Evidence shows a strong correlation between human mortality/morbidity and atmospheric ultrafine carbon particle (UFCP with aerodynamic diameter <18 nm). Theoretical and experimental studies have attempted to use mass concentration/dose as exposure dosimetry to construct the dose-response relationships. Yet little attention has been given to the problem of using surface area dosimetry in UFCP-related risk assessment. We introduced an integrated risk assessment framework based on surface area dosimetry to estimate the adverse health potential risk exposed to atmospheric UFCP. We used the neutrophil cells elevation effect as adverse health effect endpoint. We reanalyzed the published data of UFCP particle diameter ($d_p$) and associated specific surface area (SSA) to reconstruct their relationship through log-linear regression method. Our results show that smaller particle size ($d_p < 51$ nm) demonstrated steep slope ($\ln SSA = 11.0 - 2.03 \ln d_p$), whereas larger particle size ($d_p > 51$ nm) was found close to the theoretical relationship ($\ln SSA = 8.65 - 1.20 \ln d_p$). We applied the modeled relationships to estimate the surface area doses of human inhaled particles in specific scenarios or subgroups. Our findings show that Adult and Youth subgroups in northern Taiwan region posed the highest potential risk, indicating that the median 10% exceedance risks are 39.6 (95%CI: 36.4–42.9) fold compared to control based on neutrophil cells elevation effect. The result provides a preliminary aspect for discussing the human health adverse effect exposed to atmospheric UFCP for specific groups based on particle surface area dosimetry.

1. Introduction

Epidemiological studies have revealed that ambient particulate matter (PM) concentrations and negative health impacts, such as asthma, hospital admissions, and premature mortality, are inextricably linked. The relationships between air pollutants and respiratory/cardiac diseases were found in several emergency room studies in Taiwan (Hwang et al., 2004), France (Nerriere et al., 2005), Canada (Yang et al., 2005), and USA (Wilson et al., 2005). Utell and Samet (1996) indicated that the potential mechanisms for induction of an inflammatory response attributed to: (i) ultrafine particle (UFP, i.e. particle with aerodynamic diameter less than 0.18 μm (Fine et al., 2004)), (ii) transition metal ions and (iii) aerosol acidity. Hence, a lot of UFPs play an important role for causing human adverse effects. Chow and Watson (2006) categorized the toxicological endpoints for human or animal exposing to UFPs into five classes of cell counts, enzymes, cytokines, bronchoalveolar lavage fluid (BALF), and DNA damage.

Generally, the carbon content is the most abundance element of particle in the ambient environment. Particulate matter contained nearly 50% (Li and Lin, 2003), 33–42% (Chio, 2005), and 17–25% (Lee et al., 2005) of carbonaceous materials measured in northern, central and southern
Taiwan, respectively. Therefore, PM contained carbon materials should be severe problem to human, especially for ultrafine carbon particle (UFCP) due to their relative large surface area compared to fine or coarse carbon particle (Kim et al., 2005; Stoeger et al., 2006). The large surface area of UFP or UFCP might play an important role causing more severe health effect (Oberdörster et al., 2005; Stoeger et al., 2006). In light of the above mentioned studies, surface area was selected as the key dosimetry in our study.

The carbon contents included organic matter, elemental carbon (EC) and inorganic carbon emitted largely from fossil fuel combustion, biomass burning, residential coal combustion, biogenic emissions, and secondary sources (Watson et al., 2006). Yet carbon black (CB) is a powdered form of EC that is manufactured by using hydrocarbons pyrolysis process. Over 90% CB production in the world is used for the reinforcement of rubber, and the other remaining is used for printing inks, paints, lacquer, and additives for industry (IARC, 1996; Tsai et al., 2001; Wellmann et al., 2006). The annual consumption rate of CB in Taiwan was nearly 113,000 tons yr\(^{-1}\). 90% of them were used domestically (Tsai et al., 2001). The International Agency for Research on Cancer (IARC, 1996) have addressed the CB as a group 2B (possible human carcinogen) based on the development of lung tumors in rats expose to CB chronically.

