Abstract

The evaporation coefficients of water in air and nitrogen were found as a function of temperature, by studying the evaporation of pure water droplet. The droplet was levitated in an electrodynamic trap placed in a climatic chamber maintaining atmospheric pressure. Droplet radius evolution and evaporation dynamics were studied with high precision by analyzing the angle-resolved light scattering Mie interference patterns. A model of quasi-stationary droplet evolution, accounting for the kinetic effects near the droplet surface was applied. In particular, the effect of thermal effusion (a short range analogue of thermal diffusion) was discussed and accounted for. The evaporation coefficient $\alpha$ in air and in nitrogen were found equal. $\alpha$ was found to decrease from $\sim 0.18$ to $\sim 0.13$ for the temperature range from 273.1 K to 293.1 K and follow the trend given by Arrhenius formula. The agreement with condensation coefficient values obtained with essentially different method by Li et al.\textsuperscript{1} was found excellent. The comparison of experimental conditions used in both methods revealed no dependence of evaporation/condensation coefficient upon
droplet charge nor ambient gas pressure within experimental parameters range. The average value of thermal accommodation coefficient over the same temperature range was found to be $1 \pm 0.05$.

**Keywords:** Mie scattering, evaporation model, Arrhenius formula.

## 1 Introduction

Many problems of science and technology are related to the evaporation from droplets and condensation on them. Cloud and aerosol microphysics together with construction of climate models, electrospaying, combustion, jet printing (compare) and spray painting (compare) are just some areas of relevance. Though they concern large sets of coexisting droplets, the understanding of transport processes at the surface of a single droplet is vital for solving them properly. Mass and heat transport processes at (nearly) flat interface can be efficiently modeled as a diffusion phenomenon. However, the evolution of small droplets is significantly influenced by effusion, which takes place in effectively collision-free region in the very vicinity of the interface (up to the mean free path of surrounding gas molecules). In order to account for this phenomenon, a so called evaporation (condensation) or mass accommodation coefficient $\alpha$ is introduced besides the diffusion coefficient. Likewise the thermal conductivity coefficient should be accompanied by a thermal accommodation coefficient $\alpha_T$. These coefficients describe transport properties of the liquid-gas interface. The mass accommodation coefficient can be perceived as the probability that a molecule (e.g. water) impinging on the interface from the gaseous phase side enters into the liquid phase and does not rebound. Analogically, the thermal accommodation coefficient determines the probability that a molecule impinging on the interface attains thermal equilibrium with the medium on the opposite side. The considerations of evaporation and condensation coefficients are considered equivalent and the values of these coefficients - equal. Both $\alpha$ and $\alpha_T$ coefficients are phenomenological and should describe only the properties of the very interface. All other processes influencing mass and heat transport, such as chemistry of the interface or the electrostatic interactions should be accounted for separately.

It is agreed, however, that $\alpha$ might possibly exhibit some temperature dependence. Many attempts have been made over nearly a century, to determine experimentally the values of $\alpha$ and $\alpha_T$ for water, but the results obtained by
different authors spanned from $\sim 0.001$ to 1 for $\alpha$ and from $\sim 0.5$ to 1 for $\alpha_T$ (see e.g.\textsuperscript{1,12–19} and\textsuperscript{9,11,20} for reviews). A variety of experimental methods was used. Both condensation on and evaporation from the surface of bulk liquid, liquid films, jets, and droplets were investigated in various environments (vacuum, standard air, passive or reactive atmospheres) under various pressures and for various water vapor saturations. Small droplets, such as encountered in clouds, have been favored, since kinetic effects manifest strongly for them. Suspended droplets, trains of droplets, clouds of droplets and single trapped droplets were studied.

We must admit that in our studies we have also experienced the flow of kinetic coefficients values in time. We have tried to overcome it. We will discuss the possible sources of the divergence of results in section \[5.1\].

