Characteristics and Receptor Modeling of Atmospheric PM_{2.5} at Urban and Rural Sites in Pingtung, Taiwan

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Abstract

Suspended particles of PM_{2.5} in air were sampled concurrently at an urban site and a rural site in Pingtung County in southern Taiwan, in the spring, the summer and the fall of 2005. All samples were analyzed to identify eight water-soluble ions, carbonaceous contents, and 19 metal elements. Measurements reveal that the overall means of PM₁₀ (and PM_{2.5}) are 59.2 (47.4) μ g/m³ at Pingtung (urban) site, and 63.6 (45.7) μ g/m³ at Chao-Chou (rural) site. Although both sites exhibited strong correlations (*R* = 0.98 at Pingtung, and *R* = 0.78 at Chao-Chou) between PM₁₀ and PM_{2.5} masses, the mean PM_{2.5}/PM₁₀ ratio was 0.81 at Pingtung, higher than 0.68 at Chao-Chou, suggesting that relatively large bare lands and outdoor burning on farms may have caused more coarse particles to be present in PM_{2.5} at a rural site (Chao-Chou) than at an urban site (Pingtung). Results of CMB (chemical mass balance) modeling show that the main contributors to PM_{2.5} mass at Pingtung are vehicle exhaust (49.3–62.4%) and secondary aerosols (SO₄²⁻, NO₃⁻ and NH₄⁺) (31.2–37.8%), while those at Chao-Chou are the outdoor burning (25.3–50.4%) of agricultural waste, secondary aerosols (27.2–34.3%) and vehicle exhaust (12.0–26.9%), depending on the seasons.

Keywords: PM_{2.5}; Water-soluble ions; Carbonaceous species; Receptor modeling; CMB analysis.

INTRODUCTION

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Primary particulate matter (PM) is introduced directly into the atmosphere by



Fig. 1. Locations of measuring sites in Pingtung County.

anthropogenic sources (e.g., industry, vehicles and coal combustion) and natural sources (e.g., volcanic eruption, wildfire and marine aerosols). However, secondary aerosols have particles that are formed in the atmosphere by chemical reactions of gaseous components (gas-toparticle conversion) (Hinds, 1982). The most important secondary aerosols are sulfate (SO_4^{2-}) , nitrate (NO_3^{-}) and ammonium (NH_4^{+}) , formed mainly by photochemical reactions of the precursor gases, including sulfur dioxide (SO₂), nitrogen oxides (NO_x = NO + NO₂), ammonia (NH₃) or nitric acid (HNO₃). Some of the particulate organic carbon is formed from reactions that involve volatile organic compounds (Meng et al., 1997; Chow et al., 1998; Seinfeld and Pandis, 1998). Importantly, fine particles $PM_{2.5}$ (particles with an aerodynamic diameter of under 2.5 µm), have

attracted much interest since 1990s, because they can easily be inhaled and deposited in the respiratory organs of the human body and are thus very detrimental to human health (Needleman et al., 1990; Little, 1995: Oberdorster et al., 1995; Schwartz, 2000). The speciation and concentrations of PM_{2.5} are therefore critical in understanding their source contributors and developing effective methods of reducing their atmospheric levels. Currently, Taiwan's EPA (Environmental Protection Administration) has been establishing several PM_{2.5} monitoring networks around the country and expects to regulate PM_{2.5} in the near future.

Pingtung County (with a population of around 0.91 million and an area of around 2,775 km²) is located at the southern end of Taiwan (Fig. 1). Except in some densely populated areas in the northern parts of the

