_e} I_e(t),$$

(18)

It follows that the total ion current $I_{\text{ion}}(t) = I_p(t) + I_n(t)$ is given by

$$I_{\text{ion}}(t) = \left( \nu_i \frac{v_p}{v_e} + \nu_a \frac{v_n}{v_e} \right) \int_0^t I_e(t') dt'.$$

(19)

3.5.2. After electron transit

For the calculation of the ion currents at times $t \geq T_e$ the ion charge densities at time $t = T_e$ are needed. To obtain them, a simpler model of the electron current during the interval $[0,T_e]$ is used than the one described in section 3.4. The diffusion of the electron swarm is set to zero, yielding the electron charge density

$$\rho_e(x,t) = n_0 \exp(\nu_{\text{eff}} t) \delta(x - v_e t).$$

(20)
According to equation (14) and (15) static cation and anion charge densities are created until time $T_e$. At time $T_e$, the ion charge densities in the gap are

$$\rho_p(x, T_e) = n_0 \frac{\nu_i}{v_e} \exp \left( \frac{\nu_i}{v_e} x \right),$$  \hspace{1cm} (21)

$$\rho_n(x, T_e) = n_0 \frac{\nu_n}{v_e} \exp \left( \frac{\nu_n}{v_e} x \right).$$  \hspace{1cm} (22)

After the electron transit, between $T_e$ and the transit times $T_p = d/v_p$ and $T_n = d/v_n$ of cations and anions, the currents originating from positive and negative ions can be expressed as a function of time $t' = t - T_e$ [23]:

$$I_p(t') = \frac{q_0}{T_p} \int_{v_p t'}^{d} \rho_p(x, T_e) dx = I_0 \frac{T_e}{T_p} \frac{\nu_i}{\nu_{eff}} \left( e^{\nu_{eff}T_e} - e^{\nu_{eff}T_p} \right),$$  \hspace{1cm} (23)

$$I_n(t') = \frac{q_0}{T_n} \int_{0}^{d-v_n t'} \rho_n(x, T_e) dx = I_0 \frac{T_e}{T_n} \frac{\nu_n}{\nu_{eff}} \left( e^{\nu_{eff}T_e(1 - \frac{t'}{T_n})} - 1 \right).$$  \hspace{1cm} (24)

### 3.6. Electron current analysis

#### 3.6.1. Separation of the electron and ion contribution

The measured current $I_{exp}$ is the sum of the electron and ion currents. For analyzing the electron current on the basis of the swarm model from section 3.4 the pure electron current $I_e$ is necessary. In this section, the method for extracting $I_e$ from $I_{exp}$ is described. For zero electron diffusion, all electrons arrive at the anode at the same time $T_e$. However, in case of diffusion some electrons arrive earlier and some delayed. Therefore, a time $T_3$ is defined, at which "nearly" all electrons arrived. We determine $T_e$ from the measured current and we set $T_3 = 2 \cdot T_e$. For typical gap distances and pressures in our experiments $T_e$ is two orders of magnitude larger than $\tau_D$, that is $T_e/\tau_D \approx 10^2$. In this case, the electron current at time $T_3$ drops to $I_e(T_3)/I_0 \propto (1 - \text{erf} \left( 0.5 \sqrt{T_e/\tau_D} \right)) \approx 10^{-12}$ of its initial current, according to equation (12). Thus, from this time point on, no more electrons are present and the measured current is only from ions $I_{exp}(T_3) = I_{ion}(T_3)$. There is no sensitivity of $I_{ion}(T_3)$ on the exact value of $T_3$, since $I_{ion}(t)$ is basically constant on the time scale of $T_e$. Using this considerations together with equation (19), the measured current can be written as

$$I_{exp}(t) = I_e(t) + \left( \frac{\nu_i}{v_e} + \frac{\nu_n}{v_e} \right) \int_0^t I_e(t') dt'.$$  \hspace{1cm} (25)