Evidence showed that the workers in CB manufacture plant, highway toll station, and dockyard posed potential human health risks (Morfeld et al., 2006; Wellmann et al., 2006). Mauderly et al. (2000) demonstrated the similar pulmonary carcinogenetics of CB and diesel soot in rats exposed heavily for two years. Soot is the residue from incomplete combustion of carbon-containing material, and varies widely in composition and physicochemical properties. These contained carbon particles would be released through tire wearing and oil burning processes emitted from vehicles (Mauderly et al., 2000), ten Brink et al. (2005) also indicated that the fraction of EC in the ultrafine level is approximately 40% by using an approach with filter and impactor sampling methods. Because the effects of ultrafine CB and UFCP inhaled could be similar, we intended to assess the UFP- and UFCP-induced adverse health risk from ambient environment using the biological effects exposed to CB.

The objectives of this study are threefold: (i) to assess the surface area-based cumulative dose of UFP exposure from atmospheric environment; (ii) to reconstruct and verify the suitable dose-response profiles to describe the relationship between UFP and human health effects; and (iii) to estimate the exceedance risk curves to susceptible subgroups or specific scenarios. Toward these goals, we reanalyzed several datasets measured throughout Taiwan regions and used an integrated method to estimate cumulative doses of different settings. Monte Carlo (MC) analysis and human respiratory tract (HRT)/multiple path particle dosimetry (MPPD) models (ICRP, 1994; CIIT Centers for Health Research, 2006) are introduced. HRT model is used for determining the probabilities and uncertainties of exposure doses from atmospheric environment. The MPPD model is used for assessing the internal dose of human lung. We also provided a framework to assess the exceedance risk profiles based on the toxicological endpoints from bioassays for protecting and early warning of the human health.

2. Materials and methods

We divided the probabilistic risk assessment (PRA) framework into four parts (Fig. 1) and will be described in detail in the subsequent sections.

2.1. Problem formulation

Studied designs included the northern, central, and southern Taiwan regions, denoting as NT, CT, and ST, respectively (Figs. 1A and 2). Three age groups (Adult, Youth, and Infant) and four activities (indoor (In), sleep (Sl), outdoor (Ou), and in-traffic (Tr)) are taken into account for exposure assessment. We divided the study area into three regions of Taiwan (NT, CT, and ST) because these three regions had different characteristics, especially for populations, vehicles, and industrial sources. Hence, we considered that assessing the risk in individual city or region was necessary. Based on the volume of air breathed rate, we also considered that these age groups (Adult, Youth, and Infant) are significantly different. On the other hand, different activities with associated time spent should be taken into account for modeling the real conditions.

The excellent available published PM\(_{0.18}\) data in Taiwan conducted from the previous study (Chio et al., 2007) was divided into several settings representing (i) different regions in Taiwan (NT, CT, and ST) and (ii) different activities (In, Sl, Ou, and Tr) within those regions (Fig. 2).

2.2. Exposure assessment

The precise estimation for particle surface area is crucial, because it affect the quantification of the surface area-based exposure dose significantly (Fig. 1B). We could obtain the surface area of particle through BET (Brunauer, Elmmett, and Teller) method (Brunauer et al., 1938). The particle diameter can be measured using transmission electron microscopy (Renwick et al., 2004; Stoeger et al., 2006). Here, we reconstructed the empirical relationship between particle size and corresponding surface area based on previous published data (Table 1). The specific surface area (SSA) of PM is defined as particle surface area (SA) divided by particle mass (m),

\[
\text{SSA} = \frac{\pi \eta d_p^2}{6 \rho \pi \eta d_p^3} = \frac{6}{\rho} (\eta d_p)^{-1} (d_p)^{-1},
\]

where \( \rho \) is the density of particle, \( \eta = 1/6m \) is the sharp parameter of particle diameter (d). A natural log linear function between SSA and dp can be rewritten, leading to

\[
\ln [\text{SSA}] = \ln \left[ \frac{6}{\rho} (\eta d_p)^{-1} \right] - \ln ([d_p]).
\]
\[ \ln Y = a + b \ln X, \]  
(3)

where \( Y \) is the size-dependent SSA of particle (dependent variable), \( X \) is the particle diameter (\( dp \)) (independent variable), \( a \) is the interception of the linear fitted model, and \( b \) is slope of the linear fitted model.