The measurement of temperature dependence of $\alpha$ was rarely attempted, since the large divergence of obtained $\alpha$ values obscures the effect. Two recent studies by Li et al.\textsuperscript{1} and by Winkler et al.\textsuperscript{12} can serve as an example. The authors of the first study (Boston College/Aerody ne Research Inc. group) found that $\alpha$ decreases with temperature within the temperature range between 257 and 280 K. The authors of the second study (University of Vienna/University of Helsinki group) claim that $\alpha$ and $\alpha_T$ exhibit no temperature dependence between 250 and 290 K. The comparison of these results can be found in.\textsuperscript{20}

In this paper we present our new experimental results of evaporation coefficient of water in air, as well as our reprocessed results for water in nitrogen (compare\textsuperscript{21}), versus temperature; both under atmospheric pressure. The results for air and for nitrogen are consistent, which also indicates that the presence of small amounts of such soluble and/or reactant gases as CO$_2$ in the ambient air, does not influence the value of kinetic coefficients. In comparison to our previous works we refined our data processing which enabled us to determine the droplet radius with higher accuracy and trace its evolution with higher confidence. Smoother radius derivatives enabled us, in turn, to employ direct fitting of the model in finding the kinetic coefficients and avoid most approximations. We also operated on a larger set of experimental runs. This yielded correction of the values of kinetic coefficients we obtained previously, and revealed different temperature dependence. These results turned out to be in excellent agreement with the results of BC/ARI group: the values of $\alpha$ coincide within the temperature range from $\sim 273$ to $\sim 281$ K and our results extending towards higher temperature follow the temperature dependence found by BC/ARI group. Since BC/ARI group
and our results together span over larger temperature range, the accuracy of finding temperature dependence of \( \alpha \) could be improved.

## 2 Experiment

![Experimental setup - top view.](image)

The experimental setup is presented in figure 1. It consisted of a hyperboloidal electrodynamic quadrupole trap (see e.g. [22]), kept in a small climatic chamber. The high resistivity electric circuit of the trap drive enabled us to operate in a humid atmosphere. A detailed description of this apparatus can be found in [21,23] and of further modifications in [24,25].

The droplets were introduced into the trap with a piezo injector, similar to constructed e.g. by Lee et al. [26] or Zoltan. [27] The injection timing was...
controlled relative to the trap driving AC signal. By choosing the proper injection phase, the sign and, to a certain extent, also the value of the charge of the injected droplet could be controlled. The initial temperature of the droplet was that of the chamber.

In each experiment, the chamber was first flushed with clean, dry nitrogen, and then filled with a mixture of nitrogen/air and water vapor. The temperature in the chamber was monitored and stabilized. A zone type temperature control enabled us to eliminate vertical temperature gradients. Horizontal gradients were found negligible.

The humidity in the chamber, but outside of the trap was monitored continuously with semiconductor sensors. Due to poor vapor exchange through trap openings accompanied by injecting liquid water into the trap, the humidity inside the trap could not be inferred directly from those measurements. The value of humidity in the trap found as fitting parameter (see section 4) turned out to be higher by several percent than sensors readings. Resorting to fitting method was inevitable, since the humidity accuracy required for the correct assessment of kinetic coefficients was inaccessible via any on-line sensor measurement. On the contrary, analyzing droplet radius evolution seems a highly accurate method of measuring relative humidity, surpassing any on-line methods.

In our experiments we used ultra-pure water. The details about its initial parameters and sample preparation can be found in,21 where we discussed also the absorption of impurities by ultra-pure water, and their influence upon the experimental results there.

Droplet evolutions were studied with time resolved static light scattering, with green or red laser light. We found no inconsistency between the results obtained for both and we infer that the light wavelength had no influence upon the results.

