



1 Mapping the Performance of a Small Mixing-type 2 Condensation Particle Counter

3 Jitong Zhou^{1,2}, Gehang Huang^{1,2}, Fajun Yu^{3,4}, Xiaoqi Lei^{2,6}, Xiujuan Wang⁷, Huaqiao Gui^{3,4},
4 Huanqin Wang^{1,3,4,*}, Da-Ren Chen^{5,*}

5 ¹ University of Science and Technology of China, Hefei, China;

6 ² Institute of Intelligent Machines, Hefei Institutes of Physical Science, Chinese Academy of
7 Sciences, Hefei, China;

8 ³ Key Laboratory of Environmental Optics and Technology, Anhui Institute of Optics and Fine
9 Mechanics, Chinese Academy of Sciences, Hefei 230031, China

10 ⁴ Institute of Environment, Hefei Comprehensive National Science Center, Hefei 230031, China

11 ⁵ Particle Laboratory, Department of Mechanical and Nuclear Engineering, Virginia
12 Commonwealth University, 401 West Main Street, Richmond, VA 23220, USA

13 ⁶ School of Biological Resources and Environmental Sciences, Jishou University, Jishou, 416000,
14 China

15 ⁷ School of Microelectronics, Hefei University of Technology, Hefei 230009, China

16

17 *Correspondence to:* Da-Ren Chen (dchen3@vcu.edu), Huanqin Wang (hqwang@iim.ac.cn)

18



Abstract

The performance of a small mixing-type condensation particle counter (sMCPC) was numerically evaluated. The modeling calculated the fields of turbulent flow and temperature, and species transport in the particle channel of sMCPC, and the growth of particles included the effects of Kelvin, non-continuum and latent heat. Upon the validated, the model was applied to investigate the effects of temperature difference ($\Delta T = T_s - T_c$, where T_c and T_s are the temperature setting for working fluid saturation and sampled aerosol cooling, respectively), total flow rate (Q_g), and vapor fraction (f) on the working-fluid-governed supersaturation and particle activation in the sMCPC. It is found that the supersaturation ratio is increased, and the critical activation diameter ($D_{p,50}$) is lowered by increasing ΔT ; the excessive increase of Q_g reduces the supersaturation ratio and shifts the ratio peak towards the downstream of carrier flow; both the supersaturation ratio and the $D_{p,50}$ -slope are increased by increasing f . Under specific thermal and flow conditions, minimum activation diameters obtained in the cases with working fluids of ethylene glycol (EG), diethylene glycol (DEG), and dimethyl phthalate (DMP) is less than that in the case with *n*-butanol (B). Because of the particle growth after the activation, final sizes of particles exiting the particle growth tube are in micrometers in the case with *n*-butanol (B), and ~ 700 nm in case with EG; in contrast, final particle sizes in cases with DEG and DMP generally remain below the detection limit of typical optical particle counters (OPCs), i.e., ~ 0.3 μm .

Keywords: Condensation particle counter (CPC); Mixing-type condensation particle counter; CPC modeling; Particle activation; Particle growth.



1 Introduction

Condensation Particle Counters (CPCs) are widely used for measuring the number/concentration of aerosol particles, particularly for ultrafine particles defined as those with sizes less than 100 nm (below the lower detection limit of a typical optical particle counter, OPC). It is because of the "size amplification (via condensation) and optical counting" in CPCs, which significantly reduces the lower size detection limit of CPCs although at the expense of losing the original particle size (Heathman & Ensor, 2019). CPCs have been extensively applied in atmospheric pollution monitoring, environmental exposure assessment, roadside vehicle emission monitoring and many other applications in addition to scientific aerosol research (Takegawa et al., 2022; Pradhan et al., 2022; Li et al., 2024a).

Two types of CPCs capable of continuous counting (i.e., laminar flow and mixing types) are available. For laminar-flow CPCs with n-butanol as working fluid, aerosol particles first pass through a vapor-saturated chamber containing hot vapor and then transport to the condensation (or particle growth) channel at a temperature setting less than that of the vapor-saturated chamber (Wlasits et al., 2020). A stable temperature gradient established in the particle growth channel develops a supersaturated vapor environment, allowing particles to grow larger in sizes (ideally to optically detectable sizes) (Li et al., 2024b). Commercial CPCs, e.g., TSI CPC model 3025/3076, are capable of detecting particles in sizes as small as 3 nm (or even smaller with a particle amplifier, Iida et al., 2009). Furthermore, by increasing the temperature gradient or using a working fluid with a high saturated vapor pressure (e.g., diethylene glycol), the detection limit of laminar CPCs can be extended to approximately 2 nm (Vanhanen et al., 2011). Water-based CPCs, which exhibit performance comparable to butanol-based CPCs, have also been developed. Note that the temperature setting for establishing supersaturation conditions and growing the particle size in water-based CPCs is different from those of butanol-based CPCs (Hering et al., 2005; Mei et al., 2021; Hao et al., 2023).

MCPCs (Mixing-type CPCs or particle amplifier) amplify the particle size by rapidly mixing saturated vapor of the working fluid at high temperature with aerosol stream at cold temperature in a chamber (Fuchs, 1964). The mixing chamber can be small, e.g., a Swagelok cross used in the work of Wang et al. (2002). The above mixing creates a steep temperature gradient driving the working fluid vapor to the particle surface that activates the growth of particles in the sizes down to sub-5 nm and increases their sizes to optically detectable particle size limit (Mavliev, 2002; Kousaka et al., 1982). Vanhanen et al. (2011) introduced a mixing-type particle size magnifier that generates high supersaturation in a short channel by combining heated saturated vapor with a cold aerosol flow, achieving sub-3 nm detection. Wang et al. (2002) introduced a rapid-mixing condensation particle counting technique. Building on the above principle, Brechtel 9403a MCPC has been demonstrated to be UAV-deployable for *in-situ* aerosol monitoring (Bates et al., 2013). Overall, the MCPC development trajectory has been toward miniaturization while preserving its activation and reliable performance. Compared with laminar flow CPCs, MCPCs offer enhanced mixing and reduced particle residence time prior to their growth, which suppresses the particle loss (due to the Brownian diffusion) and counting errors induced by wall condensation (Sgro & de la Mora, 2003). The above features make MCPCs good candidates for applications requiring mobile particle monitoring.

Studies have employed COMSOL to calculate the flow and temperature fields, and species transport in laminar-flow CPCs. Kangasluoma et al. (2015) used COMSOL to quantify how the saturator–condenser



temperature difference shapes the supersaturation field in CPCs, affecting the overall CPC performance. Building on the above modeling framework, Barmounis et al. (2017) showed that reducing the operating temperature window can enhance the detection efficiency for sub-3 nm particles. Thomas et al. (2018) further employed COMSOL to examine how carrier-gas composition influences the particle activation in CPCs. Hao et al. (2021) modeled a laminar-flow CPC and identified an operational condition optimizing the activation efficiency of sub-3 nm ultrafine particles. However, the studies on the modeling of mixing-type CPC performance are limited.

In this modeling of MCPCs, COMSOL was applied to calculate both flow and temperature fields, and working fluid vapor concentration distribution in the aerosol flow channel of a small MCPC (Wang et al., 2025). The effect of operational parameters, including the temperature setting, flow velocity, flowrate ratio, and working fluid type, on the supersaturation establishment and particle activation efficiency were quantitatively evaluated. Under consideration of the effects of the Kelvin, non-continuum heat-transfer corrections, and latent heat, the condensational growth of particles in a small MCPC with different working fluids was computed in MATLAB. The result obtained in this modeling provides a solid foundation for the design and optimization of a small MCPC.

