The transport of mass at the nano-scale during evaporation of droplets: the Hertz-Knudsen equation at the nano-scale

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Abstract

The applicability of Hertz-Knudsen equation to evolution of droplets at the nano-scale was investigated upon analysis of existing molecular dynamics (MD) simulations [Holyst et al., PRL 100; Yaguchi et al., J. Fluid. Sci. Tech. 5; Ishiyama et al., Phys. Fluids 16]. The equation was found satisfactory for radii larger than ∼4 nm. Concepts of Gibbs equimolecular dividing surface and the surface of tension were utilized in order to accommodate the surface phase density and temperature profiles, clearly manifesting at the nano-scale. The equimolecular dividing surface was identified as the surface of the droplet. A modification to the Tolman formula was proposed in order to describe surface tension for droplet radii smaller than ∼50 nm. We assumed that the evaporation coefficient for a system in and out of equilibrium may differ. We verified that this difference might be attributed to surface temperature change

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only. The empirical dependencies of the evaporation coefficient and the surface tension for flat interface, of liquid Ar in Ar gas at equilibrium, at the nano-scale, upon temperature was taken from existing MD data. Two parameterizations of the Hertz-Knudsen equation were proposed: (i) one using off-equilibrium condensation coefficient and effective density and (ii) another one using effective density and temperature at the interface. The second parametrization leads to an approximate solution of the Hertz-Knudsen equation requiring no free parameters. Such solution is suitable for experimental use at the nano-scale if only the temperature of the droplet (core) can be measured.

**Keywords:** nano-droplet evaporation/condensation, Hertz-Knudsen equation, condensation coefficient.

**Introduction**

The ubiquitous processes of evaporation and condensation arouse continued interest. A detailed modeling of these processes, especially concerning evaporation of droplets, is of interest, because of their role in the Earth ecosystem and in technology. Molecular dynamics (MD) simulations provide valuable data on these processes. However, the large scale, engineering applications of MD is still far from being feasible, while it seems that the continuous-medium descriptions of evaporation/condensation can provide many valuable information even at the sub-micro-scale (see eg. 2, 3).

In this work, we examine the extent of applicability of the (kinetic theory of gases) Hertz-Knudsen (HK) model to nano-scale evaporation. We used available in literature MD data to test our hypotheses.

The HK model considers free, ballistic motion of vapor particles near the droplet surface and introduces the permeability of the gas-liquid interface to these particles in a form of the accommodation (evaporation, condensation) coefficient as an empirical parameter. The HK model utilizes the concept of geometrical droplet radius and well localized (bulk) physical properties such as surface tension or density. It works well enough at the micro-scale though may not be sufficient...
to describe droplet evaporation/condensation at the nano-scale where sharp boundaries are lost. The surface region is discernible at the nano-scale (see figure 1) while it would be invisible at the micro-scale. If the HK model is to be adapted to the nano-scale, it is necessary to (re)define a few fundamental concepts: where the surface of the droplet is located, where the surface tension is exerted, which values from the density and temperature profiles should be used in transport equations, etc. Similarly, the applicability of the Antoine, Kelvin and Tolman equations at the nano-scale must be verified.

![Figure 1: Fluid density profiles for various stages of nano-droplet evaporation (MD simulation from\textsuperscript{6}) shown as solid dots and dashed lines. Middle stage (solid dots) accompanied by the corresponding temperature profile ((red) solid triangles and solid line).](image)

We used concepts of the Gibbs equimolecular dividing surface and the surface of tension (see eg.\textsuperscript{10}). They were introduced by Gibbs and developed independently by Tolman and Koenig, who derived approximate formulas linking the positions of these surfaces.\textsuperscript{11,12} The equimolecular (equimolar) dividing surface is a boundary in a hypothetical system of equal total number of molecules, at which the density changes discontinuously from liquid to vapor.\textsuperscript{10} The surface of tension is defined as the surface for which the Laplace equation holds exactly for all droplet radii. The equimolecular dividing surface was identified as the surface of the droplet and was used as a reference surface in model investigations. It is worth noticing (see figure 1) that the droplet tem-
perature is practically constant below the Gibbs surface. Other possible reference surfaces can be defined, however they don’t provide easy solution of the surface tension problem.