	Tomporatura	Wind speed	Deriod of	Ralativa humiditu		
Date	$\binom{^{0}C}{}$	(m/s)	sunshine ^a (h)	(%)		
	(C) (10.5) Substitute (1) (70) (a) Pinotung site					
3/08	20.6	<u>1.4</u>	5.7	52.1		
3/09	23.1	1.8	6.6	61.7		
4/29	28.2	1.6	10.2	79.4		
4/30	29.3	1.7	12.2	69.9		
5/01	29.6	1.9	11.8	68.5		
5/02	29.3	2.0	8.9	77.6		
6/21	29.1	1.8	7.9	75.2		
6/22	28.9	1.8	6.4	75.9		
7/12	31.6	2.2	11.2	66.0		
7/13	30.4	1.9	10.1	66.6		
8/09	31.2	1.8	7.4	67.9		
8/10	30.0	2.1	7.9	70.5		
9/13	30.0	1.6	3.5	72.3		
9/14	30.5	1.9	8.6	67.4		
10/12	28.6	1.8	7.5	75.4		
10/13	29.4	1.9	9.8	70.8		
		(b) Chao-Chou	site			
3/08	25.0	1.3	5.7	60.8		
3/09	25.4	1.6	6.6	68.4		
4/29	25.4	1.6	10.2	74.8		
4/30	25.4	1.7	12.2	66.8		
5/01	25.5	1.7	11.8	65.5		
5/02	25.5	1.8	8.9	72.8		
6/21	26.5	1.6	7.9	79.3		
6/22	26.3	1.5	6.4	80.5		
7/12	26.7	1.7	11.2	74.5		
7/13	27.4	1.5	10.1	74.8		
8/02	26.4	1.9	10.3	69.1		
8/03	26.9	2.3	9.9	67.9		
9/07	26.1	1.3	9.3	75.8		
9/08	26.0	1.5	7.6	73.4		
10/12	27.7	1.2	7.5	76.3		
10/13	27.9	14	98	73 9		

Table 1. Meteorological conditions at the two sites in Pingtung County in 2005.

^{*}Data were collected from the Hsiung-Kong station of Central Weather Bureau in Kaohsiung city.

county, with several small industrial parks, it is mainly agricultural, with touring and

sightseeing. However, the air quality in the northern (e.g., Pingtung city) and central parts

(e.g., Chao-Chou town) of Pingtung County is as bad as that in Kaohsiung metropolitan area, despite the fact that their population densities and emissions are much lower than those of the neighboring areas, mainly because the northern and central parts of Pingtung County are south or southwest of, and thus downwind of, the Kaohsiung City and Kaohsiung County, whenever a northerly or north-easterly wind prevails, including in autumn and winter (Chen *et al.*, 2003; 2004). However, the concentrations and compositions of PM_{2.5} in Pingtung County have not been measured, which are essential for estimating possible source contributions and/or assessing potential health risks.

This work presents measurements of concentrations and constituents in $PM_{2.5}$, including water-soluble ions, carbonaceous contents and metal elements, at an urban site and an agricultural town in Pingtung County, made in the spring, summer and fall of 2005. The CMB (chemical mass balance) receptor model (Watson *et al.*, 1997) was then applied to identify the potential source contributions to $PM_{2.5}$ mass.

MEASUREMENTS

Sampling sites and periods

The sampling sites are located in Pingtung city (urban site) and Chao-Chou town (rural site), in the northern and central parts of Pingtung County, respectively (Fig. 1). Pingtung city, with a population of about 216,222 and an area of 65.1 km², at a distance of about 22 km from the west coast, is the capital of Pingtung County. Chao-Chou is

primarily an agricultural town, with a population of about 57,189 and an area of 42.4 km² and is at a distance of about 14 km from the west coast. Taiwan's EPA has set up air-quality monitoring stations at the two sites. Therefore, hourly air quality and meteorological data. including PM_{10} on (particles with an aerodynamic diameter of under 10 µm), temperature and winds, are available. The two sites are approximately 16 km apart.

Samples were obtained over 16 days simultaneously at both sites, in spring (March 8–9, April 29–May 2), summer (June 21–June 22, July 12–13, August 9–10) and fall (September 13–14, October 12–13) in 2005. Each day comprised two 12h sampling periods–daytime (07:00–19:00) and nighttime (19:00–07:00). Given the limited budget and period of the project, the experiments were not conducted in winter. Table 1 presents the meteorological conditions at the sampling sites, including temperature, wind speed, period of sunshine and relative humidity.