The upper and lower 95% confident limit level for the fitted linear model is also considered as,

\[ \hat{Y}_b \pm t(1 - \alpha/2; N - 2)s\{\hat{Y}_b\}, \]  
(4)

\[ s\{\hat{Y}_b\} = \text{SQRT} \left[ \text{MSE} \times \frac{1}{N} + \frac{(X_b - \bar{X})^2}{\sum(X - \bar{X})^2} \right], \]  
(5)

where \( \hat{Y}_b \) is the predicted mean value, \( t(1 - \alpha/2; N - 2) \) is a value of the \( t \)-distribution with \( N - 2 \) degrees of freedom, \( \alpha \) is the proposed significant level (here is equal to 0.05), \( N \) is sample sizes in regression model, \( s\{\hat{Y}_b\} \) means the standard deviation of the predicted mean value \( \hat{Y}_b \), SQRT means the square root operator, \( \text{MSE} = \sum(Y_i - \hat{Y}_i)^2/(N - 2) \) is the mean square error with \( \hat{Y}_i = a + bX_i \), and \( X_o, X, \) and \( X \) are the random, observed, and averaged independent-variables, respectively. \( Y_i \) and \( \hat{Y}_i \), however, are the random depend-variables responded to random variable \( X \) and \( Y \) variables are referred as particle diameter \( dp \) and specific surface area SSA in this study. Eqs. (4) and (5) are derived from Walpole et al. (1998). We applied the HRT model (ICRP, 1994) to estimate the internal exposure doses in different lung regions (Fig. 1B). The HRT model, varied with particle size ranges of PM\(_{0.18}\) and equilibrium time to each regional compartment and represented by a linear dynamic equation (Liao et al., 2006), was used to estimate surface area-based UFCP.

Fig. 1. Schematic diagram of the proposed probabilistic risk assessment framework in this study. The full names of all the abbreviated words in the figure are presented in the main content.
concentration in lung alveolar-interstitial (AI) region. We solved the linear dynamic equation to obtain the mass lung/indoor (L/I) ratio based on an equilibrium state in each compartment. The specific fractional deposition ($d_F$) for PM$_{0.18}$ in each compartment was then estimated, with a focus on the lung tissue of the lower respiratory tract (AI region). We also incorporated the MPPD model version 2 (CIIT Centers for Health Research, 2006) with HRT model (ICRP, 1994) to quantify the $d_F$ value.

We reconstructed an exposure model based on surface area dosimetry, involving the active surface area concentration of UFCP ($SA, \text{m}^2 \text{m}^{-3}$), volume of air breathed ($AB, \text{m}^3 \text{h}^{-1}$), time spent ($TS, \text{h d}^{-1}$), and exposure duration ($ED, d$),

$$D = SA \times AB \times TS \times ED,$$

with $SA = C_{\text{UFCP}} \times SSA \times d_F = PM \times f \times SSA \times d_F$, where $D$ is surface area-based cumulative dose of UFCP ($\text{m}^2$), $C_{\text{UFCP}}$ is the mass concentration of UFCP exposure and can be obtained from UFP concentration (PM) multiple by an apportion factor ($f$) of UFCP to UFP, SSA is the specific surface area of size-specific UFCP, $d_F$ is the deposition fraction deposited on AI region by HRT and MPPD models.

We treated AB and PM probabilistically and had lognormal distributions. The $f$ is assumed to be a solid value of 0.4 (ten Brink et al., 2005). The reliable values of factor $f$ in different settings are extremely difficult to obtain, if it is not impossible. Cass et al. (2000) and Ning et al. (2007) reported that the likelihood values of $f$ in urban area ranging from 0.44 ± 0.12 (mean ± sd) to 0.49 ± 0.13. Size-specific SSA can be obtained from our proposed particle surface area approach, TS is a scenario-specific constant value, ED is constant value of 365 days (i.e. 1-year duration for chronic exposure).