3 Evaporation model

In order to interpret the experimental results, a model of evaporation was necessary. The model of evaporation we used was based on a generally accepted model which can be found in such textbooks as.9,28,29 It was a slightly rephrased and numerically reexamined version of what we had used previously.21,30 Below we discuss the details of the model equations we used, since the results may depend significantly on the apparently minute approxima-
tions made. We also point to a certain approximation typically made that we found weighing heavily upon the results.

The quasi-stationary evaporation of a free, motionless droplet larger than the mean free path of vapor molecules, can be easily described with the diffusion equations with boundary conditions defined by the thermodynamic conditions in the reservoir (far from the droplet). This part of the model does not rise much difficulties as long as the characteristic times of the process justify the quasi-stationary approach.\textsuperscript{31,32}

For small droplets, of the size comparable with the mean free path of vapor molecules, the language of diffusion is not appropriate. The transport of mass and heat below the mean free path distance from the surface must be perceived as effusion or evaporation into vacuum. The net effusive flow of vapor can be expressed as difference between outgoing and incoming effusive flows:\textsuperscript{9,33}

\[
J = \pi a^2 \alpha \left[ \rho_e(r = a) \overline{v}(T_a) - \rho(r = a + \Delta) \overline{v}(T_{a+\Delta}) \right],
\]  

(1)

where $\overline{v}(T) = \sqrt{8RT/\pi M}$ is the mean absolute thermal velocity of vapor molecules for the temperature $T$; $T_a$ is the temperature of the droplet (surface), $T_{a+\Delta}$ is the temperature of vapor at the distance $\Delta$ (comparable with the mean free path of the vapor molecule) from the surface. $\rho(r)$ is the vapor density at the distance $r$ from the droplet center while $\rho_e(r = a)$ is the vapor density at the droplet surface for the equilibrium conditions (steady state, no net flow).

The usual approximation made is $T_a = T_{a+\Delta}$ (see e.g.\textsuperscript{9}). It implies neglecting the slowing down of the mass transport by thermal effusion (a short range analogue of thermal diffusion). It should also be noted, that lifting the temperature dependence of $\overline{v}$ introduces some additional temperature dependence into $\alpha$. Unfortunately, discarding this usual approximation excludes using standard solutions and substantially complicates calculations. To overcome such difficulties, we decided to introduce a simple correction of $\alpha$ at the end. We shall address this issue in detail later. Following the standard route, we compare effusive and diffusive flows at $r = a + \Delta$. Since these flows are equal and both are proportional to the vapor density gradient it is possible to write a compact expression:

\[
da{da}{dt} = \frac{MD_k(a, T_a)}{R\rho_L} \left[ S_p \frac{p_c(T_R)}{T_R} - \frac{p_a(T_a)}{T_a} \right]
\]  

(2)
\[
= \frac{M D_k(a, T_a)}{R \rho_L} \frac{p_\infty(T_R)}{T_R} \left[ S - \frac{p_a(T_a)}{p_\infty(T_a)} \frac{p_\infty(T_a)}{T_a} \right],
\]

where

\[
\frac{p_a(T_a)}{p_\infty(T_a)} = \exp \left[ \frac{M}{RT_a \rho_L} \left( \frac{2 \gamma}{a} - \frac{Q^2}{32 \pi^2 \varepsilon_0 a^4} \right) \right] \tag{4}
\]
is the Kelvin equation, accounting for the modification of equilibrium vapor density near the droplet surface due to the surface curvature and charge effects,\textsuperscript{29} and

\[
\frac{p_\infty(T_a)}{p_\infty(T_R)} = \exp \left[ \frac{qM}{R} \left( \frac{1}{T_R} - \frac{1}{T_a} \right) \right] \tag{5}
\]
is the Clausius-Clapeyron equation. The effective diffusion coefficient \(D_k\) accounts for the effect of effusion:

\[
D_k = \frac{D}{a/(a + \Delta C) + D \sqrt{2 \pi M/(RT_a)/(a \alpha)}}. \tag{6}
\]