Methods of sampling and analysis

At the Pingtung and Chao-Chou sites, a manual dichotomous sampler (G241 Model, Graseby Anderson) and a Universal Air Sampler (Model 310, MSP Corporation), packed with Teflon and Quartz filters, respectively, were adopted. The dichotomous sampler was maintained at a total air flow rate of 16.7 L/min–15 L/min for PM_{2.5} and 1.7 L/min for PM_{2.5-10} (particles with diameters of between 2.5 and 10 μ m). The Universal Air Sampler was maintained at a total air flow rate

of 285 L/min–270 L/min for $PM_{2.5}$ and 15 L/min for $PM_{2.5-10}$. Notably, the concentration of PM_{10} is the sum of those of $PM_{2.5}$ and $PM_{2.5-10}$. Before sampling, filters were placed in box at temperature of $25 \pm 3^{\circ}C$ and relative humidity of $40 \pm 5\%$, and then weighed after two days. An electrical balance with a precision of 0.01 mg was adopted to weigh the blank filters and particles.

Eight water-soluble ions including Na⁺, K⁺, Mg^{2+} , Ca^{2+} , NH_4^+ , Cl^- , NO_3^- and SO_4^{2-} in PM_{2.5} were analyzed. The water-soluble ions were extracted using an ultrasonic bath (UC-300, Sturdy) for 120 minutes. All extraction solutions were filtered through a cellulose acetate filter (ADVANTEC MFS, Inc., USA cat No., CO20A025A; pore size, 0.2 µm; diameter, 25 mm). The inorganic species were analyzed by ion chromatography (IC) (DIONEX 120 with an electrochemical detector (DIONEX CDM-1)). Cations were analyzed using a guard column (Shim-pack CG12A-SC, DIONEX) and an analytical column (Shim-pack CS12A-SC, DIONEX). Anions were analyzed using a Shim-pack AG4A guard column and a Shim-pack AS4A analytical column. The eluents used in cation and anion analyses were 20 mM methane sulfonic acid and 1.8 mM sodium carbonate + 1.7 mM sodium bicarbonate, respectively. Analytical drift was monitored throughout the analytical procedure. Recovery efficiencies were measured and analyzed using diluted samples spiked with known quantities of the studied ions. Recovery efficiencies of 93-107% were obtained. Both field and laboratory blank samples were prepared and analyzed; all data were corrected with reference to a blank. The coefficient of determination, R^2 , of the calibration lines made by electrical conductivity measurements exceeded 99.9% for all species. The detection limits of the analysis for each species, when converted to atmospheric concentration in ng/m³, were 28 (Na⁺), 48 (K⁺), 52 (Mg²⁺), 42 (Ca²⁺), 108 (NH₄⁺), 186 (Cl⁻), 114 (NO₃⁻) and 246 (SO₄²⁻).

Total carbon (TC) and elemental carbon (EC) were analyzed using an elemental analyzer (TOC-500A, Shimadzu). A quarter of each sample filter was heated in advance in a 340°C oven for 100 min to expel the organic carbon (OC) content, and then fed into the elemental analyzer to determine the EC content. The heating time adopted herein (100 min) was close to that (120 min) adopted by Cachier et al. (1989) and longer than the reduced heating time (43 min) used by Lavanchy et al. (1999). Another quarter of each sample filter was fed directly into the elemental analyzer without any pretreatment to determine the TC concentration. OC was then determined by subtracting EC from TC: OC = TC – EC. Similar methods were used by others, for example, Cadle and Mulawa (1990), Harrison et al. (1997) and Choi et al. (2004). The detection limits of the analysis for OC and EC were 0.024 µg. Notably, since O associated with OC mostly exists in carbonyl compounds in atmosphere, their concentrations are usually much smaller than those of alkane compounds. Also, since mass contribution by H is also much smaller than that by C, the mass contribution to OC due to H and O was therefore neglected here.

Nineteen metal elements were analyzed by

Profile name/number	Abbreviation	Descriptions and references
Marine source	Marin	Marine source— Southern Taiwan; Chen <i>et al.</i> (2001)
Vehicle exhaust	Auto1	Gasoline vehicle—motorcycles; Chen <i>et al.</i> (2001)
Vehicle exhaust (profile #31107)	Auto2	Light duty vehicle with catalyst; USEPA (2002)
Vehicle exhaust (profile #32208)	Auto3	Heavy duty diesel trucks; USEPA (2002)
Vehicle exhaust (profile #32103)	Auto4	Light duty vehicles —diesel; USEPA (2002)
Vehicle exhaust	Auto5	Heavy duty diesel trucks; Chen et al. (2006)
Municipal incinerator	Incer1	Municipal incinerator—Kaohsiung, Taiwan; Chen <i>et al.</i> (2001)
Municipal incinerator (profile #17105)	Incer2	Municipal incinerator—Philadelphia, USEPA (2002)
Field burning (profile #42304)	Burn1	Agricultural field burning; USEPA (2002)
Field burning (profile #42324)	Burn2	Field burning—Fescue; USEPA (2002)
Field burning	Burn3	Agricultural field burning; Lin (2005)
Secondary reactant (profile #25409)	Amnit	Ammonium nitrate; USEPA (2002)
Secondary reactant (profile #25410)	Amsul	Ammonium sulfate; USEPA (2002)

 Table 2. Possible PM2.5 source profiles in CMB model.