2 Material and Methods

2.1 Modeling of Mixing-type CPC

COMSOL was applied to calculate both the flow and temperature fields, and vapor transport in a small MCPC designed in the axisymmetric configuration. Shown in Fig. 1 is the schematic diagram of the model sMCPC. The governing equations used to calculate the flow and temperature fields, vapor distribution and particle growth in the small MCPC are described in this section.

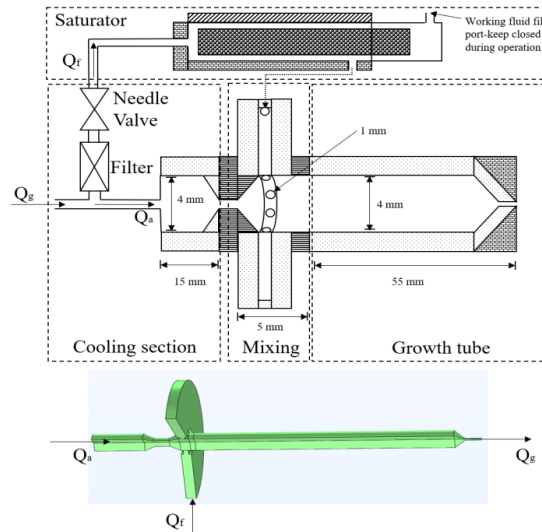


Figure 1. Schematic diagram of the model sMCPC and its 3D solid model for this study

In this study, the total inlet flow rate to the sMCPC is denoted as Q_g , which is separated into two streams: a portion of total inlet flow is used as vapor carrier flow, Q_h , (vapor carrier flow $Q_h = Q_g \times f$) which



carries hot vapor to the mixing chamber after passing through a HEPA filter and a working fluid reservoir; the other portion of total flow, $Q_a = Q_g \times (1 - f)$, which carries aerosol particles to the mixing chamber.

2.1.1 Problem Formulation

a) Governing Equations for Particle Growth

Both the supersaturation ratio and Kelvin diameter play crucial roles in the nucleation and size enlargement of particles. In a supersaturated vapor environment, vapor molecules condense onto the surface of particles, resulting in their size enlargement. Supersaturation ratio is the key factor governing the saturation rate (S_r), defined as the ratio of the partial pressure of the condensing vapor (P) to the saturated vapor pressure (P_s) at the flow temperature (T):

$$S_r = \frac{P}{P_s} \quad (1)$$

For the condensation growth of particles, the Kelvin diameter D_k defines as the smallest particle size that can be activated to start the growth of particles under a specific supersaturation ratio, S_r . Equation 2 is derived from the Kelvin equation, which relates the equilibrium vapor pressure over a curved droplet surface to that over a flat surface,

$$D_{p, kel} = \frac{4\sigma v_m}{k_b T \ln S_r} \quad (2)$$

where σ is the liquid's surface tension, v_m its molecular volume, k_b the Boltzmann constant, T the absolute temperature.

To quantify the particle activation of a CPC, the activation efficiency (η_{act}) was introduced as a key performance metric, defined as the ratio of particles which are successfully activated, and their sizes are grown to an optically detectable size, to the total number of particles entering the CPC. The activation efficiency can thus be calculated as:

$$\eta_{act} = \frac{\int_0^{R_{act}} 2\pi r w(r) N(r) dr}{Q_a N_0} \quad (3)$$

where r is the radial coordinate, $w(r)$ is the axial velocity profile, and $N(r)$ is the local number concentration of activated particles with the radius r . Q_a denotes the sampled aerosol flowrate; N_0 is the number concentration of sampled particles; and R_{act} represents the maximal radius of the activation zone in the growth tube.

While the supersaturation ratio is the key driver for condensation growth of particles in a CPC, excessively high supersaturation levels may increase the chance of new particle generation via the homogeneous nucleation of working fluid vapor. The formation of new particles can result in false counts in the CPC reading. It is therefore necessary to quantitatively assess whether the operational supersaturation leads to homogeneous nucleation or not through the evaluation of homogeneous nucleation rate (based on classical nucleation theory). By limiting the contribution of homogeneously nucleated particles to less than one particle per second, the maximal supersaturation avoiding the homogeneous nucleation can be determined. This provides a basis for determining the optimal CPC temperature in the modeling (Iida et al, 2009). The nucleation rate I can be calculated by (Friedlander, 2000)



$$I = 2 \times \left[\frac{p}{(2\pi m k_b T)^{\frac{1}{2}}} \right] \times \left(n v_m^{\frac{2}{3}} \right) \times \left[\frac{\sigma v_m^{\frac{2}{3}}}{k_b T} \right]^{\frac{1}{2}} \times \exp \left[-\frac{16\pi\sigma^3 v_m^2}{3(k_b T)^3 (\ln S)^2} \right] \quad (4)$$

where m is the molecular mass of working fluid and n is the molecular concentration of the condensing vapor. The total nucleation rate within the particle growth chamber is calculated by integrating I over the chamber volume.

Once particles are activated in a vapor-supersaturated environment, their subsequent condensational growth can be described by the diffusion-driven mass transport of vapor molecules to the particle surface. With the assumption of a spherical particle, and neglecting both the particle coalescence and curvature-dependent effects beyond the Kelvin correction, the temporal growth of particle diameter D_p can be estimated by:

$$\frac{dD_p}{dt} = \frac{4D_v v_m \chi (p - p_d)}{k_b T D_p} \quad (5)$$

where D_v is the diffusion coefficient of the condensing vapor in air, v_m is the molecular volume of the condensable vapor, χ is the Fuchs–Sutugin correction factor accounting for non-continuum effects, p is the surrounding vapor pressure, and p_d is the equilibrium vapor pressure at the droplet surface. The value of p_d is computed as a function of the droplet surface temperature T_d :

$$p_d = p_s(T_d) \exp \left(\frac{4\sigma v_m}{k_b T_d D_p} \right) \quad (6)$$

as proposed by Butt et al. (2013), incorporating the Kelvin effect for curved interfaces. The correction factor χ depends on the Knudsen number ($K_n = 2\lambda/D_p$) and is given by (Hegg & Larson, 1990):

$$\chi = \frac{1 + K_n}{1 + 1.71K_n + 1.33K_n^2} \quad (7)$$

To evaluate the thermal balance at the droplet surface, the temperature evolution of the droplet, T_d , is governed by a coupled energy conservation equation taking into consideration of the latent heat release and conductive heat-transfer between the droplet and surrounding gas:

$$\frac{dT_d}{dt} = \frac{3}{c_p \rho_v D_p} \left(H_{vap} \frac{dD_p}{dt} - 4k_g \frac{(T_d - T)\chi_h}{D_p} \right) \quad (8)$$

where c_p is the specific heat capacity of the droplet, ρ_v is the droplet density, H_{vap} is the latent heat of vaporization, and k_g is the thermal conductivity of the surrounding gas. χ_h corrects for the non-continuum effects in heat-transfer, calculated as

$$\chi_h = \left[1 + \frac{2k_g}{\alpha_T D_p \rho_g c_{p,g}} \left(\frac{2\pi m_g}{kT} \right)^{\frac{1}{2}} \right]^{-1} \quad (9)$$

where m_g , ρ_g , and $c_{p,g}$ are the molecular weight, density, and heat capacity of air, respectively. α_T is the thermal accommodation coefficient, and was assumed to be unity (Seinfeld and Pandis, 2016). Equation (8) captures the dynamic interplay between heat and mass transfer during the particle growth for accurately predicting the droplet evolution under transient supersaturation conditions.