\(D\) is the diffusion constant for water vapor in nitrogen/air, \(T_R\) is the temperature of the reservoir, \(Q\) is the droplet charge, \(p_\infty\) and \(p_a\) are the equilibrium (saturated) vapor pressure above the flat interface and above the interface of the curvature radius \(a\) at a given temperature. \(S\) is relative humidity. \(\gamma\), \(\rho_L\), \(M\) and \(q\) are the surface tension, density, molecular mass and the latent heat of evaporation of liquid water, \(\varepsilon_0\) is the permittivity of vacuum, \(R\) is the universal gas constant. \(\Delta C\) defines the effective range of the gas kinetic effects. It is comparable with the mean free path of particles of surrounding gaseous medium \(\lambda_a\). We assumed \(\Delta C = 4\lambda_a/3.9\).

The change of droplet mass by evaporation (condensation) is associated with heat absorption (release), which manifests as temperature drop (rise) toward the droplet. The equation for the transport of heat can be presented in a convenient form:

\[
\frac{a}{a} \frac{da}{dt} = \frac{\lambda_K(a, T_a)}{q \rho_L} (T_a - T_R), \tag{7}
\]

where

\[
\lambda_K = \frac{\lambda}{a/(a + \Delta_T) + \lambda \sqrt{2 \pi M/(RT_a)/(a \alpha_T \rho_N c_P)}}, \tag{8}
\]

7
is the effective thermal conductivity of moist nitrogen (air) and $\lambda$, $\rho_N$, $M_N$ and $c_P$ are thermal conductivity, density, molecular mass and specific heat capacity under constant pressure of moist air/nitrogen respectively. $\Delta_T$ plays role analogous to $\Delta_C$ and was assumed as $\Delta_T = \Delta_C + 4\lambda/(\bar{v}c_P\rho_N)$. Since in the vicinity of standard temperature and pressure, the partial pressure of water vapor can be neglected in comparison to that of air/nitrogen, it can be assumed that the heat is conducted to the droplet mostly by the molecules of air/nitrogen. In consequence, the flux of mass can be considered independently of the flux of heat and $\Delta_C$ associated with the transport of mass should be distinguished from $\Delta_T$ associated with the transport of heat.

The direct influence of the droplet charge, through charge-dipole interaction, upon the mass (or heat) transport was estimated to be negligible for droplet charge ensuring Coulomb stability (compare$^{34}$). Similarly, field emission did not take place for surface charge densities encountered in our experiments (see e.g.$^{35}$). The Coulomb explosion of the droplet is a threshold process and does not need accounting in the transport equations.

4 Experimental data processing

The procedure of the numerical processing of experimental data, which we found the most stable and yielding the most consistent results, relies on the direct fit of the model equations to the experimentally obtained droplet radius change rate $\dot{a} \equiv da/dt$ as a function of droplet radius $a$. The data preparation procedure is presented below.

The running radius of the droplet $a_i(t_i)$ was obtained (off line) from the angularly resolved Mie scattering pattern for the time $t_i$ with the help of a gradientless fitting procedure ("library method"). Each droplet evolution yielded a sequence of a few hundreds data points indexed with $i$ (see figure 2). We had found that in order to obtain reliable results, significant care must be taken to ensure a high signal to noise ratio of the measurement. There happen data points misplaced to incorrect "evolution branch", associated with the Mie resonances that could not be handled with the method used (see description of the method$^{21}$). The accuracy of a single value of droplet radius $a_i$ (except for misplaced points) was estimated as $\pm$15 nm. The $a_i(t_i)$ sequence was stripped to main "evolution branch" (indicated with arrow in figure 2) and interpolated in order to obtain regularly spaced data points. The time derivative $\dot{a}_i(t_i)$ was calculated (figure 2). The $a_i(t_i)$ evolution was
Figure 2: An example of temporal droplet radius evolution, before and after processing - top and bottom curve respectively. Derivative calculated from processed data - middle curve. $N = 395$, $T_R = 283$ K, $p_{atm} = 1006$ hPa, $S_{sens} = 0.9$.

smoothed with low pass FFT filter and combined with the derivative in order to obtain $\dot{a}_i(a_i)$. Finally $\dot{a}_i(a_i)$ was smoothed (figure 3).