ICP-MS (Inductively Coupled Plasma with Mass Spectroscopy, Agilent 7500 series) and ICP-AES (Atomic Emission Spectrometer, Perkin Elmer Optima 2000DV for Si) after being treated in a microwave digest oven and by acid-stripping. The microwave process was two cycles; each cycle involved two operating powers, which were 1% of 300 W (twice, 2.5 min per time) and 100% of 1200 W (three times, 2.5 min per time), following the power series 1200-300-1200-300-1200W. The rest time between each cycle was also 2.5 min. This process yields an operational temperature of over 160°C to ensure the complete digestion of the particles collected on the Teflon filters. Then, the digested solution was diluted to a volume of 25 mL using ultra-pure water (specific resistance ≥ 18.3 M Ω cm) to perform the metal analysis of Ag, Al, Ba, Ca, Cd, Cr, Cu, Fe, K, Mg, Mn, Na, Ni, Pb, S, Si, Sr, Ti and Zn by ICP-MS and ICP-AES. The calibration was made using multi-element (metal) standards (certified reference materials (CRMs); Spex, Metuchen, USA) in a 1% (v/v) HNO₃ solution. tenth sample was Everv spiked with liquidstandards with particular amounts of identified metal elements (Baker Co.). The efficiency of acid digestion was in 90-95% (Allen et al., 2002; Lin et al., 2005). The CRMs were also used as quality control standards. Analytical drift was monitored throughout the efficiencies procedure. Recovery were determined and analyzed using a diluted sample spiked with a known quantity of metal. Recovery efficiencies from 80 to 120% were achieved. The detection limits of the analysis for metal elements, when converted to atmospheric concentration in ng/m^3 , were 67.15

Source profile	Base	1	2	3	4	5
(a) Pingtung site						
Marin	16.19	29.06	14.38	14.46	14.67	13.34
Autol		-2.85				
Auto2	11.69		49.31	53.89	46.74	43.52
Auto3		42.54	7.82			
Auto4					6.73	7.38
Auto5	24.96			2.41		
Incer1						1.80
Incer2	1.89					
Amnit	16.13	15.14	12.39	12.14	12.51	12.67
Amsul	17.45	12.43	13.82	13.89	14.08	14.30
R^2	0.94	0.93	0.98	0.98	0.98	0.98
χ^2	2.08	1.47	0.21	0.22	0.19	0.23
Mass accounted (%)	101.9	110.2	111.9	110.8	108.4	106.5
(b) Chao-Chou site						
Marin	7.22	13.06	11.89	5.58	10.48	11.31
Autol	5.56	5.74		6.35	5.20	0.93
Auto2			37.98			33.58
Auto3					24.79	
Auto4				6.18		
Auto5	13.89	21.18	7.31			7.64
Burn1		17.12				
Burn2	31.77		11.15	38.61	13.08	13.57
Amnit	14.18	13.99	12.92	14.03	14.27	13.04
Amsul	15.12	14.00	14.40	15.08	14.52	14.47
R^2	0.96	0.96	0.98	0.85	0.95	0.98
χ^2	2.00	1.79	0.30	6.23	1.69	0.39
Mass accounted (%)	100.4	97.4	109.5	98.2	94.2	108.2

Table 3. Sensitivity of CMB results to combinations of source profiles.

(Na), 0.45 (Mg), 2.88 (Al), 83.50 (Si), 250.0 (S), 3.85 (K), 9.93 (Ca), 0.63 (Ti), 0.26 (Cr), 0.20 (Mn), 1.06 (Fe), 0.17 (Ni), 0.07 (Cu), 0.62 (Zn), 1.25 (Sr), 0.16 (Ag), 0.17 (Cd), 0.19 (Ba) and 0.13 (Pb).