The above-outlined framework provides the basis for calculating the particle activation and condensational growth under vapor-supersaturated conditions in a CPC. However, the local values of supersaturation and vapor concentration are highly dependent on the flow and temperature fields in the



187 mixing chamber of a MCPC. Therefore, an axisymmetric modeling was carried out by solving the
 188 governing equations for turbulent flow, heat transfer, and vapor transport in the model sMCPC.

189

190 **b) Governing Equations for Transport Phenomena in a MCPC**

191 This section describes the formulation used in the modeling of transport phenomena in MCPC. It includes
 192 the SST k - ω model for turbulent flow, energy equation for heat transfer, and dilute species transport
 193 equations for vapor distribution. All these equations are fully coupled in the modeling to calculate the
 194 local thermodynamic and fluid dynamics in the sMCPC.

195

196 The SST (Shear Stress Transport) turbulent flow model combines the advantages of both the k - ω model
 197 (providing improved accuracy near walls and in boundary layers) and the k - ε model (offering robustness
 198 in free-shear flows, i.e., better performance in fully developed turbulence regions) (Menter, 1993; Menter
 199 et al., 2003). By using a blending function to transition between two models based on distance from the
 200 wall, the SST model effectively captures the flow characteristics in both near-wall and core flow regions.
 201 This feature makes the flow model well-suited for calculating turbulent flows with strong shear stress
 202 and separation, such as that in the mixing chamber of a MCPC. Accurate prediction of velocity variation,
 203 shear layer, and recirculation zone is essential for the calculation of vapor transport and supersaturation
 204 distribution. The governing equations for mass and momentum conservation under the Reynolds-
 205 Averaged Navier–Stokes (RANS) framework are expressed as:

$$206 \quad \rho \nabla \cdot \mathbf{u} = 0 \quad (10)$$

$$207 \quad \rho(\mathbf{u} \cdot \nabla) \mathbf{u} = -\nabla p + \nabla \cdot [(\mu + \mu_t)(\nabla \mathbf{u} + \nabla \mathbf{u}^T)] + \mathbf{F} + \rho \mathbf{g} \quad (11)$$

208 where μ_t is the turbulent viscosity derived from the SST model, and $\rho \mathbf{g}$ represents the buoyancy force,
 209 which could be important due to the thermal gradients in a MCPC.

210

211 The transport of turbulence quantities in the SST model is governed by the following two additional
 212 partial differential equations: one for the turbulent kinetic energy k , and one for the specific dissipation
 213 rate ω :

$$214 \quad \rho(\mathbf{u} \cdot \nabla) k = \nabla \cdot [(\mu + \sigma_k \mu_t) \nabla k] + P_k - \beta^* \rho k \omega \quad (12)$$

$$215 \quad \rho(\mathbf{u} \cdot \nabla) \omega = \nabla \cdot [(\mu + \sigma_\omega \mu_t) \nabla \omega] + \frac{\gamma}{\mu_t} \rho P - \rho \beta_0 \omega^2 + 2(1 - F_1) \frac{\rho \sigma_{\omega 2}}{\omega} \nabla k \cdot \nabla \omega \quad (13)$$

216 where $P_k = \mu_t \mathbf{S}^2$ is the turbulence production term, $\mathbf{S}$ is the mean strain rate magnitude, $\mu_t = \frac{\rho k}{\omega}$ is the
 217 eddy viscosity, F_1 is the blending function between k - ω and k - ε models near the wall. σ_k , σ_ω , β^* , γ , and
 218 β_0 are empirical model constants.

219

220 The temperature distribution in a MCPC significantly influences the local saturation vapor pressure and
 221 thus the supersaturation field. The heat transfer in the flow is modeled using the steady-state energy
 222 conservation equation:

$$223 \quad \rho c_p (\nabla T) = \nabla \cdot (k_{\text{eff}} \nabla T) + Q_{vd} \quad (14)$$

224 where ρ is the fluid density, c_p is the specific heat capacity at constant pressure, T is the absolute
 225 temperature, $k_{\text{eff}} = k + \frac{c_p \mu_t}{Pr_t}$ is the effective thermal conductivity, incorporating both molecular and

226 turbulent heat conduction, $Q_{vd} = \tau : \nabla \mathbf{u} + q_{turb}$ accounts for viscous and turbulent dissipation. The
 227 inclusion of turbulent thermal diffusivity via the turbulent viscosity μ_t is crucial for capturing enhanced



convective heat transport in the mixing region.

The transport of the working fluid vapor is modeled using the dilute species transport equation, which governs the convection and diffusion of vapor molecules:

$$\mathbf{u} \cdot \nabla c = D_{\text{eff}} \cdot \nabla^2 c \quad (15)$$

where c is the vapor molar concentration, $D_{\text{eff}} = D + \frac{\mu_t}{\rho Sc_t}$ is the effective diffusivity, Sc_t is the

turbulent Schmidt number.

c) Boundary Conditions

Appropriate boundary conditions are required to solve the above governing equations for modeling the flow mixing, temperature distribution, and vapor concentration in the sMCPC. The computational domain for modeling the sMCPC is shown in **Fig. 2**. The no-slip and specific temperature conditions (i.e., either saturator temperature T_s , or cooling temperature T_c) were set for all the solid walls for the flow and temperature fields, respectively. The vapor concentration of working fluid was set to the saturated concentration c_s at the inlet and to c_g on the walls of the growth tube. From the operation viewpoint, the temperature setting for the working fluid vapor and the growth tube determines the achievable supersaturation ratio in the growth tube, which is one of the key driving parameters. The sMCPC performance with different working fluids were further compared to investigate their effects on hygroscopic particle growth. Detailed boundary conditions are summarized in **Tab. 1** for reference.

Tab. 1 - Boundary conditions used in the modeling of a sMCPC

Region/Surface	Condition Description	Parameter(s)	Value
Inlet	Fully developed flow; parabolic velocity profile	Q_g	(0.1~0.4) LPM
		f	0.8~0.95
All solid walls	No-slip condition	—	—
Saturator walls	Fixed high temperature (saturation zone)	T_s	(35~50) °C
Cooling and Growth tube walls	Fixed low temperature (cooling zone)	T_c	(5~15) °C
Vapor stream inlet	Saturation vapor concentration at T_s	c_s	$C_v^{\text{sat}}(T_s)$
Growth tube walls	Saturation vapor concentration at T_c	c_g	$C_v^{\text{sat}}(T_c)$
Outlet	Reference zero pressure, convective flux (as applicable)	—	—

2.1.2 Computational Domain and Its Meshing

Fig. 2 shows the used computational domain including the aerosol inlet, saturated vapor inlet, turbulent mixing zone, particle growth tube, and outlet. The overall length of the domain is approximately 75 mm and an inner diameter of 4 mm. The mixing and growth tubes are 4 and 55 mm in length, respectively.