On the other hand, subtracting equation 2 from equation 7 leads to an equation binding $T_a$ and $a$. For every experimental $a_i$ this equation can be unambiguously numerically solved for $T_a$, yielding $T_a(a_i)$. This, on insertion into equation 2 yields at every experimental data point a numerically solvable equation binding $\dot{a}$ and $a_i$. Thus, a model prediction of $\dot{a}(a_i)$ could be obtained.

In order to find $\alpha$, $\alpha_T$, $S$ and $Q$, we minimized the function

$$
\chi^2 = \frac{X^2}{N} \sum_{i=1}^{N} [\dot{a}_i(t_i) - \dot{a}(a_i(t_i), \alpha, \alpha_T, S, Q)]^2
$$

(9)

using a gradient method. $N$ was the total number of experimental data points.
Figure 3: Droplet radius temporal derivative versus droplet radius, corresponding to figure 2 before and after filtering (points). The result of model fitting - solid line. Fitting parameters: $S_{fit} = 0.9762$, $Q = 3.7 \times 10^6$ elementary charge units, $\alpha = 0.155$, $\alpha_T = 1$.

of the evolution, and $\chi_0$ was an arbitrarily chosen normalizing factor. $\alpha$ and $S$ were found to be the essential parameters and could be unambiguously determined, while $\alpha_T$ and $Q$ could be determined only with limited confidence. Since $\alpha$ and $S$ had seemed partially interconnected, the minimization was performed very carefully, starting from various combinations of $\alpha$ and $S$ ($\alpha$ larger, $S$ smaller versus $\alpha$ smaller $S$ larger) and accepted only if leading to the same results. The less relevant parameters were initialized as follows: $\alpha_T = 1$, (values above 1 were allowed; compare$^{12}$), and $Q = 8\pi\sqrt{\varepsilon_0\gamma a_i^3}$, where $a_i$ corresponded to the smallest droplet radius observed in the evolution (no Coulomb instabilities during evolution). The resulting $Q$ was much approximate, and we couldn’t detect the eventual droplet charge loss (see eg.$^{36}$) by analyzing the evolution of the droplet radius. The minimization was also
hardly sensitive to $\alpha_T$, however a value close to unity could be inferred. Since for larger droplets ($a > 6 \ \mu m$) the kinetic effects as well as the effect of the droplet charge were negligible, only $S$ was fitted in this range, as a first step, and then the minimization was extended towards smaller radii with $\alpha$ added as a parameter. Finally $\alpha_T$ and $Q$ parameters were added. The whole procedure exhibited best stability for $S > 95\%$, since the evaporation was slower then (compare equation [2]) and thus: (i) the evolution of the droplet radius could be determined with high precision and (ii) the temperature jump at the interface $\Delta T$ was so small (compare equation [7]) that the model equations used were exact enough. It would be valuable to validate the procedure of finding kinetic coefficients using other liquids (such as ethylene glycol). Unfortunately, the parameters such as diffusion constant are usually not known with adequate precision. On the other hand, after slight modification of the procedure, it should be possible to look just for diffusion constant, which we intend to do soon.