The analysis presented in this work concentrates on the transfer of mass. The full model describes also the energy transfer. The full description is necessary for the system consisting of two or more components, especially when the heat is primarily carried by non-evaporating component, e.g. in a case of evaporation of water droplets to moist air. In case of one-component system, equations of mass and energy transport are equivalent.

**MD simulation and HK model**

The MD simulation that we relate to, described in detail in, concerned evaporation of a nanodroplet into its own vapor. A monatomic, Lennard-Jones (LJ) fluid, corresponding to argon was used. The potential representing an LJ atom was truncated for radii larger than 0.875 nm. The LJ diameter was taken as 0.35 nm. Each simulation run started with a liquid droplet equilibrated with surrounding gas and located at the center of a sphere of radius $R_b$. Gas particles were reflected elastically from the boundary of the sphere. During the evaporation process, the center of the droplet was fixed at the center of the sphere. The parameters of the simulation were selected so that the simulated process was fully self-controlled and only the temperature at $R_b$ was fixed. The number of LJ atoms used in simulations ranged from $\sim 7.5 \times 10^5$ to $\sim 2.7 \times 10^6$ and the droplet temperature found in different simulations ranged from $\sim 95$ to $\sim 150$ K. The nano-droplet radius decreased, due to evaporation, from $\sim 12$ to $\sim 3$ nm. The studied system can be perceived as consisting of three phases of LJ fluid: droplet core, interface region (droplet surface layer) and surrounding vapor.

We start our considerations with a standard HK model and make use of the density and temperature profiles from the MD simulation. In general, mass transport from the evaporating droplet can be expressed as

$$\frac{dm}{dt} = -S(j_{out} - j_{in}),$$

(1)
where $m$ is the droplet mass, $j_{out}$ and $j_{in}$ are the outgoing and ingoing mass fluxes respectively and $S$ is the droplet surface area. The ratio between $j_{in}$ and the mass flux of molecules colliding with the surface $j_{col}$ defines the mass accommodation (condensation) coefficient $\alpha_c$ and reflects the permeability of the interface:

$$\alpha_c = \frac{j_{in}}{j_{col}}.$$  \hspace{1cm} (2)

In general, $\alpha_c$ may be expected to be a function of state of the surface phase.\textsuperscript{16–18} Combining (1) and (2) we obtain

$$\frac{dm}{dt} = -S(j_{out} - \alpha_c j_{col}) \cdot$$  \hspace{1cm} (3)

In the frame of kinetic theory of gases, $j_{col}$ corresponds to the thermal flux in the gas phase for the current state of the system:

$$j_{col} = \frac{\rho_v(T_v)u(T_v)}{4}, \text{ where } u = \sqrt{\frac{8k_BT_v}{\pi m_p}}$$  \hspace{1cm} (4)

is the average thermal velocity of molecules in the gas phase, $\rho_v$ and $T_v$ are the density and temperature of vapor respectively, $m_p$ is the mass of molecules (LJ atom in the MD simulation), and $k_B$ is the Boltzman constant. Similarly, classical approach at the micro-scale is based on the assumption that the outgoing flux is a function of temperature only. It is inferred\textsuperscript{4,5} that $j_{out}$ must be equal to the ingoing flux at equilibrium corresponding to $T_d$:

$$j_{out} = \alpha_{ce} j_{ine} = \alpha_{ce} \frac{\rho_{ve}(T_d)u(T_d)}{4}.$$  \hspace{1cm} (5)

The subscript ‘$e$’ indicates the equilibrium state and we abandon the usual assumption of $\alpha_{ce} = \alpha_c$ (eg.\textsuperscript{5,19}). It remains to be verified, whether these assumptions hold at the nano-scale. It is also worth noting that though the description is valid for both evaporation and condensation it uses only the condensation coefficients.