RECEPTOR MODELING

Analysis of source contributions

Calculations of the contributions of sources were based on the CMB8 receptor model (Watson *et al.*, 1997). This model comprises a solution to linear equations that express the concentration of each species at a receptor site as a linear sum of products of source profile abundances and source contributions. The source profile abundances (or fingerprints) and the concentrations of the species must first be



Date

Fig. 2. Mean concentrations of PM_{10} and $PM_{2.5}$ and $PM_{2.5}/PM_{10}$ vs. sampling date in 2005 at (a) the Pingtung site, and (b) the Chao-Chou site.

determined as the inputs to the CMB model. Then, the individual source contributions can be determined using multiple regression techniques. The performance of the CMB8 model is evaluated using several statistical indices, including the coefficient of determination ($R^2 \ge 0.8$), CHI squared ($\chi^2 \le$ 4.0), the percentage of mass accounted for (0.8-1.2), T-statistics (≥ 2.0) and the uncertainty ratio (-2.0 to 2.0). These values are employed herein.

Source profiles and sensitivity analysis

Multivariate factor analysis was adopted to

help identify dominant source categories, from which three principal factors were obtained that explained about 61% and 56% of the total variance at the Pingtung and the Chao-Chou sites, respectively (Chen et al., 2006). Based on the dominant species in individual factors and current source profiles, the three principal sources are considered to be vehicle exhaust, marine aerosols and incinerators in Pingtung, and vehicle exhaust, outdoor burning of agricultural plants and marine aerosol in Chao-Chou. A pool of 12 profiles associated with these three pollution sources was selected (Table 2); five files (Marin, Auto1, Auto5, and Incer1) were obtained from the local data, and others were obtained from the SPECIATE 3.2 library files (USEPA, 2002).

Prior to performing receptor modeling, sensitivity analyses were performed to select quasi-optimal combinations of source profiles from the pool to generate overall best model performance (Fujita et al., 1994). Table 3 presents typical sensitivity results for the daytime (07:00-19:00) data on 8 March 2005 at Pingtung and Chao-Chou sites, indicating that the "Base" case yielded better results than the other five cases. Therefore, the CMB results presented herein are all derived from the six source profiles (Marin, Auto2, Auto5, Amsul, Amnit, Incer2) in the "Base" case at the Pingtung site in Table 3; and from the six source profiles (Marin, Auto1, Auto5, Amsul, Amnit, Burn2) in the "Base" case at the Chao-Chou site. Notably, the mass fractions in Amsul (ammonium sulfate, (NH₄)₂SO₄) are 72.7% for SO_4^{2-} and 27.3% for NH_4^+ , while those in Amnit (ammonium nitrate, NH₄NO₃) are 77.5% for NO_3^- and 22.5% for NH_4^+ .

RESULTS AND DISCUSSION

Concentrations of PM₁₀ and PM_{2.5}

Figs. 2(a) and 2(b) plot the concentrations of PM_{10} and $PM_{2.5}$ and the $PM_{2.5}/PM_{10}$ ratio vs. the sampling date, for the Pingtung and Chao-Chou sites, respectively. The results were obtained over three seasons (spring, summer, and fall) in 2005. The highest concentrations of PM_{10} and PM_{25} were 104 and 91.6 μ g/m³ on March 9 at Pingtung site, and 140 and 122 μ g/m³ on March 8 at Chao-Chou site. The overall means of PM_{10} and PM_{2.5} were 59.2 and 47.4 μ g/m³ respectively at Pingtung, and 63.6 and 45.7 μ g/m³ respectively at Chao-Chou. The concentrations of PM₁₀ and PM_{2.5} in the early spring, fall and winter were about two to three times those in the late spring and summer, because fall, winter and/or early spring are typically the worst periods of air quality in southern Taiwan, in which dryness, little rain and low mixing heights reduce the ability of the atmosphere to dilute the airborne pollutants (Chen et al., 2004).