4.1 Correction of $\alpha$

In order to estimate the influence of $\Delta T$ upon the obtained value of $\alpha$, we apply an approximation $T_{a+\Delta} = T_R$ to formula [1] which is opposite to usually applied $\Delta T = 0$, and we compare the results of both approximations. The approximation that we introduce means that we account only for thermal effusion while neglecting thermal diffusion. Since for our experimental conditions the temperature gradient was highest in the very vicinity of the interface (see [37]), our approximation was legitimate. For simplicity we also assumed that the shape of distribution of vapor density was spatially constant and temperature independent. It implied $\rho(r = a + \Delta) = S_{a+\Delta} \rho(T_R)$, where $S_{a+\Delta} = \text{const}$ represented relative humidity at $r = a + \Delta$. If we require that the effusive flows calculated with each of the approximations are equal, we have:

\[
\frac{\alpha}{\alpha(\Delta T = 0)} = \frac{S_{a+\Delta} - \frac{\rho_c(T_a)}{\rho_c(T_R)} \sqrt{\frac{T_R}{T_b}}}{S_{a+\Delta} - 1} \approx \frac{S_{a+\Delta} - \frac{\rho_c(T_a)}{\rho_c(T_R)}}{S_{a+\Delta} - 1} = \frac{S_{a+\Delta} - \rho_c(T_a)}{S_{a+\Delta} - 1}. \tag{10}
\]

Introducing $T_a(a_i)$ (see section [4]) into equation [10] we can find a correction of $\alpha$, where $S_{a+\Delta}$ is a (scaling) parameter. It is initiated as $S_{a+\Delta} = S$ and optimized so that $\alpha/\alpha(\Delta T = 0) \to 1$ for $\Delta T \to 0$ (larger $T_a$ in case of our experiment; see the inset in figure [4]). The results presented in figure [3] are
already corrected. In our case a significant (by a factor of nearly 2) correction was near the freezing point and by several percent at 276.5 K. The equation is essentially approximate and leads to underestimation of $\alpha$. It can be seen in figure - our data points seem to lie slightly below the trend line. It turns out that for many reasonable experimental conditions the correction factor can be higher than 2. We shall discuss a few examples in section 5.1. Considering the approximations made, we estimate that for thermodynamic conditions encountered in atmosphere the accuracy of the correction factor should not be worse than several percent.

5 Results and Discussion

The raw results are presented in figure 4 as a function of the droplet (surface) temperature. The kinetic coefficients should be presented as a function of the droplet (surface) temperature, since in general, due to evaporative cooling, it may differ significantly from the temperature of the reservoir. In case of BC/ARI group experiments, $T_R - T_a \leq 2$ K. Though, in our case $T_R - T_a \leq 0.7$ K only, it is sufficient that some of our results correspond to supercooled water as well.

The kinetic coefficients found for water droplets in nitrogen and in air were mutually compatible (see figure 4). It implies, that the gases absorbed by water from the air had negligible impact upon our measurements and generally there is no strong dependence upon the composition of the ambient atmosphere.

The final results are shown in figure 5. There are values of evaporation coefficient we obtained (solid circles) and values obtained by BC/ARI group, taken from (hollow circles). The values of thermal accommodation coefficient we obtained are also presented (solid squares). Data points corresponding to our results were obtained by averaging the raw results (compare figure 4). We also followed BC/ARI group and used the formula they derived basing on Transition State Theory (TST) (e.g. equation 7 in). Such formulation enables expressing the results in the language of thermodynamic potentials:

$$\frac{\alpha}{1 - \alpha} = exp (\Delta G_{obs})$$

(11)

where $\Delta G_{obs}$ is the Gibbs free energy and its temperature dependence can be expressed as $\Delta G_{obs} = \Delta H_{obs} - T \Delta S_{obs}$. $\Delta H_{obs}$ and $T \Delta S_{obs}$ are treated just as
Figure 4: Non-averaged experimentally obtained values of $\alpha$ as a function of droplet surface temperature. Solid and open circles represent results obtained for nitrogen and air respectively. The corresponding calculated evaporation coefficient correction factors, due to the thermal effusion, are presented in the inset.

parameters, their physical meaning is not clear (see discussion below). This formula is derived on an assumption, well justified with elegant experiments by Nathanson et al. described in\textsuperscript{38} that the particles from the gaseous phase enter liquid via an intermediate surface state. Dashed line in figure corresponds the fit we made to the results of BC/ARI group and our data points together. It yielded $\Delta H_{\text{obs}} = 4830 \pm 150$ cal/mol and $\Delta S_{\text{obs}} = 20.3 \pm 0.5$ cal/mol, which is within the limits of error equal to the values given in\textsuperscript{1}, i.e. $\Delta H_{\text{obs}} = 4.8 \pm 0.5$ kcal/mol and $\Delta S_{\text{obs}} = 20.3 \pm 1.8$ cal/mol. The accuracy of our fit (and so of the values obtained) is higher due to the larger number of data points.