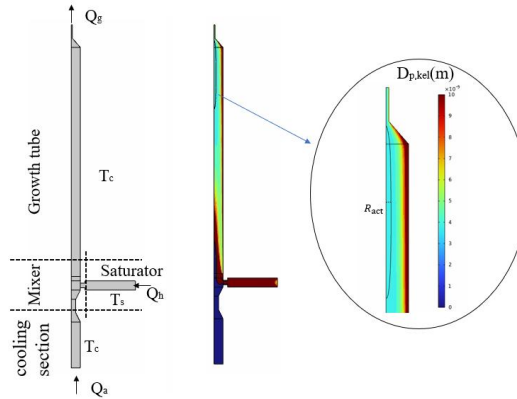


Figure 2. Computational domain for modeling the sMCPC performance and a typical distribution of the equilibrium Kelvin-diameter

Structured meshes with local refinement in the regions near the walls, in the mixing zone, and in the particle growth region were used. The thickness of the first layer meshes near solid walls was set to 0.002 mm to ensure the dimensionless wall distance $y^+ < 1$, (where $y^+ = \frac{\rho u_\tau y}{\mu}$, y is the distance from the wall, u_τ is friction velocity, ρ is density and μ is dynamic viscosity) in accordance with the requirement for using the SST $k-\omega$ turbulence flow model. In regions where saturated vapor and aerosol flows intersect, and within the mixing zone, the mesh size was refined to 0.03 mm. A total of approximately 24366 elements were used in the computational domain, with average element quality greater than 0.98 and a maximum mesh aspect ratio less than 10 (shown in **Fig. 3**).

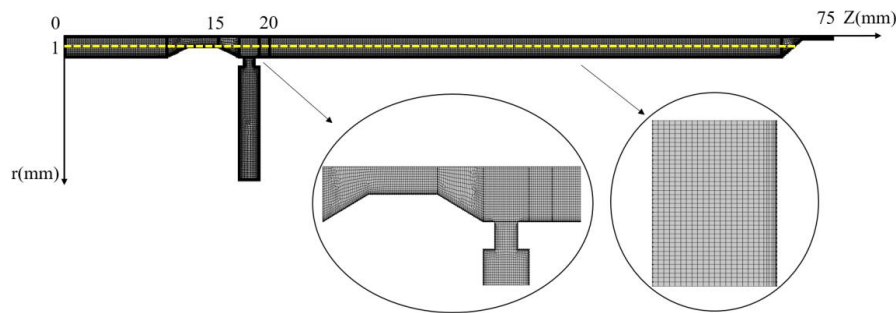


Figure 3. Computational domain for modeling the sMCPC performance and a typical mesh layout near walls and in the mixing/growth zones

To ensure the mesh independence of calculation results, a mesh sensitivity analysis was conducted by comparing the axial temperature profiles obtained using medium and fine meshes. As shown in **Fig. 4**, the temperature distributions along the centerline ($r = 0$) and near the wall ($r = 1$ mm) show excellent agreement between the two meshing schemes. The fine-meshed results closely overlap the medium-meshed results, with negligible deviation throughout the domain, including regions with steep thermal gradients in the region where two streams are first encountered. The above comparison confirms that the



medium meshing offers sufficient resolution to capture the key features of temperature. Medium meshing was therefore adopted for all subsequent modeling to save computational time.

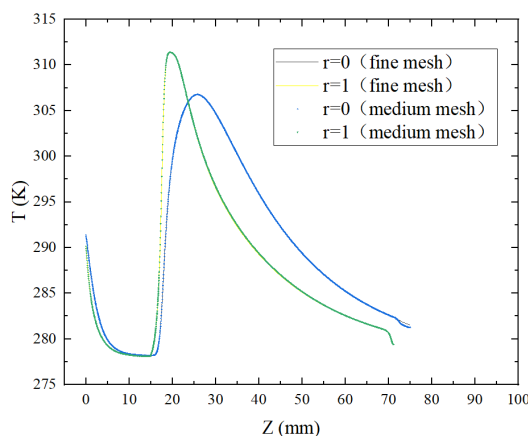


Figure 4. Comparison of axial temperature profiles along the centerline of the MCPC (calculated using medium and fine meshes)

2.1.3 Model Validation

To validate the modeling of the sMCPC, supplementary experiment to measure the temperature profile along the axis of the sMCPC was conducted. Under the operational conditions ($T_c = 10\text{ }^{\circ}\text{C}$, $T_s = 40\text{ }^{\circ}\text{C}$, $Q_g = 0.3\text{ LPM}$, $f = 0.9$), the axial temperature distribution along the centerline of the particle growth region was measured and compared with the calculated result. As shown in **Fig. 5**, reasonable agreement between the experimental and calculated temperature profiles along the MCPC centerline was observed ($R^2 = 0.927$). Overall, the modeling reproduces the temperature gradient and its spatial distribution. Minor discrepancies between two results could be attributed to the slight misalignment of measurement locations with the centerline when probing the thermocouples into the sMCPC.

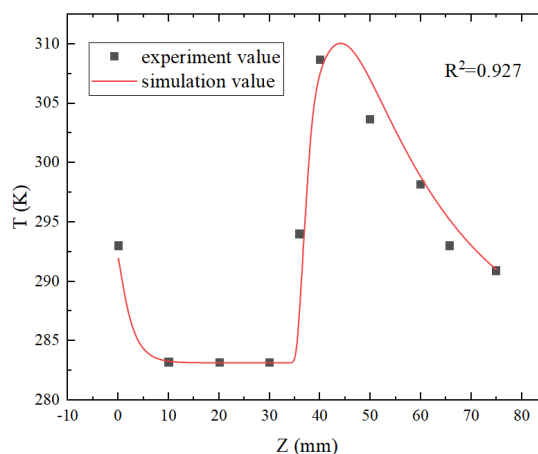


Figure 5. Comparison of measured and calculated axial temperature profile of the sMCPC



3 Results and Discussion

3.1 Effect of temperature setting

The differential temperature between T_c (for cooling section and growth tube) and T_s (vapor saturation chamber) is one of key factors influencing the performance of the sMCPC. The maximal supersaturation threshold to prevent working fluid vapor from homogeneous nucleation in the sMCPC is calculated using Eq. (4). Accordingly, the temperature setting for saturation vapor, T_s , is 35, 40, 45, and 50 °C; and the setting for both cooling section and growth tube, T_c , is 5, 10, and 15 °C, when using n-butanol as the working fluid. **Fig. 5** shows the temperature profile and supersaturation ratio along the axis of the sMCPC.

As evidenced in **Fig. 6**, significant variation was observed in both temperature (**Fig. 6a**) and supersaturation profile, S_r , (**Fig. 6b**) along the sMCPC axis. The increase of S_r was pronounced between $z = 20$ mm and $z = 60$ mm, peaked at $z \sim 65$ mm for all the studied temperature settings. Note that, although the setting with the largest temperature difference (i.e., $T_c = 15$ °C, $T_s = 50$ °C) resulted in the highest thermal peak, its maximal supersaturation was actually lower than that observed in the setting of $T_c = 5$ °C, $T_s = 40$ °C. It is attributed to the elevated vapor pressure results in the enhanced vapor loss at the high temperature setting. The calculated result indicates that $S_{r,max}$ reaches approximately 3.2 for the setting of $T_c = 5$ °C and $T_s = 40$ °C, compared to ~ 2.8 for the setting of $T_c = 15$ °C and $T_s = 50$ °C, suggesting that simultaneous increase in both T_s and T_c actually limits the achievable max. supersaturation in the sMCPC. Moreover, under minimal temperature differentials (e.g., $T_c = 5$ °C, $T_s = 35$ °C), the rate of increase in the supersaturation ratio was significantly low, and peaked at a relatively low value (compared with other settings), indicating weak driving force for the activation of ultrafine particles in sMCPC.