The factor in front of the integral can be obtained by the constraint $I_e(T_3) = 0$, and equation (25) becomes

$$I_{exp}(t) = I_e(t) + \frac{I_{exp}(T_3)}{\int_0^{T_3} I_e(t') dt'} \int_0^t I_e(t') dt'.$$  \hspace{1cm} (26)

This integral equation for $I_e$ is solved iteratively by starting in zero order with $I_e^{(0)}(t) = I_{exp}(t)$ and applying the operation

$$I_e^{(i)}(t) = I_{exp}(t) - \frac{I_{exp}(T_3)}{\int_0^{T_3} I_e^{(i-1)}(t') dt'} \int_0^t I_e^{(i-1)}(t') dt'.$$  \hspace{1cm} (27)
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3.6.2. Applying the electron swarm model to $I_e$ The electron current $I_e$ is evaluated on the basis of the electron swarm model from section 3.4 in order to obtain the electron drift velocity $v_e$, the characteristic time for longitudinal diffusion $\tau_D$ and the effective reaction rate $\nu_{\text{eff}}$. The rate $\nu_{\text{eff}}$ and the initial current $I_0$ are determined by means of a linear regression of $\log(I_e)$ versus $t$ in an interval $[T_1, T_2]$ where $I_e$ is unaffected by diffusion, as shown in figure 2 (a). The electron drift time $T_e$ and the characteristic time for longitudinal diffusion $\tau_D$ are determined from a fit of equation (12) to the falling edge of $I_e$ in the interval $[T_2, T_3]$.

3.7. Ion current analysis After the electron current analysis, the ion current is analyzed on the basis of the ion swarm model from section 3.5. In this section, we will write temporal parameters as vectors, since all currents will be treated as vectors for the fit routine. We seek a minimum number of fit parameters, in order to perform an efficient fit of the measured current $I_{\text{exp}}$ to the ion current model. Here, the fit minimizes the $L^2$-norm $||I_{\text{exp}} - I_p - I_n||$. At this point, the initial electron current $I_0$, the effective rate $\nu_{\text{eff}}$ and the time $T_e$ are already known from the electron current analysis, see section 3.6. For the ion current analysis they are treated as constant terms. After eliminating $\nu_a$ by using $\nu_{\text{eff}} = \nu_i - \nu_a$, three unknown parameters remain: $\nu_i$, $T_p$ and $T_n$. Notice that $I_p$ and $I_n$ depend linearly on $\nu_i$. We therefore rewrite the norm $||I_{\text{exp}} - I_p - I_n||$, using equations (23) and (24), to the form $||a - \nu_i b||$, with [24]

$$a = I + I_0 \frac{T_e}{T_n} (e^{\nu_{\text{eff}} T_e (1 - \frac{T_p}{T_n})} - 1).$$  \hspace{1cm} (28)$$

$$b = I_0 \frac{T_e}{\nu_{\text{eff}}} \left( e^{\nu_{\text{eff}} T_p} - e^{\nu_{\text{eff}} T_n} - \frac{e^{\nu_{\text{eff}} T_e (1 - \frac{T_p}{T_n})} - 1}{T_p} \right).$$  \hspace{1cm} (29)$$

Thus, only two independent fit parameters, $T_p$ and $T_n$, remain for the final fit, whereas the parameter $\nu_i$ that minimizes $||a - \nu_i b||$ is obtained by the linear inversion $\nu_i = a \cdot b / b^2$, with the constraint $\nu_i \geq \nu_{\text{eff}}$.