The comparison of our results with those of BC/ARI group indicates
also that there was no perceivable influence of droplet charge upon kinetic coefficients. Vibrating orifice injector generates, at least in average, neutral droplets, while in our experiments with evaporating charged droplets it could be assumed that the charge was approaching its maximum value - the Rayleigh limit. Similarly, the comparison of aforementioned experiments reveals no measurable influence of ambient atmosphere pressure upon the value of kinetic coefficients.

The temperature dependence of $\alpha$, though obtained with essentially different method, coincides with the results of BC/ARI group (see eg. 20). Our result extends into higher temperature range. Furthermore, we measured
evaporation coefficient while BC/ARI group measured condensation coefficient. It supports the notion of equivalence of these coefficients.

The thermal accommodation coefficient we obtained $\alpha_T = 1 \pm 0.05$ (figure 5) agrees with both BC/ARI and UV/UH groups’ results. However it is hard to assess the real uncertainty of $\alpha_T$; the statistical error we found may be too small (see section 4). Thus it is not possible to derive information on its temperature dependence. Recently, there seems to arise a general consensus that $\alpha_T$ is close to 1, which means, that all the particles striking the interface thermalize.

5.1 An attempt of results coordination

Since it is quite improbable that all the kinetic coefficients measurements performed over the years were loaded with random errors, it must be assumed that the experiments, though accurate by themselves, measured different quantities. Many authors have tried to coordinate the results by pointing out what was really measured (see e.g.:9,11). However there is no consensus. We shall also try to address this issue.

The divergence of results obtained by different authors has been usually attributed to: (i) difficulties in accounting for various physical and chemical interfacial processes; (ii) effects of impurities, and especially surface active agents;39 (iii) structure of the interface (dynamic surface tension, reaching the balance by the interface) and (iv) dependence of the coefficient value upon the model used (indirectness of measurement). It has been pointed out9,11 that two classes of experiments could be distinguished: (i) with a quasi-static interface, yielding $\alpha < 0.1$ and (ii) with a continuously renewing surface, yielding $\alpha \geq 0.1$. However, such categorization requires defining the time scale. Such scale has not been agreed yet, neither the leading mechanism responsible for interface aging. For example, the characteristic times used in Molecular Dynamics (MD) studies are only hundreds of ps. This falls into a non-stationary interval, when the transients in the temperature and vapor density fields are starting to form. The Transition State Theory (TST) considerations of Nagayama et al.,40 seem to be in agreement with MD calculations and predict $\alpha \simeq 1$ around room temperature. However, it is worth noting, that, for example, stationary values of the surface tension are reached within milliseconds11 and all these time scales are far below the characteristic timescale of cloud droplet growth process, which lie in the range of seconds (or even minutes).41
Recently, Fukuta and Myers\textsuperscript{17} have noticed, that accounting for the effect of moving gas-liquid interface ("moving boundary effect") can change the resulting value of kinetic coefficients by several percent. In their work they managed to account for this effect in an elegant way. Though the thermodynamical conditions and the velocity of the interface in our experiment were similar to theirs, in present work, we have decided to neglect the moving boundary effect, since the correction of mass accommodation coefficient we propose is much larger.