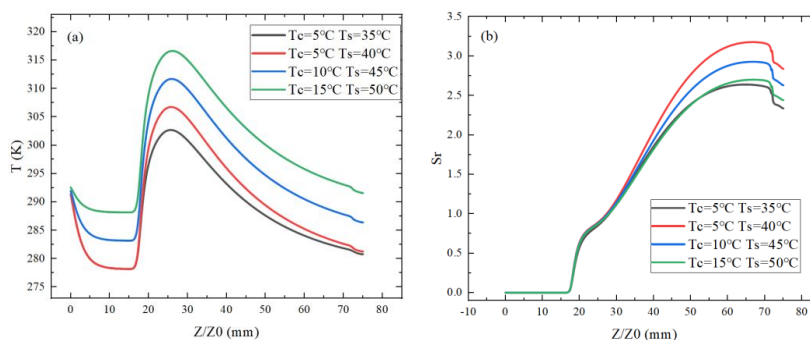


Figure 6. Temperature distribution (a) and Supersaturation ratio distribution (b) along the axis of the sMCPC under different temperature settings for T_c and T_s .

Fig. 7 shows the calculated activation efficiency for ultrafine particles in the sMCPC under different temperature settings. Under the condition of $T_s = 40$ °C and $T_c = 5$ °C, the particle diameter corresponding to 50% activation efficiency ($D_{p,50}$) is ~ 3.55 nm. In contrast, under the setting of $T_s = 35$ °C and $T_c = 5$ °C, this value increased to ~ 4.25 nm, indicating a notable difference in the activation performance between two temperature settings. The above observation further confirms that the temperature difference between T_s and T_c is a key parameter governing the strength and spatial distribution of the supersaturation in sMCPC, playing a dominant role in the particle growth activation. Not only does large



temperature difference facilitate the formation of high supersaturation ratio in the growth tube but also broaden the effective activation zone, thereby enhancing the CPC's sensitivity to sub-4 nm particles (Kuang et al., 2012; Barmounis et al., 2017). The condition of $T_s = 40^\circ\text{C}$ and $T_c = 5^\circ\text{C}$ was therefore adopted as the temperature setting in subsequent runs.

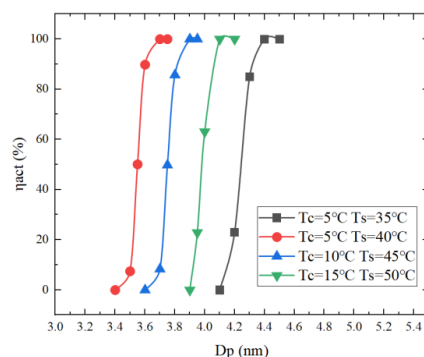


Figure 7. Effect of temperature settings (T_c and T_s) on the calculated activation efficiency and critical activation diameter of particles in sMCPC

3.2 Effect of total flowrate

Fig. 8 shows the effect of total flowrate, Q_g , on the temperature, saturation ratio, and activation efficiency of the sMCPC under a fixed vapor flow fraction of $f = 0.9$. As shown in the temperature profiles (**Fig. 8a**), the increase Q_g from 0.1 LPM to 0.4 LPM results in a noticeable decrease in the peak temperature and the peak location is moved downstream. This observation indicates that at a high flowrate, the residence time for mixing saturated vapor and cool air in the sMCPC is reduced, thereby diminishing heat-transfer efficiency.

The saturation distribution (**Fig. 8b**) exhibits similar trend as observed in the temperature distribution. At either a low total flowrate or high vapor fractions, sMCPC can rapidly establish a high supersaturation environment over a short axial distance. However, when Q_g increases from 0.1 to 0.3 LPM, the value of the maximal supersaturation ratio does not vary significantly. The notable change was observed for the downstream location shift of peaked supersaturation ratio in the growth tube. It is also noteworthy that, when the aerosol flow rate Q_a is very low, the diffusional loss of particles in the cooling section is significant (Balendra et al., 2023). In the consideration of both activation performance and diffusion losses, 0.3 LPM is thus selected as the operational total flow rate for the sMCPC.

Furthermore, the activation efficiency curves (**Fig. 8c**) reveal that the $D_{p,50}$ remains approximately 3.55 nm for Q_g between 0.1 and 0.3 LPM. When Q_g increases to 0.4 LPM, $D_{p,50}$ increased to above 3.60 nm, indicating that excessively high total flow rates hinder the effective activation of sub-4 nm particles.

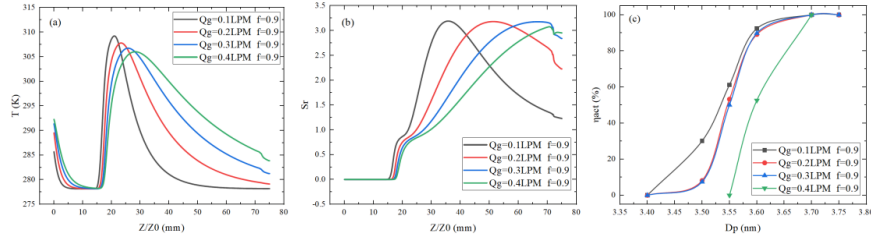


Figure 8. Effect of Q_g (at fixed $f = 0.9$) on (a) temperature, T ; (b) supersaturation ratio, S_r ; and (c) activation efficiency, η_{act} , in the sMCPC

Fig. 9 shows the effect of f (the fraction of total flowrate used as the vapor carrier flowrate) on the temperature, supersaturation ratio, and activation efficiency of the sMCPC, at a fixed total flowrate of $Q_g = 0.3$ LPM. From **Fig. 9a**, it is evident that the increase of f resulted in a significant elevation of the temperature peak in the mixing zone. It indicates that a high f value facilitates the formation of strong temperature difference over a short axial distance, which is conducive to enhance the supersaturation.

The corresponding supersaturation distribution (**Fig. 9b**) shows that at $f = 0.95$, a higher supersaturation region was established closer to the inlet, and the maximal S_r is greater than that under low f value condition. It indicates that not only does a high vapor fraction increase the vapor concentration but also enhance the mixing efficiency between hot vapor and cooled aerosol-laden air.

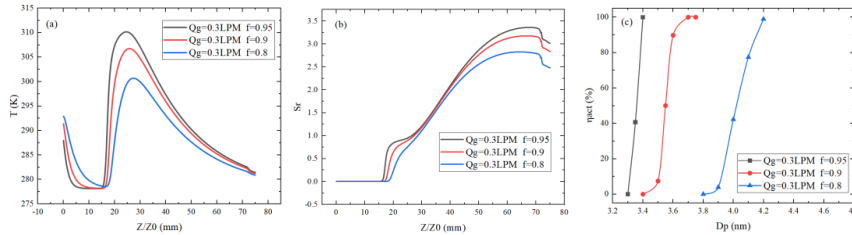


Figure 9. Effect of f ($Q_g = 0.3$ LPM) on (a) temperature, T ; (b) supersaturation ratio, S_r ; and (c) activation efficiency, η_{act} in the sMCPC

As shown in **Fig. 9c**, the effect of f on activation efficiency is dramatically pronounced. When the vapor carrier flowrate ratio, f , increases from 0.80 to 0.95 under a fixed total flow rate ($Q_g = 0.3$ LPM), the $D_{p,50}$ significantly decreased from ~ 4.2 nm to ~ 3.4 nm. This observation clearly demonstrates that a high vapor carrier flow fraction substantially enhances the sMCPC's sensitivity for ultrafine particles. Moreover, the steepness of the corresponding activation curves increased as f increased. The observed steep slope implies the establishment of a stable and well-defined supersaturated environment, enabling particles with the sizes above the critical size to be effectively activated without going through gradual transition. Not only does the increase of the vapor carrier flowrate boost the working fluid vapor concentration and enhance heat and mass transfer in the mixing zone but also leads to more consistent and sharply defined condensation growth behavior in the growth tube.