However, unlike the PM_{10} or the $PM_{2.5}$ mass, the $PM_{2.5}/PM_{10}$ ratio varied insignificantly with the season. On average, $PM_{2.5}/PM_{10}$ was $0.81 \pm$ 0.07 at Pingtung site, and 0.68 ± 0.11 at Chao-Chou site. The correlation coefficient (*R*) between $PM_{2.5}$ and the PM_{10} masses was 0.98 at Pingtung site and 0.78 at Chao-Chou site, indicating that the mass of $PM_{2.5}$ dominated that of PM_{10} . Although Pingtung site has a higher $PM_{2.5}/PM_{10}$ ratio than Chao-Chou site, the presented ratios are toward the highest end of the



Fig. 3. Mean concentrations of eight water-soluble ions in $PM_{2.5}$ vs. sampling date in 2005 at (a) the Pingtung site, and (b) the Chao-Chou site.

reported results. For instance, the $PM_{2.5}/PM_{10}$ ratios were 0.41–0.81 at 14 sites in the Central California (Chow *et al.*, 1996; 1999), 0.71–0.77 at eight sites in and around metropolitan Philadelphia (Burton *et al.*, 1996), 0.57–0.71 at

seven sites in Kaohsiung (Lin, 2002), about 0.72 at an urban site and a remote site in the eastern United States (Vukovich and Sherwell, 2002), 0.61–0.78 at two sites in Hong Kong (Ho *et al.*, 2003), 0.63–0.77 at five sites in Nanjing (Wang,

Species	Pingtur	ng	Chao-Chou	
	Mean (%)	SD (%)	Mean (%)	SD (%)
Na^+	1.01	1.49	0.92	1.24
K^+	0.81	0.41	0.98	0.43
Mg^{2+}	0.34	0.48	0.25	0.38
Ca^{2+}	0.58	1.26	0.42	0.32
$\mathrm{NH_4}^+$	9.65	3.03	8.17	4.86
Cl^{-}	2.96	2.47	3.15	1.81
$\mathrm{NO_3}^-$	11.10	4.07	10.4	5.24
$\mathrm{SO_4}^{2-}$	17.80	4.58	18.2	4.81
EC	9.60	3.16	10.00	2.49
OC	21.40	5.24	22.30	7.86

Table 4. Mass fraction and standard deviation (SD) of eight water-soluble ions and carbon contents (OC and EC) in $PM_{2.5}$ at the Pingtung and Chao-Chou sites.

Table 5. Mass fractions and standard deviation (SD) of 19 metal elements in $PM_{2.5}$ at the Pingtung and Chao-Chou sites.

Species	Pingtung		Chao-Chou		
	Mean (%)	SD (%)	Mean (%)	SD (%)	
Na	0.950	0.770	0.810	0.660	
Mg	0.230	0.190	0.230	0.210	
Al	0.560	0.380	0.670	0.580	
Si	0.590	0.410	1.520	0.970	
S	7.600	1.620	7.530	2.210	
Κ	0.760	0.460	0.580	0.410	
Ca	0.770	0.790	0.820	1.590	
Ti	0.010	0.010	0.010	0.004	
Cr	0.010	0.010	0.020	0.030	
Mn	0.030	0.020	0.030	0.040	
Fe	0.490	0.370	0.360	0.430	
Ni	0.210	0.520	0.070	0.090	
Cu	0.030	0.020	0.020	0.030	
Zn	0.300	0.280	0.340	0.240	
Sr	0.010	0.001	0.010	0.005	
Ag	0.020	0.080	0.002	0.002	
Cd	0.001	0.001	0.001	0.001	
Ba	0.010	0.010	0.010	0.010	
Pb	0.090	0.140	0.060	0.050	

et al., 2003), 0.75–0.76 at six sites in Switzerland (Gehrig *et al.*, 2003), 0.63–0.73 in

Taichung (Fang *et al.*, 2003), 0.45–0.73 at three sites in Beijing (Sun *et al.*, 2004), and 0.4–0.6 in

Santiago in Chile (Koutrakis et al., 2005). Many factors, including traffic and industrial activity, emission patterns/strengths and meteorological conditions affect the $PM_{2.5}/PM_{10}$ ratios. The PM_{2.5}/PM₁₀ ratios are typically (though not always) low (< 0.6) near traffic sites, in relatively dry areas or in the seasons with relatively strong winds such that coarse particles (particles with diameter larger than 2.5 µm) are inclined to being re-suspended in the air (Sun et al., 2004; Koutrakis et al., 2005). Notably, although daytime concentrations of PM₁₀, PM_{2.5} or PM_{2.5}/PM₁₀ ratios may differ from those at night (not plotted), the bulk differences between the two periods were negligible (with a difference of about $\pm 3\%$) at each site.