3.8. Derivation of the electron drift velocity and diffusion coefficient

The electron drift time $T_e$ and the characteristic time for longitudinal diffusion $\tau_D$ are in a first step obtained from fits to single current measurements, see section 3.6. One could derive the drift velocity simply by $v_e = d/T_e$. However, to increase precision, and to eliminate the possible offset in the initial swarm position or width due to the rather undefined non-equilibrium phase of the photo-electrons or experimental imprecision of the gap distance, the final values of $v_e$ and $\tau_D$ are extracted from several measurement at the same $E/N$-value but different gap distances $d$. Figure 12 (a) shows electron currents,
measured at gap distances $d = [d^{(1)}, d^{(2)}, d^{(3)}, d^{(4)}]$, which are fitted to derive the drift times $T_e = [T_e^{(1)}, T_e^{(2)}, T_e^{(3)}, T_e^{(4)}]$ as well as the characteristic time for longitudinal diffusions $\tau_D = [\tau_D^{(1)}, \tau_D^{(2)}, \tau_D^{(3)}, \tau_D^{(4)}]$. The values of $\tau_D$ should be equal but for the measurement uncertainty. The drift velocity is derived by linear regression of $d$ versus $T_e$, corresponding to the slope in Figure 12 (b). Similarly, we obtain the final value for $\tau_D$ from linear regression of $\tau_D T_e$ versus $T_e$, as shown in Figure 12 (c).

Figure 12: Sketch of how to obtain the electron drift velocity $v_e$ and the characteristic time for longitudinal diffusion $\tau_D$ for measurements taken at the same $E/N$ value. (a) Measured electron current $I_e$ vs time for 4 different gap distances $d^{(1)}, d^{(2)}, d^{(3)}, d^{(4)}$ resulting in 4 drift times $T_e^{(1)}, T_e^{(2)}, T_e^{(3)}, T_e^{(4)}$. The full line is the fit of equation (12) to $I_e$. (b) Gap distance vs drift time (circles) for the electron currents from (a) and linear regression (full line). (c) Characteristic time for longitudinal diffusion multiplied by drift time $\tau_D T_e$ vs drift time $T_e$ (circles) from (a) and linear regression (full line).
3.9. Detection and correction of a gap distance offset

The exact value of the reduced electric field $E/N$ as well as the derivation of $v_e$, as described in section 3.8, might be influenced by a possible inaccuracy of the measured gap distance $d$. The latter determines, together with the gap voltage $U$ and the particle density $N$, the value $E/N = U/dN$. In the following we describe how to detect and determine a small offset $d_0$ in the measured value of $d$. Accordingly, we correct the value of $E/N$ and the procedure of determining $v_e$ from section 3.8. There we assumed that the measurements are exactly at the same $E/N$-value, see figure 12. However, if there is an offset $d_0$, those measurements are not exactly at the same $E/N$-value, but at

$$\left( \frac{E}{N} \right)_c = \frac{E}{N} \left( 1 + \frac{d_0}{d} \right). \quad (30)$$

We denote $(E/N)_c$ as the corrected, or real $E/N$-value. It follows, that the measured values of the drift velocity are not exactly equal, but

$$v_e = (\mu N)(E/N)_c, \quad (31)$$

assuming that the reduced mobility $\mu N$ is constant over a small $E/N$-interval. The linear relation between $d$ versus $T_e$, assumed in the section 3.8, must be replaced by

$$d = d_0 + (\mu N) \frac{E}{N} \left( 1 + \frac{d_0}{d} \right) T_e. \quad (32)$$

This equation for $d_0$ can be solved iteratively by starting in zero order with $d_0 = 0$ and applying the linear regression

$$\Delta d^{(i+1)} = d - (\mu N)^{(i+1)} \frac{E}{N} \left( 1 + \frac{d_0^{(i)}}{d} \right) T_e. \quad (33)$$

where

$$d_0^{(i)} = \gamma \sum_{j=0}^{i} \Delta d^{(j)} \quad (34)$$

where $\gamma$ is a damping parameter comprised in $]0, 1[$. We continue the iterations until $d_0^{(i)}$ converges. The final values of the distance offset $d_0$ and of the reduced mobility $(\mu N)$ are obtained from the last iteration. Then, the final $(E/N)_c$ values are obtained with equation (30). The final value of $E/N$ that corresponds to the obtained reduced mobility $(\mu N)$ is taken as the mean of the final $(E/N)_c$ values.

References

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