We notice (see figure 1) that the shape of the fluid density profile in the interfacial region hardly
changes during the evaporation. Thus the left-hand-side of equation 3 can be expressed as

\[ \rho_d \frac{dV_d}{dt} + \rho_s \frac{dV_s}{dt}, \quad (6) \]

where \( V_d \) and \( \rho_d \) are the volume of the droplet core and the bulk liquid density respectively, while \( V_s \) and \( \rho_s \) are the volume and the effective density of the interface region respectively. As it has been mentioned, it is convenient to adopt the Gibbs surface (sphere) radius as the droplet (surface) radius \( a \). We introduce the droplet core radius as \( a - \delta a \) (see figure 1), where \( \delta a \approx \text{const} \) defines the cut-off position in the fluid density profile. We chose it to obey the equality \((d\rho/dr)_{r=a-\delta a} = 0.01(d\rho/dr)_{r=a}\), where \( r \) is the distance from the droplet center and \( \rho \) is the corresponding fluid density. Thus, \( dV_d = 4\pi(a - \delta a)^2 da \), \( dV_s = 4\pi a^2 da \) and \( S = 4\pi a^2 \). Therefore, equation 3 can be rewritten in a form describing the evolution of the droplet radius:

\[ 4 \left[ \rho_d \left( \frac{a - \delta a}{a} \right)^2 + \rho_s \right] \dot{a} = \alpha_c(T_s)\rho_v(T_v)u(T_v) - \alpha_{ve}(T_d)\rho_ve(T_d)u(T_d). \quad (7) \]

It is natural to link \( \alpha_c \) to the droplet surface (Gibbs surface) and the temperature there, hence the notation: \( \alpha_c(T_s) \). However, as it has been already mentioned, \( T_s \approx T_d \). The values of \( \rho_d \) and \( T_d \) were taken as averages of the density and temperature profiles over the droplet core, and \( \rho_v \) and \( T_v \) as averages over the mean free path at the beginning of the vapor phase. The cut-off radius for the vapor phase \( R \) was chosen to obey the equality \((d\rho/dr)_{r=R} = 0.05(d\rho/dr)_{r=a}\). Averaging minimizes the error due to limited number of LJ atoms. Since the mass contained in the droplet core is proportional to \( a^3 \) while the mass contained in the transient (interface) region is proportional to \( a^2 \delta a \), the term corresponding to interface region can be omitted for the droplet larger than \( \sim 10 \) nm.

The value of \( \rho_{ve} \) must be estimated from the relation describing the equilibrium state of the system at the temperature \( T_d \). At the micro-scale and macro-scale a good estimation of \( \rho_{ve} \) is delivered by the set of three equations: the (extended) Antoine equation (see the Appendix), the Kelvin equation and the van der Waals (or even ideal gas) equation. It turns out that these equations can be applied at the nano-scale as well. In particular, the extended Antoine equation given in\(^{20} \)}
and the Kelvin equation were verified with the MD simulations\textsuperscript{9} for $a > 3$ nm at the temperature range from $\sim 85$ K to $\sim 105$ K.

In order to find $\rho_{ve}$ over the curved surface correctly, the Kelvin equation must be applied. At the nano-scale, a corrected value of the surface tension (as a function of droplet radius) must be used. This value is provided by Tolman formula, which is an approximate solution of the Tolman equation.\textsuperscript{11,12} The Tolman formula fails for droplet radii smaller than $\sim 50$ nm.\textsuperscript{8} However, using data from,\textsuperscript{8} we found that the Tolman formula can be heuristically modified to accommodate all radii greater than $\sim 3$ nm. We propose the following improved formula in the Koenig’s notation:\textsuperscript{12}

$$\frac{\gamma(a_{st})}{\gamma_{\infty}} = \left(1 - 2\frac{\delta_T}{a_{st}}\right)^w,$$

where $\gamma(a_{st})$ is the surface tension at the surface of tension, $\gamma_{\infty}$ is the surface tension of the flat interface, $\delta_T = a - a_{st}$ is the Tolman length, related to the Gibbs surface (droplet) radius $a$ and the radius of the surface of tension $a_{st}$, and $w$ is the added exponent. The empirical formula for the temperature dependence of $\gamma_{\infty}$ for the LJ liquid was found from\textsuperscript{6,8} (see the Appendix) for the temperature range from $\sim 85$ K to $\sim 135$ K. Formula 8 was very satisfactorily fitted to data published in,\textsuperscript{8} yielding $w = 1.6 \pm 0.2$ and $\delta_T = 0.41 \pm 0.06$ nm. So far we can attribute no physical meaning to $w$ parameter.