3.3 Effect of working fluid



We selected ethylene glycol (EG), diethylene glycol (DEG), dimethyl phthalate (DMP), and n-butanol (B) to investigate the combinational effect of vapor pressure and surface tension on the $D_{p,50}$ and droplet growth relative to the detection limit of optical particle counters (OPCs) (Hao et al. 2021). The calculated result of activation efficiency as a function of particle sizes is shown on **Fig. 10**.

Under the selected flow and temperature settings ($Q_g = 0.3\text{LPM}$, $f = 0.95$, $T_c = 5^\circ\text{C}$ and $T_s = 40^\circ\text{C}$), activation efficiency–diameter curves for four working fluids exhibit pronounced difference in their critical activation diameters (**Fig. 10**). EG exhibited the smallest critical diameter ($D_{p,50} \approx 2.5\text{--}2.6\text{ nm}$), followed by DEG and DMP ($D_{p,50} \approx 2.9\text{--}3.0\text{ nm}$), while B showed the largest diameter ($D_{p,50} \approx 3.3\text{--}3.4\text{ nm}$). The steep transition near $D_{p,50}$ in the cases with EG/DEG/DMP indicates sharp discrimination between non-activated and fully activated particles. Mechanistically, the minimal activation diameter is jointly determined by the curvature of the working fluid, governed by its surface tension and molecular volume, and the maximal supersaturation ratio sustained in the sMCPC. Working fluids capable of producing high and stable supersaturation level can effectively reduce the Kelvin diameter. Compared with B, both EG and DEG exhibited lower saturated vapor pressures and greater temperature sensitivity, making it easier to establish a stable and high supersaturation in the growth tube, thereby resulting in smaller critical activation diameters. However, the “activation” of particle growth does not always lead to the growth of particle size to an optically detectable size.

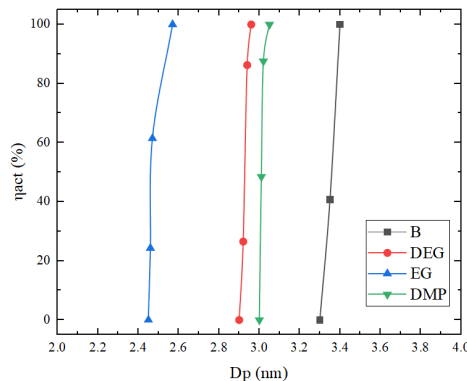


Figure 10. Calculated activation efficiency for the four working fluids (B, DEG, EG and DMP)

As shown on Fig. 11, the particle growth through the growth tube (exposure to high supersaturation environment) differed dramatically among four selected working fluids. The final particle sizes exiting the growth tube was in micrometers in the case with n-butanol (B); ~700 nm in the case with EG; 200–300 nm in the case with DEG. The final size was the smallest in the case of DMP. Therefore, in the sMCPC with limited growth tube length, both DMP and DEG would not be good candidates for working fluids due to insufficient droplet size growth. In contrast, the final particle sizes in cases with both EG and B are more likely to exceed the OPC detection limit although the activation diameter in the case with B is higher (compared with those in the other cases).

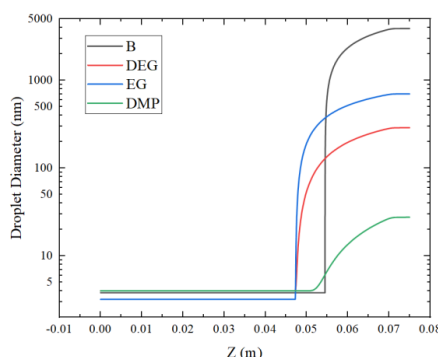


Figure 11. Growth of the particle size as a function of distance along the axis of sMCPC when using the four selected working fluids (B, DEG, EG and DMP)

4 Conclusion

An axisymmetric model was developed on the COMSOL platform to investigate the performance of a MCPC (mixing -type CPC) through the calculation of flow, temperature, and working fluid vapor concentration fields in the device. With the calculated fields, MATLAB was applied to obtain the activation efficiency and condensational growth of individual particles in the MCPC, in which the effects of Kelvin, non-continuum heat-transfer and mass-transfer, and latent-heat release were considered in the calculation of particle trajectories. Once validated by experimental data, the developed model enables us to numerically study the performance of a MCPC in general. In this study, we applied the developed model to study the performance of a specific sMCPC.

Two core performance metrics, i.e., $D_{p,50}$ (the activation size of particles for condensational growth) and the final size after the condensational growth as the functions of the temperature difference between the temperatures of cooling/growth tube and saturation vapor temperature, ΔT , total flowrate (Q_g), and the vapor carrier flow fraction (f), and working fluids were systematically evaluated. The result is summarized in the following:

- (i) **Temperature setting.** Moderate increase of ΔT with a low T_c enhances the supersaturation ratio in the sMCPC, resulting in the reduction of $D_{p,50}$. Under the condition of $T_c = 5^\circ\text{C}$ and $T_s = 40^\circ\text{C}$, both the peak supersaturation along the sMCPC axis and activation efficiency of the sMCPC are superior to those at the condition of $T_c = 5^\circ\text{C}$ and $T_s = 35^\circ\text{C}$, resulting in the decrease of $D_{p,50}$ from 4.25 nm to 3.55 nm.
- (ii) **Total Flowrate setting.** $D_{p,50}$ remains unchanged as Q_g varied from 0.1 to 0.3 LPM but obviously increases when Q_g increased to 0.4 L min⁻¹ (> 3.60 nm). Moreover, the location of the maximal supersaturation shifts downstream (toward the outlet), indicating that excessively high flowrate weakens the activation and reduces the effective particle growth path.
- (iii) **Vapor-carrier flow fraction setting.** The increase of f strengthens supersaturation and steepens the activation curve of the sMCPC. When f increased from 0.80 to 0.95 at $Q_g = 0.3$ LPM, $D_{p,50}$ decreased from 4.2 nm to 3.4 nm, resulting in the reduction of lower detection limit.
- (iv) **Working-fluid selection.** sMCPC with working fluids of EG, DEG, and DMP exhibited lower activation thresholds than that with B (with the smallest $D_{p,50} = 2.5\text{--}2.6$ nm for EG). However, the small activation particle size of the sMCPC did not always guarantee the final particle size for



optical detection. It is found that sMCPC with B as working fluid has the fastest droplet growth rate (with the final size in micrometers). sMCPC with EG grew the size of particles to ~ 700 nm. The final size in the cases with DEG/DMP failed to reach the OPC detection threshold (~ 0.3 μm), leading to the loss of particle detection efficiency because of the activated-but-undetected" events.

The above findings provide general guidance to specify the temperature settings and selection of, Q_g/f , and working fluids in the sMCPC design to achieve satisfactory performance for the detection of particle numbers in ambient aerosol monitoring.

Data availability

The sMCPC data in the study are available upon request to Huanqin Wang (hqwang@iim.ac.cn).

Author contributions

JZ, GH, and DC designed the research. JZ led the simulation and data analyses. JZ led the writing, with significant input from DC, XL and FY as well as further input from all other authors. HW, XW, and HG provided suggestions on the revision.