*Water-soluble ions and carbonaceous contents in PM*_{2.5}

Figs. 3(a) and 3(b) present the concentrations of eight water-soluble ions vs. the sampling date at the Pingtung and Chao-Chou sites, respectively, indicating that the total concentrations of water-soluble ions generally followed the variation in PM2.5. Table 4 presents the mass fractions of individual constituents (including OC and EC) in PM2.5 mass for all samples at both sites. Both sites had the same leading abundant species in PM_{2.5} mass, although contributions varied as follows: OC (21.4%), SO₄²⁻ (17.8%), NO₃⁻ (11.1%), NH₄⁺ (9.70%) and EC (9.6%) at Pingtung, and OC (22.3%), EC (10.0%), SO_4^{2-} (8.3%), NO_3^{-} (4.7%) and NH_4^+ (1.4%) at Chao-Chou. Therefore, three secondary aerosols $(SO_4^2)^{-1}$, NO_3^{-} and NH_4^{+}) (and TC = OC + EC) represented about 38.4% (31.0%) of the PM_{2.5}

mass at Pingtung, and 36.5% (32.4%) of the PM_{2.5} mass at Chao-Chou, suggesting that traffic exhausts (relating to OC and EC) and three secondary aerosols were the most important sources of atmospheric PM_{2.5} at both sites; altogether, they represented about 70% of PM_{2.5} mass at both sites. The sub-leading abundant species at the two sites were Cl⁻ (1.8-2.5%) and Na⁺ (0.14-1.5%); Ca²⁺, K⁺ and Mg²⁺ had abundances of 0.1-1.3%. In summary, water-soluble ions and carbonaceous species represented about 70% of the PM_{2.5} mass at both sites.

The presented results are similar to and/or slightly higher than those of Chen *et al.* (2001), which indicated that the three secondary aerosols (SO_4^{2-} , NO_3^{-} and NH_4^{+}) contributed about 30% of the PM_{2.5} mass, while OC and EC contributed about 20% of the PM_{2.5} mass in neighboring Kaohsiung city.

Metals in PM_{2.5}

Table 5 presents the mass fractions of 19 metals in PM_{2.5} mass for all samples at the two sites. The ten most abundant species are S (7.6%, 7.53%), Na (0.95%, 0.81%), Ca (0.77%, 0.82%), K (0.76%, 0.58%), Si (0.59%, 1.52%), Al (0.56%, 0.67%), Fe (0.49%, 0.36%), Zn (0.3%, 0.34%), Mg (0.23%, 0.23%) and Ni (0.21%, 0.07%); the first value in parenthesis refers to Pingtung, and the second refers to Chao-Chou. These metals mostly originate in the earth's crust and traffic exhausts; in particularly, Zn and Ni are closely related to automobile/truck exhausts and fuel burning (Hillamo *et al.*, 1993; Hueglin *et al.*, 2005). Industrial sources were thought to have little impact on the concentrations of these



Fig. 4. Box plots of source contributions (% of calculated mass) to $PM_{2.5}$ mass during spring, summer and fall at the (a) Pingtung site (Source symbol: 1: marine aerosol; 2: gasoline vehicles; 3: diesel trucks; 4: ammonium sulfate; 5: ammonium nitrate; and 6: incinerators), and (b) Chao-Chou site (Source symbol: 1: marine aerosol; 2: motorcycles; 3: diesel trucks; 4: ammonium sulfate; 5: ammonium nitrate; and 6: outdoor burning).

metals since no large industrial factories are located here. Meanwhile, nine other metals are minor, ranging from 0.001 to 0.09%. Altogether, the 19 metals contributed about 12.5% and 13.1% of the $PM_{2.5}$ mass at Pingtung and Chao-Chou, respectively.

In summary, water-soluble ions, carbonaceous

contents (OC and EC) and metal elements represented about 82% of the $PM_{2.5}$ mass at both sites. Some of the 18% of the $PM_{2.5}$ mass that was not accounted for was associated with the unanalyzed species and some was associated with measurement errors, such as the gap between gravimetric and chemical



Fig. 5. Scatter plots of calculated vs. measured concentrations for five species at (a) the Pingtung site, and (b) the Chao-Chou site. Solid line represents the best fit.

measurements (Andrews et al., 2000).