The value of the condensation coefficient at equilibrium $\alpha_{ce}(T_d)$ as the function of droplet temperature is known from independent MD simulations\textsuperscript{9} (see Appendix) for the temperature range from $\sim 70$ K to $\sim 130$ K.

Thus, for $a \gtrsim 10$ nm, $\alpha_c$ can remain the only free parameter of the model, while for smaller $a$, the effective interface density $\rho_s$ also has to be fitted. Since in our case $a \lesssim 10$ nm for the most part of the evolution, $\rho_s$ was always fitted.
Results and discussion

Two examples of a fit of equation 7 to molecular dynamics data are presented in figure 2 in $\dot{a}(a)$ form.

The density and temperature of fluid (used in equation 7) yielded by MD simulation for limited number of (LJ) atoms fluctuate significantly. In consequence, $\dot{a}(t)$ calculated from $a(t)$ fluctuates significantly as well. Thus, 9-points moving average was applied to $\dot{a}(a)$ and the difference between the raw and the averaged $\dot{a}$ served as an estimate of the uncertainty of $\dot{a}$ obtained from MD simulation. Then, the optimization can be performed unambiguously and the quality of fit is good for $a \gtrsim 4$ nm (compare also our results for larger droplets\textsuperscript{14,15}). It proves, that the continuous-medium HK approach can be adopted at the nano-scale. The detailed analysis showed that the approach is applicable as long as the surface layer (interface region) density profile is conserved (which is accompanied by $T_d \simeq \text{const}$).

Figure 2: Two examples of nano-droplet evaporation in $\dot{a}(a)$ form (derivative of the droplet radius with respect to time versus the radius); (black) solid dots represent MD data from,\textsuperscript{6} (red) triangles correspond to fits of equation 7.

We present $\alpha_c(T_d)$ and $\alpha_{ce}(T_d)$ in figure 3. $\alpha_c(T_d)$ obtained for different temperatures is self-consistent and consistent with $\alpha_{ce}(T_d)$. It is also generally consistent with previous results at the
micro-scale,\textsuperscript{14,21,22} though the Arrhenius relation is not followed. $\alpha_c(T_d)$ and $\alpha_{ce}(T_d)$ dependence on the liquid temperature $T_d$ are very similar, but $\alpha_c(T_d) < \alpha_{ce}(T_d)$ for all $T_d$. Such relation seems justified, since $\alpha_c$ is the condensation coefficient for an evaporating droplet. The second parameter of the fit, $\rho_s$, is presented in figure 4 versus $\rho_d$. All parameters of the fit are presented in table 1.

![Figure 3: Condensation coefficients, presented versus droplet temperature $T_d$: (black) open circles and (black) solid line - $\alpha_{ce}$ calculated with equation 5 and ((red) solid triangles) - $\alpha_c$ found as the free parameter of equation 7.](image)

Another parametrization of the fit is also possible. Now we assume that the functional dependence of $\alpha_{ce}$ and $\alpha_c$ upon temperature is the same. By introducing an effective temperature at interface $T_{s(\text{eff})}$ and substituting $\alpha_c = \alpha_{ce}(T_{s(\text{eff})})$ (using equation 11), $T_{s(\text{eff})}$ and $\rho_s$ become the free parameters of the fit. The new parameter $T_{s(\text{eff})}$ of the fit is presented in figure 5 as a function of the liquid temperature $T_d$.