Competing interests

The authors declare that they have no conflict of interest.

Acknowledgements

This work was supported by National Key Research and Development Program of China under Grant 2025ZD1201200; National Natural Science Foundation of China under Grant U2133212; National Key Research and Development Program of China under Grant 2023YFC3705400, and the Key Research and Development Plan of Tianjin under Grant 22YFYSHZ00260.

Reference

- Barmounis, K., Ranjithkumar, A., Schmidt-Ott, A., Attoui, M., & Biskos, G. Enhancing the detection efficiency of condensation particle counters for sub-2 nm particles. *J. Aerosol Sci.*, 117, 44–53, 10.1016/j.jaerosci.2017.12.005, 2017.
- Balendra, S., Kale, A., Pongetti, J., Kazemimanesh, M., Haugen, M., Weller, L., & Boies, A. Condensation particle counters: Exploring the limits of miniaturisation. *J. Aerosol Sci.*, 175, 106266, <https://doi.org/10.1016/j.jaerosci.2023.106266>, 2023.
- Bates, T. S., Quinn, P. K., Johnson, J. E., Corless, A., Brechtel, F. J., Stalin, S. E., Meinig, C., & Burkhardt, J. F. Measurements of atmospheric aerosol vertical distributions above Svalbard, Norway, using unmanned aerial systems (UAS). *Atmos. Meas. Tech.*, 6(8), 2115–2120, <https://doi.org/10.5194/amt-6-2115-2013>, 2013.
- Butt, H.-J., Graf, K., & Kappl, M. *Physics and chemistry of interfaces*. John Wiley & Sons, 10.1002/3527602313, 2003.
- Friedlander, S. K. *Smoke, dust, and haze*. New York: Oxford University Press, 10.5860/choice.38-2208, 2000.
- Fuchs, N. A. *The Mechanics of Aerosols*. Pergamon Press, 10.1063/1.3047354, 1964.
- Hao, W., Stolzenburg, M., Attoui, M., Zhang, J., & Wang, Y. Optimizing the activation efficiency of sub-3 nm particles in a laminar flow condensation particle counter: Model simulation. *J. Aerosol Sci.*, 158,



- 105841, <https://doi.org/10.1016/j.jaerosci.2021.105841>, 2021.
- Hao, W., Mei, F., Hering, S., Spielman, S., Schmid, B., Tomlinson, J., & Wang, Y. Mapping the performance of a versatile water-based condensation particle counter (vWCPC) with numerical simulation and experimental study. *Atmos. Meas. Tech.*, 16(17), 3973–3986, <https://doi.org/10.5194/amt-16-3973-2023>, 2023.
- Hegg, D. A., & Larson, T. V. The effects of microphysical parameterization on model predictions of sulfate production in clouds. *Tellus, Ser. B, Chem. phys. meteorol.*, 42, 272–284, <https://doi.org/10.3402/tellusb.v42i3.15220>, 1990.
- Heathman, A. M. D., & Ensor, D. Monitoring of nanoscale particles in cleanrooms: ISO 14644-12. *Journal of the IEST*, 62(1), 50–59, 10.17764/1557-2196-62.1.50, 2019.
- Hering, S. V., Stolzenburg, M. R., Quant, F. R., Oberreit, D. R., & Keady, P. B. A Laminar-Flow, Water-Based condensation particle Counter (WCPC). *Aerosol Sci. Technol.*, 39(7), 659–672, <https://doi.org/10.1080/02786820500182123>, 2005.
- Iida, K., Stolzenburg, M. R., & McMurry, P. H. Effect of working fluid on sub-2 nm particle detection with a laminar flow ultrafine condensation particle counter. *Aerosol Sci. Technol.*, 43, 81–96, <https://doi.org/10.1080/02786820802488194>, 2009.
- Kangasluoma, J., Ahonen, L., Attoui, M., Vuollekoski, H., Kulmala, M., & Petäjä, T. Sub-3 nm Particle Detection with Commercial TSI 3772 and Airmodus A20 Fine Condensation Particle Counters. *Aerosol Sci. Technol.*, 49(8), 674–681, <https://doi.org/10.1080/02786826.2015.1058481>, 2015.
- Kuang, C., Chen, M., McMurry, P. H., & Wang, J. Modification of laminar flow ultrafine condensation particle counters for the enhanced detection of 1 nm condensation nuclei. *Aerosol Sci. Technol.*, 46, 309–315, <https://doi.org/10.1080/02786826.2011.626815>, 2012.
- Kousaka, Y., Niida, T., Okuyama, K., & Tanaka, H. Development of a mixing type condensation nucleus counter. *J. Aerosol Sci.*, 13(3), 231–240, [https://doi.org/10.1016/0021-8502\(82\)90064-7](https://doi.org/10.1016/0021-8502(82)90064-7), 1982.
- Li, Y., J. Wu, H. Wu, X. Liu, Q. Zhou, Y. Lu, Y. Wu, M. Liu, H. Ou, S. Mai, et al. A bipolar SMPS network for measuring atmospheric aerosols using natural air ions. *Atmos. Environ.* 325:120462, <https://doi.org/10.1016/j.atmosenv.2024.120462>, 2024a.
- Li, Y., Chen, X., Wu, J., Zhang, Q., Zhang, Z., Hao, J., & Jiang, J. A convertible condensation particle counter using alcohol or water as the working fluid. *Aerosol Sci. Technol.*, 59(2), 185–194, <https://doi.org/10.1080/02786826.2024.2395939>, 2024b.
- Mavliev, R. Turbulent mixing condensation nucleus counter. *Atmos. Res.*, 62(3–4), 303–314, [https://doi.org/10.1016/S0169-8095\(02\)00016-9](https://doi.org/10.1016/S0169-8095(02)00016-9), 2002.
- Mei, F., Spielman, S., Hering, S., Wang, J., Pekour, M. S., Lewis, G., Schmid, B., Tomlinson, J., and Havlicek, M. Simulation-aided characterization of a versatile water-based condensation particle counter for atmospheric airborne research, *Atmos. Meas. Tech.*, 14, 7329–7340, <https://doi.org/10.5194/amt-14-7329-2021>, 2021.
- Menter, F.R. Zonal two equation k-w turbulence models for aerodynamic flows. AIAA Pap, <https://doi.org/10.2514/6.1993-2906>, 1993.
- Menter, F.R.; Kuntz, M.; Langtry, R. Ten years of industrial experience with the SST turbulence model. In *Turbulence, Heat and Mass Transfer*; Hanjalic, K., Nagano, Y., Tummers, M., Eds.; Begell House Inc.: Danbury, CT, USA, Volume 4, pp. 625–632, 2003.
- Pradhan, B., Jayaratne, R., Thompson, H., Buonanno, G., Mazaheri, M., Nyarku, M., Lin, W., Pereira, M. L., Cyrys, J., Peters, A., & Morawska, L. Utility of outdoor central site monitoring in assessing