2.3. Effect assessment

Here we tested several target biomarkers from the literature by which limited data points can be constructed as surface area-based dose-response profiles. The neutrophil cells elevation effect (Renwick et al., 2004) is the available endpoints for assessing the precursor of adverse human health in the present study. Meanwhile, the
The dose-response profile with significance at its level of immune response while determination of neutrophil cells for a given surface area-based UFCP dose can be expressed mathematically as,

$$P(R, D) = P(D) \times P(R|D),$$

(9)

where $P(R|D)$ is the probabilistic risk for a certain cumulative dose $D$, $P(D)$ is the CDF of having surface area-based UFCP cumulative dose $D$ on human lung AI region, and $P(D)$ is the probability of exceeding the cumulative dose associated with the related response and can be expressed mathematically as,

$$P(R, D) = P(D) \times P(R|D),$$

(9)

where $P(R|D)$ is the probabilistic risk for a certain cumulative dose $D$, $P(D)$ is the CDF of having surface area-based UFCP cumulative dose $D$ on human lung AI region, and $P(D)$ is the probability of exceeding the cumulative dose associated with the related response and can be expressed mathematically as,

$$P(R, D) = P(D) \times P(R|D),$$

(9)

where $P(R|D)$ is the probabilistic risk for a certain cumulative dose $D$, $P(D)$ is the CDF of having surface area-based UFCP cumulative dose $D$ on human lung AI region, and $P(D)$ is the probability of exceeding the cumulative dose associated with the related response and can be expressed mathematically as,

$$P(R, D) = P(D) \times P(R|D),$$

(9)

where $P(R|D)$ is the probabilistic risk for a certain cumulative dose $D$, $P(D)$ is the CDF of having surface area-based UFCP cumulative dose $D$ on human lung AI region, and $P(D)$ is the probability of exceeding the cumulative dose associated with the related response and can be expressed mathematically as,

$$P(R, D) = P(D) \times P(R|D),$$

(9)

where $P(R|D)$ is the probabilistic risk for a certain cumulative dose $D$, $P(D)$ is the CDF of having surface area-based UFCP cumulative dose $D$ on human lung AI region, and $P(D)$ is the probability of exceeding the cumulative dose associated with the related response and can be expressed mathematically as,

$$P(R, D) = P(D) \times P(R|D),$$

(9)

where $P(R|D)$ is the probabilistic risk for a certain cumulative dose $D$, $P(D)$ is the CDF of having surface area-based UFCP cumulative dose $D$ on human lung AI region, and $P(D)$ is the probability of exceeding the cumulative dose associated with the related response and can be expressed mathematically as,

$$P(R, D) = P(D) \times P(R|D),$$

(9)

where $P(R|D)$ is the probabilistic risk for a certain cumulative dose $D$, $P(D)$ is the CDF of having surface area-based UFCP cumulative dose $D$ on human lung AI region, and $P(D)$ is the probability of exceeding the cumulative dose associated with the related response and can be expressed mathematically as,

$$P(R, D) = P(D) \times P(R|D),$$

(9)

where $P(R|D)$ is the probabilistic risk for a certain cumulative dose $D$, $P(D)$ is the CDF of having surface area-based UFCP cumulative dose $D$ on human lung AI region, and $P(D)$ is the probability of exceeding the cumulative dose associated with the related response and can be expressed mathematically as,

$$P(R, D) = P(D) \times P(R|D),$$

(9)

where $P(R|D)$ is the probabilistic risk for a certain cumulative dose $D$, $P(D)$ is the CDF of having surface area-based UFCP cumulative dose $D$ on human lung AI region, and $P(D)$ is the probability of exceeding the cumulative dose associated with the related response and can be expressed mathematically as,

$$P(R, D) = P(D) \times P(R|D),$$

(9)

where $P(R|D)$ is the probabilistic risk for a certain cumulative dose $D$, $P(D)$ is the CDF of having surface area-based UFCP cumulative dose $D$ on human lung AI region, and $P(D)$ is the probability of exceeding the cumulative dose associated with the related response and can be expressed mathematically as,

$$P(R, D) = P(D) \times P(R|D),$$

(9)

where $P(R|D)$ is the probabilistic risk for a certain cumulative dose $D$, $P(D)$ is the CDF of having surface area-based UFCP cumulative dose $D$ on human lung AI region, and $P(D)$ is the probability of exceeding the cumulative dose associated with the related response and can be expressed mathematically as,

$$P(R, D) = P(D) \times P(R|D),$$

(9)

where $P(R|D)$ is the probabilistic risk for a certain cumulative dose $D$, $P(D)$ is the CDF of having surface area-based UFCP cumulative dose $D$ on human lung AI region, and $P(D)$ is the probability of exceeding the cumulative dose associated with the related response and can be expressed mathematically as,
The AB values associated with different age groups and activities are shown in Table 2. In exposure assessment, the AB values and TS parameters in Table 2 are significantly affecting the surface area-based dose estimations. The assumed TS parameters for the Adult, Youth, and Infant subgroups in indoor activity are 13, 10, and 8 h, respectively. There are 8 h sleep activity for Adult and Youth subgroups that are also considered, whereas 14 h in sleep activity for Infant subgroup is reasonably assumed. For TS in indoor and in-traffic activities, the value for Youth (6 h) subgroup is higher than those of Adult (3 h) and Infant (2 h) subgroups.