In this paper we would like to point to a mechanism which falls within the 4th category - model dependent, however it is related to the issue of the characteristic timescale of the process and its distance from the thermodynamic equilibrium. Usually, authors are careful to estimate the characteristic times of mass and heat transport processes involved, in order to assure the proper description. It seems, that in some cases this alone can be somewhat misleading, because of the thermal effusion which we already mentioned. We shall consider four examples.

In case of BC/ARI group experiments, the vapor-liquid contact lasts several milliseconds but the droplet is essentially in equilibrium with the reservoir. In order to achieve temperatures below 273 K the evaporative cooling was used which inevitably caused temperature jump near the surface (up to 2 K) and thermal effusion as a consequence. However, since the value of $\alpha$ was not obtained from the evolution of droplet radius, its value should be safe and no correction is needed.

In our case, we selected for the analysis the droplet evolutions which lasted a few seconds which guaranteed that the process had been quasi-stationary in the diffusion time scale. For faster evolutions the temperature jump approached 1 K, and since $\alpha$ was obtained from the evolution, it had to be corrected by means presented above.

In case of the experiment of UV/UH group,\textsuperscript{12} the evolution lasted $\sim 50$ ms, which is shorter than in our case, but for the thermodynamic conditions they had, the process still could be regarded as quasi-stationary. However, the temperature jump of $\sim 3$ K could be expected for such evolution. This alone would require a correction of $\alpha$ by a factor of 2. Further overestimation might be caused by uncertainty of water vapor saturation. There are also rather few data points lying on a relatively flat curve, which as we know from our experience, causes the increase of measurement uncertainty.

Lastly, in case of very interesting Fukuta and Myers experiment\textsuperscript{17} the evolution (condensation) lasted $\sim 3$ s (similarly as in our experiment). Since
the final droplet radius was $\sim 2 \mu$m, it can be inferred that $\dot{a} \approx 1 \mu$m/s, which in turn yields temperature jump of only $\sim 0.2$ K. However, since the mass transport was relatively slow (supersaturation used was very small), the effect of even small temperature jump at the interface could be relatively large. According to our estimation (see expression 10) the correction of mass accommodation coefficient should be as high as 5! This would bring Fukuta and Myers result for NaCl and (NH$_4$)$_2$SO$_4$ at 277 K to $\alpha \approx 0.2$, which agrees within the limits of error with ours and BC/ARI group results, even allowing for moving boundary effect which we neglected.

6 Conclusions

We conclude that it is feasible to obtain reliable values of evaporation coefficient by analyzing the evaporation of a small droplet. It requires however several tens of data points per evolution and droplet radius measurement accuracy of several nanometers. Generally accepted model of quasi-stationary evaporation seems sufficient for experimental data analysis in most cases. We found however that when evaporative cooling of the droplet becomes of the order of 1 K, it is necessary to consider the effect of thermal effusion, which is a short distance analogue of thermal diffusion. The kinetic coefficients found for water droplets in nitrogen and in air were mutually compatible. The evaporation coefficient for the temperature range from 293.1 K down to 273.1 K was found to increase from $\sim 0.13$ to $\sim 0.18$ and follow the trend given by Arrhenius formula (see III) with the parameters $\Delta H_{obs} = 4830 \pm 150$ cal/mol and $\Delta S_{obs} = 20.3 \pm 0.5$ cal/mol. This temperature dependence is in excellent agreement with the results of BC/ARI group, which concern condensation coefficient, were obtained with essentially different technique for much lower ambient gas pressure and extend toward lower temperatures. The comparison with BC/ARI group experiments enables to draw a few additional conclusions: (i) the evaporation and condensation coefficients are essentially equivalent; (ii) there was no measurable influence of ambient atmosphere pressure upon the value of kinetic coefficients in the range from $\sim$ kPa to $\sim$ 100 kPa; (iii) there was no measurable influence of droplet charge upon the value of kinetic coefficients up to the Rayleigh stability limit. The value of thermal accommodation coefficient we obtained $\alpha_T = 1 \pm 0.05$ agrees well with recent results of many authors.
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