- 541 exposure of school children to ultrafine particles. *The Sci. Total Environ.*, 859, 160162,
 542 <https://doi.org/10.1016/j.scitotenv.2022.160162>, 2022.
- 543 Seinfeld, J. H., & Spyros, N. P. *Atmospheric chemistry and physics: From air pollution to climate change*.
 544 John Wiley & Sons, ISBN: 978-1-118-94740-1, 2016.
- 545 Sgro, L. A., & De La Mora, J. F. A Simple Turbulent Mixing CNC for Charged Particle Detection Down
 546 to 1.2 nm. *Aerosol Sci. Technol.*, 38(1), 1–11, <https://doi.org/10.1080/02786820490247560>, 2003.
- 547 Takegawa, N., Nagasaki, A., Fushimi, A., Fujitani, Y., Murashima, Y., & Sakurai, H. Volatility of aircraft
 548 exhaust ultrafine particles inferred from field measurements at Narita International Airport. *Atmos.*
 549 *Environ.*, 292, 119391, <https://doi.org/10.1016/j.atmosenv.2022.119391>, 2022.
- 550 Thomas, J. M., Chen, X., Maißer, A., & Hogan, C. J. Differential heat and mass transfer rate influences
 551 on the activation efficiency of laminar flow condensation particle counters. *Int. J. Heat Mass Tran.*, 127,
 552 740–750, <https://doi.org/10.1016/j.ijheatmasstransfer.2018.07.002>, 2018.
- 553 Vanhanen, J., Mikkilä, J., Lehtipalo, K., Sipilä, M., Manninen, H. E., Siivola, E., Petäjä, T., & Kulmala,
 554 M. Particle size magnifier for Nano-CN detection. *Aerosol Sci. Technol.*, 45(4), 533–542,
 555 <https://doi.org/10.1080/02786826.2010.547889>, 2011.
- 556 Wang, J., McNeill, V. F., Collins, D. R., & Flagan, R. C. Fast mixing condensation nucleus counter:
 557 application to rapid scanning differential mobility analyzer measurements. *Aerosol Sci. Technol.*, 36(6),
 558 678–689, <https://doi.org/10.1080/02786820290038366>, 2002.
- 559 Wang, H., Zhou, J., Pang, A., Lei, X., Huang, C., Chen, D. *A small-scale mixing-type condensation*
 560 *particle counter and its working method*. China Patent CN 119827382 B, Oct 3. Assignee: Hefei
 561 Institutes of Physical Science, Chinese Academy of Sciences. (Application No.: ZL 2025 1
 562 0077265.X), 2025.
- 563 Wlasits, P. J., D. Stolzenburg, C. Tauber, S. Brilke, S. H. Schmitt, P. M. Winkler, and D. Wimmer.
 564 Counting on chemistry: Laboratory evaluation of seed-material-dependent detection efficiencies of
 565 ultrafine condensation particle counters. *Atmos. Meas. Tech.* 13 (7):3787–98,
 566 <https://doi.org/10.5194/amt-13-3787-2020>, 2020.

568 **Table of Nomenclature**

- 569 c : Molecular concentration of the water vapor [mol m^{-3}]
 570 c_p : Heat capacity of the vapor [$\text{J K}^{-1} \text{kg}^{-1}$]
 571 $c_{p,g}$: Heat capacity of air [$\text{J K}^{-1} \text{kg}^{-1}$]
 572 c_s : Vapor concentration equal to saturation [$\text{J K}^{-1} \text{kg}^{-1}$]
 573 c_g : Fixed vapor concentration on surface [$\text{J K}^{-1} \text{kg}^{-1}$]
 574 D : Molecular diffusion coefficient [$\text{m}^2 \cdot \text{s}^{-1}$]
 575 D_k : The Kelvin diameter[m]
 576 $D_{p,50}$: Size of particle that has a 50% activation efficiency [m]
 577 $D_{p,kel}$: Size of particle that can be activated according to the Kelvin equation [m]
 578 D_v : Diffusivity of the vapor [$\text{m}^2 \text{s}^{-1}$]
 579 D_p : Particle size [m]
 580 D_{eff} : Effective diffusivity [$\text{m}^2 \cdot \text{s}^{-1}$]
 581 f : Vapor fraction [1]
 582 F : Generic body-force density [$\text{N} \cdot \text{m}^{-3}$]
 583 F_I : The blending function between k- ω and k- ϵ models near the wall
 584 g : Gravitational acceleration [$\text{m} \cdot \text{s}^{-2}$]



585	H_{vap} : Heat of vaporization of vapor [J kg^{-1}]
586	I : Nucleation rate [$\text{m}^3 \text{s}^{-1}$]
587	K_n : Knudsen number [1]
588	k : Turbulent kinetic energy [$\text{m}^2 \text{s}^{-2}$]
589	k_g : Thermal conductivity of air [$\text{W m}^{-1} \text{K}^{-1}$]
590	k_b : Boltzmann constant, 1.38×10^{-23} [J K^{-1}]
591	k_{eff} : Effective thermal conductivity including molecular and turbulent parts [$\text{W m}^{-1} \text{K}^{-1}$]
592	k_i : Molecular thermal conductivity [$\text{W m}^{-1} \text{K}^{-1}$].
593	N : Concentration of the particles [particles m^{-3}]
594	N_0 : Concentration of particles at the inlet of the conditioner [particles m^{-3}]
595	m : Molecular mass of working fluid [kg]
596	m_g : Molecular weight of air [kg mol^{-1}]
597	n : Molecular concentration of the vapor [molecules m^{-3}]
598	P : the partial pressure of the condensing vapor [Pa]
599	P_s : Saturation vapor pressure of the vapor [Pa]
600	P_k : Production of Turbulent kinetic energy [Pa s^{-1}]
601	p : Partial pressure of the vapor [Pa]
602	p_d : Equilibrium condensing vapor pressure at the surface of the droplet [Pa]
603	p_s : Saturation vapor pressure of the vapor [Pa]
604	Q_g : The total flow rate through the sMCPC [L min^{-1}]
605	Q_a : The aerosol flow rate [L min^{-1}]
606	Q_h : The vapor flow rate [L min^{-1}]
607	Q_{vd} : Volumetric heat source/sink [W m^{-3}]
608	r : Radial coordinate in the sMCPC [m]
609	R_{act} : Maximum radius of the contour corresponding to $D_{p,ket} = D_p$ [m]
610	S_r : Saturation ratio [1]
611	$\mathcal{S}$: Mean strain-rate magnitude [s^{-1}]
612	T_c : The temperature setting for working fluid saturation [K]
613	T_s : The temperature setting for sampled aerosol cooling [K]
614	T_d : Droplet surface temperature [K]
615	T : Flow temperature in the sMCPC [K]
616	ΔT : Temperature difference between working fluid saturation and sampled aerosol cooling [K]
617	$\mathbf{u}$: Mean velocity vector [m s^{-1}]
618	v_m : Molecular volume of the water vapor [m^3]
619	w : Velocity along the axial direction in the CPC [m s^{-1}]
620	σ : Surface tension of the vapor [N m^{-1}]
621	η_{act} : Activation efficiency [1]
622	χ : Correction factor for non-continuum effects in particle condensational growth [1]
623	χ_h : Correction factor for non-continuum effects in particle heat transfer [1]
624	α_T : Thermal accommodation coefficient of air [1]
625	ρ_v : Density of vapor [kg m^{-3}]
626	ρ_g : Density of air [kg m^{-3}]
627	λ : Mean free path [m]
628	ρ : The fluid density [kg m^{-3}]



629 μ : Molecular dynamic viscosity [Pa·s]
 630 μ_t : Turbulent viscosity from the SST model [Pa·s]
 631 ω : Specific dissipation rate [s⁻¹]
 632 σ_k , σ_ω , β^* , γ , and β_0 : Empirical model constants
 633 Pr_t : Turbulent Prandtl number [1]
 634 Sc_t : Turbulent Schmidt number [1]
 635 τ : Viscous stress tensor [Pa]