These results take into account the three age groups (Adult, Youth, and Infant) with four major activities (indoor, sleep, outdoor, and in-traffic) on a daily basis. Fig. 4A presents the surface area-based UFCP dose for three age groups in NT, CT, and ST regions during one-year exposure based on subgroup-specific air breathed rates (Fig. 4B). Results indicate that the estimated doses for three age groups in NT region are higher than those in CT and ST regions. For NT region, there is highest median value of surface area-based UFCP dose for Adult (0.0914 m²), followed by Youth (0.0756 m²) and Infant (0.0159 m²), yet there is a wider range for Youth (95% CI: 0.0191–0.1486 m²). Accordingly, the Adult or Youth subgroup in NT region posed as more potential risk than those of other settings.

### Table 2

<table>
<thead>
<tr>
<th>Activities</th>
<th>Adult (A)</th>
<th>Youth (Y)</th>
<th>Infant (I)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Air breathed rate (m³ h⁻¹)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Indoor</td>
<td>1.07 (0.156)</td>
<td>0.753 (0.229)</td>
<td>0.31 (0.06)</td>
</tr>
<tr>
<td>Sleep</td>
<td>0.385 (0.092)</td>
<td>0.312 (0.073)</td>
<td>0.15 (0.03)</td>
</tr>
<tr>
<td>Outdoor</td>
<td>1.015 (0.148)</td>
<td>0.937 (0.483)</td>
<td>0.31 (0.06)</td>
</tr>
<tr>
<td>In-traffic</td>
<td>1.015 (0.148)</td>
<td>0.937 (0.483)</td>
<td>0.31 (0.06)</td>
</tr>
</tbody>
</table>

| Time spent (h d⁻¹) | | | |
| Indoor | 13 | 10 | 8 |
| Sleep | 8 | 8 | 14 |
| Outdoor | 1 | 4 | 1 |
| In-traffic | 2 | 2 | 1 |

| Data are from ICRP66 report (ICRP, 1994). |
| Data are reasonable assumed for people lived in Taiwan. |

The $d_p$ value for UFCP with 180 nm particle size deposited on AI regions was estimated to be 0.166 by HRT model (Fig. 4C). Results, however, also showed the entire distributions of $d_p$ values with polydisperse particles ($dp < 10 \mu m$) by MPPD model (estimated $dp = 0.124$, Fig. 4D). Taken together, we suggested that the $d_p$ value for UFCP deposited on the human lung AI region was accounted for 0.145 ± 0.021.

### 3.2. Dose-response curves

The maximum response $R_{\text{max}}$ for neutrophil cells elevation effect is estimated to be 47.4 fold compared to control (Fig. 5). The Hill coefficient ($n$) for the current fitting is 1.13 with a $R^2 = 0.91$. The specific cumulative dose yielding half of maximal response $K_{0.5}$ for neutrophil cells elevation effect was estimated at $2.96 \times 10^{-2} \text{ m}^2$.

### 3.3. Risk profiling

Table 3 gives that the exceedance risk (ER) matrix under 10 and 50% probabilities (ER10 and ER50) for surface area-based UFCP dose with selected endpoints onto each setting. Youth subgroup in NT region posed significant potential risk, whereas the lowest risk is appeared for Infant subgroup in ST region (Fig. 6A, B). Neutrophil cells elevation effect for Youth subgroup in NT region shows the highest potential risk, indicating the median ER10 values are 39.6 (36.4–42.9) fold compared to control.

There are highest values of median ER50 for Adult subgroup in NT region, showing 37.3 (34.7–39.9) fold compared to control for the neutrophil cells elevation effect. In contrary, for Infant subgroup in ST region, there are 15.5 (13.2–17.7) and 10.1 (7.7–12.4) fold compared to control based on the ER10 and ER50 for neutrophil cells elevation effect, respectively (Table 3).