For engineers applications the free parameters can be totally ruled out from the model, at least far from the critical point, by using linear and quadratic expressions binding them to the corresponding parameters of the droplet: $T_{s(\text{eff})} = 1.068(\pm 0.002)T_d$ and $\rho_s = 25(\pm 0.1) \times 10^{-5} \rho_d^2$ respectively. As far as in real system it would be possible to measure $a$, $T_d$ and $T_v$ all the parameters of the model can be calculated. However, the coefficients of the above relations must be found by
Table 1: Parameters $T_d$ and $\delta a$ found from MD simulations; $^6\alpha_{ce}(T_d)$ calculated with equation 11; $\alpha_c$, $T_s$ (equivalently) and $\rho_s$ found from the fit of equation 7 to the MD simulations. $^6$

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Figure 4: The effective droplet surface density $\rho_s$, found from fit, versus corresponding droplet core density $\rho_d$ (open circles). Solid line corresponds to quadratic fit (the point closest to the critical temperature was excluded).
Figure 5: The effective droplet surface temperature $T_s$ found from fit, presented versus droplet core temperature $T_d$. Solid line corresponds to linear regression.

analyzing the MD simulations.

Summary

We demonstrated that the HK equation can be successfully applied to nano-droplet evaporation for radii larger than $\sim 4$ nm. We adapted the variables and parameters in the equation to the nano-scale. The surface of the droplet was defined as the Gibbs equimolecular dividing surface and the surface of tension was distinguished. The choice of the Gibbs surface as the main reference surface was verified as beneficial. An empirical modification of the Tolman formula describing the dependence of the surface tension upon interface curvature for droplet radii smaller than $\sim 50$ nm was proposed to fit MD simulation. The modified formula holds for radii larger than $\sim 3$ nm. We assumed that the evaporation coefficient for system in and out of equilibrium may differ. The empirical dependency of the evaporation coefficient of Ar at equilibrium upon temperature in the range from 70 K to 130 K was found from MD simulation. Similarly, the dependence of the surface tension for flat interface of liquid Ar in Ar gas (at the nano-scale) upon temperature between 85 K
and 130 K was found from data published in references.\textsuperscript{6,8} The Antoine equation for saturation pressure of Ar vapor from\textsuperscript{20} was verified for temperature range from 85 K to 105 K.

Two parameterizations of the Hertz-Knudsen equation were proposed: (i) one using off-equilibrium condensation coefficient and effective density and (ii) another one using effective density and temperature at the interface. Using the first parametrization we reached the conclusion that the condensation coefficient is indeed a function of the droplet surface temperature only, which makes the condensation coefficient in and out of equilibrium seemingly different. This finding also led to the second parametrization. The second parametrization, in turn, opened way to an approximate solution of the Hertz-Knudsen equation requiring no free parameters. Such solution is suitable for experimental use at the nano-scale if only the temperature of the droplet (core) can be measured. However, such measurements cannot be done with our current apparatus, suitable only for micro-droplet experiments.\textsuperscript{14,15,22}

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### Appendix: Empirical formulas used in calculations

1. The extended Antoine equation for the saturated vapor pressure above the flat interface\textsuperscript{8,20} verified for the temperature range from $\sim 85$ K to $\sim 105$ K:

\[
\ln \frac{p_{\text{sat}}}{p_c} = (1 - |t|)^{-1}[a|t| + b|t|^{1.5} + c|t|^3 + d|t|^4] , \quad \text{where} \quad t = \frac{T - T_c}{T_c} . \quad (9)
\]

For Ar: $p_c = 48.7$ bar, $T_c = 150.8$ K and the Antoine coefficients $a = -5.90501$, $b = 1.12627$, $c = -0.76787$, $d = -1.62721$. 
2. The ‘empirical’ dependence of the surface tension for Argon for flat interface (at equilibrium) as a function of temperature, for the temperature range from $\sim 85$ K to $\sim 135$ K is as follows:

$$\gamma_{\infty}(T) = [298(\pm 4) - 2(\pm 0.04)T] \times 10^{-4}.$$  \hspace{1cm} (10)

3. The ‘empirical’ dependence of the (equilibrium) mass accommodation coefficient upon the droplet (surface) temperature, found from the data published in reference for the temperature range from $\sim 70$ K to $\sim 130$ K is as follows:

$$\alpha_{ce}(T) = 1.2(\pm 0.01) - 4(\pm 0.1) \times 10^{-5}T^2.$$  \hspace{1cm} (11)

References


Rate of radius evolution of nano-droplet

- MD simulation data
- Hertz - Knudsen equation

Radius change rate
Droplet radius

Rate of radius evolution
of nano-droplet