Source contributions to PM_{2.5} mass using CMB model

Figs. 4(a) and 4(b) are box plots of the percentage source contributions to the $PM_{2.5}$ mass for three seasons (spring, summer and fall) at Pingtung and Chao-Chou sites, respectively. The figures reveal that six sources were

identified with each site, although source categories and their contributions differ somewhat between the sites. They are marine salt, gasoline vehicles, diesel trucks, ammonium sulfate, ammonium nitrate and incinerators at Pingtung site, and marine salt, diesel trucks, ammonium sulfate and ammonium nitrate at Chao-Chou site, along with motorcycles and outdoor burning, rather than gasoline vehicles and incinerators.

At the Pingtung site, as in Fig. 4(a), gasoline vehicles (27.5 - 45.8%),diesel trucks (12.1–21.8%), ammonium sulfate (21.1–26.3%) and ammonium nitrate (6.3-14.5%) are the leading four contributors to the PM_{2.5} mass in the three seasons, although the order of the three contributors other than gasoline vehicles may vary with the season. The other contributors are marine aerosol (3.8-6.5%) and incinerators (0.7-1.4%). In summary, vehicle exhaust (49.3-62.4%) from gasoline and diesel cars and three secondary aerosols (31.2–37.8%) are the most important contributors at the Pingtung site. Meanwhile, at the Chao-Chou site in Fig. 4(b), outdoor burning (25.3-50.4%) and ammonium sulfate (16.5-25.3%) are the two leading contributors in the three seasons. In particular, the contribution of outdoor burning in the summer (47.6%) and the fall (50.4%) are thought to be related to the field-burnings of agricultural waste, usually observed during the reaping periods in Chao-Chou town. The next contributors are ammonium sulfate (7.8–11.3%), (1.8–19.4%), diesel trucks motorcycles (5.7-12.3%) and marine aerosol (6.3-8.2%) at the Chao-Chou site. In summary, outdoor burning (25.3–50.4%), secondary aerosols (27.2–34.3%) and vehicle exhaust (12.0–26.9%) from diesel trucks and motorcycles are the most important contributors at the Chao-Chou site.

The scatter plots in Figs. 5(a) and 5(b) present the correlations between the calculations and measurements of five dominant species, EC, OC, NO_3^- , SO_4^{2-} and NH_4^+ , at the two sites. The R^2 is 0.71–0.99 at Pingtung, and 0.74–0.99 at Chao-Chou,

indicating strong consistency between the calculated and the measured values.

CONCLUSIONS

Measurements reveal several similarities and differences between an urban site and a rural site. At both sites, the concentrations of $PM_{2.5}$ (also PM₁₀) varied seasonally, being worst during the fall, the winter or early spring, but the PM_{2.5}/PM₁₀ ratios varied insignificantly with the season. The abundant species in $PM_{2.5}$ mass were secondary aerosols (SO_4^{2-}, NO_3^{-}) and NH₄⁺) and carbonaceous species (OC and EC), followed by metals; together, they represented about 82% of the PM_{2.5} at each site. Although both sites exhibited strong correlations (R = 0.98 at Pingtung, and R = 0.78 at Chao-Chou) between PM₁₀ and PM_{2.5} masses, the mean $PM_{2.5}/PM_{10}$ ratio was 0.81 at Pingtung-higher than 0.68 at Chao-Chou, indicating that relatively large bare lands and outdoor burning on farms may have caused more coarse particles to be present in PM_{2.5} at a rural site (Chao-Chou) than at an urban site (Pingtung).

The similarities and differences indicated above were also revealed by the CMB calculations. For instance, at both sites, vehicle exhaust and secondary aerosols were dominant contributors to $PM_{2.5}$ mass, while marine aerosol contributed only 3.8 to 8.2%. However, the contribution of vehicle exhaust (49.3–62.4%) at Pingtung (urban) site was about 50 to 100% higher than (12.0–26.9%) at Chao-Chou (rural) site, depending on the season; the contribution (25.3–50.4%) of the outdoor burning of agricultural waste contributed greatly at the Chao-Chou (rural) site.

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