### 3.4. Sensitivity analysis

Sensitivity analysis shows that the PM in in-traffic activity (PM/Tr), PM in indoor activity (PM/In), and $d_p$ value...
(DF) for Adult subgroup in three regions are the major factor (Fig. 7A–C; 89.4–92.5% of variance), whereas the PM in indoor activity (PM/In) is the affecting factor for Infant subgroup in Taiwan (Fig. 7G–I; 46.3–65.4% of variance). Results also show that the PM in in-traffic activity (PM/Tr) and the air breathed rate in in-traffic activity (AB/Tr) are the most two factors to influence the potential risks for Youth subgroup in Taiwan, except for northern Taiwan (Fig. 7D–F). Only one major factor (AB/Tr, 78.8% of variance) is found. Summarize the results of sensitivity analyses, we could find that PM/Tr, PM/In, DF, and AB/Tr variables were the significant factors affecting the risk estimation.

4. Discussion

4.1. SSA approach and dose estimates

Theoretically, the slope \( b \) in the SSA fitted model must be \(-1\) based on Eq. (1). However, our results show that \( b \) estimates ranged from \(-2.0\) (SSA model 2 with \( dp < 51 \text{ nm} \)) to \(-1.2\) (SSA model 3 with \( dp > 51 \text{ nm} \)), indicating that the original SSA data points measured by BET method might have high variability/uncertainty caused by shape and porosity. UFCP, soot, and diesel exhaust particle (DEP) adopted from Kim et al. (2005) and Stoeger et al. (2006) will also fall within the estimation range of SSA model 1. We successfully provided the evidence for carbonaceous materials (i.e. CB, CP, DEP, and soot) to verify the intrinsic SSA-\( dp \) relationships. However, our proposed SSA-\( dp \) relationships could not be applied for other materials or metals, even the regression interceptions and slopes

![Image](image_url)
might be varied enormously compared to other studies (Brown et al., 2007; Liao et al., 2007).

Our proposed SSA model also described well for UFCP, soot, and DEP for \( dp \) ranging from 9.3 to 23.6 nm. We suggested that SSA model 2 was better applied for \( dp \) ranging from 9 to 24 nm (RMSE = 19.5), whereas SSA model 3 was better used for \( dp \) ranging from 51 to 170 nm (RMSE = 2.41). Although the target size was greater than the highest diameter of 170 nm, the estimated RSE value was only 21.5% for \( dp = 180 \) nm. We thus suggested that the fitted SSA model 2 could be applied to estimate the SSA of target particle size of \( dp = 180 \) nm.

Meanwhile, the parameters of PM level (UFP concentration), specific surface area, deposition fraction, air breathed rate, and time spent in the proposed exposure model were considered as regional-, size- and scenario-dependent variables. Moreover, the apportion factor of UFCP to UFP and exposure duration were constants based on model assumptions (ICRP, 1994; ten Brink et al., 2005). Therefore, the Adult subgroup in NT region had the maximum median of UFCP dose based on the proposed settings, whereas Youth subgroup in NT region had the widest ranges of uncertainty associated with median value.

4.2. Quantification of risk and control strategy

Through the risk estimates of these preliminary health effects, researchers or decision makers could further understand the causing factor for tumor or other chronic diseases for human exposure to UFCP or emerging nanoparticles (NP). Nevertheless, this work had been studied based on several bioassays correlated to UFCP/NP. It was believed that the rising issues for human health would be noteworthy in the near future (Nel et al., 2006).

Either air breathed rate in in-traffic activity or PM level in in-traffic activity was the major factor affecting the cumulative dose of UFCP for all settings. Because the air breathed rate in in-traffic and outdoor actives had the same impact, the major factor affecting final result was the time spent for in-traffic activity. Hence, the recommended strategy for decreasing the potential risk was to reduce the time spent for in-traffic activity, implicating that human longer time spent for in-traffic activity had higher human health adverse effect. The related epidemiological study also proved this argument (Chen et al., 2005).

<table>
<thead>
<tr>
<th>Location</th>
<th>Adult (A)</th>
<th>Youth (Y)</th>
<th>Infant (I)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ER50</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NT</td>
<td>37.3 (34.7–39.9)</td>
<td>35.5 (33.1–37.8)</td>
<td>15.8 (13.5–18.0)</td>
</tr>
<tr>
<td>CT</td>
<td>29.7 (27.6–31.8)</td>
<td>26.5 (24.4–28.6)</td>
<td>10.5 (8.2–12.9)</td>
</tr>
<tr>
<td>ST</td>
<td>29.2 (27.1–31.1)</td>
<td>26.4 (24.2–28.5)</td>
<td>10.1 (7.7–12.4)</td>
</tr>
<tr>
<td>ER10</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NT</td>
<td>39.6 (36.4–42.9)</td>
<td>39.6 (36.4–42.9)</td>
<td>20.2 (18.1–22.4)</td>
</tr>
<tr>
<td>CT</td>
<td>35.2 (32.9–37.6)</td>
<td>33.8 (31.5–36.0)</td>
<td>16.0 (13.8–18.3)</td>
</tr>
<tr>
<td>ST</td>
<td>34.7 (32.4–37.0)</td>
<td>33.2 (31.0–35.4)</td>
<td>15.5 (13.2–17.7)</td>
</tr>
</tbody>
</table>

* The highest values are in bold face. 

Reducing the PM level in in-traffic activity was suggested to be the secondary strategy for reducing the human health risk. This strategy, however, was almost an expensive, complex and time-consuming task. Mobile sources emissions standard legislation and transportation management (McCarthy et al., 2006) methods were usually the issues to reduce the PM level in in-traffic activity. Moreover, bio-diesel or alcohol-gasoline additives uses were the novel method to reduce the PM level in in-traffic activity (Durbin et al., 2007). Previous studies revealed that there were 30–50% and 5–15% of PM reduction for pure biodiesel (B100) and the blend of 20% biodiesel with 80% fossil diesel (B20), respectively (Durbin et al., 2007).

Many researches reported the real-time carbon content (e.g. black carbon or elemental carbon) measurement for ambient air (Petzold et al., 1997; Kirchstetter et al., 2008) or diesel particulate matter (Fruin et al., 2004) by using Aethalometer. The principle of detecting carbon content was based on the changing optical absorption of light transmitted through accumulated particle on filter. Petzold et al. (1997) reported a simple model to describe the relationship between light scattering effect and carbon content. Kirchstetter et al. (2008) found an interesting weekly cycle for black carbon concentration in California implying that the residents lived in the area exposed to high human health on the weekday (Monday–Friday) periods than that on the weekend (Saturday and Sunday). Therefore, we might predict and present the potential health risk to human through combining our assessment model with
5. Conclusions

We introduced an approach to estimate the cumulative surface area-based dose deposited on human lung Al region of UFCP emitted from ambient environment. We employed HRT and MPPD models to successfully obtain the size-specific deposition faction for human lung regions. Surface area dose estimations of UFCP were obtained through the relationship between particle size and surface area. We successfully constructed the suitable dose-response profiles using the neutrophil cells elevation effect to describe the relationships between surface area-based UFCP dose and human health effects. Our results show that Adult and Youth subgroup in northern Taiwan region posed the highest potential risk, indicating that the median 10% exceedance risks are 39.6 (95%CI: 36.4–42.9) fold compared to control based on neutrophil cells elevation effect. Sensitivity analyses show that PM/Tr, PM/In, DF, and AB/Tr variables were the significant factors affecting the risk estimation. These results provide a preliminary assessment model to estimate the exceedance risk profiles for inhaling UFCP-induced human adverse effect for specific groups. Furthermore, we might link our proposed model and new real-time measurement technology for atmospheric UFCP to predict the potential human health risk in the near future.

References

Chio, C.P., 2005. Chemical Compositions and Source Contributions of PM2.5 and PM2.5–10 for Urban and Coastal Areas in Central Taiwan. PhD thesis, National Chung Hsing University, Taichung, Taiwan, ROC.
Chow, J.C., Watson, J.G., 2006. Ultrafine particles and health effects. Presented at Workshop on Recent Advances in Aerosol Measurement, National Central University, and Environmental Protect Administra-

Fig. 7. Sensitivity analysis results of the contribution to variance for each setting. The full names of all the abbreviated words in the figure are presented in the main content.


