Characterization of PM, PAHs and Gaseous Pollutants Emitted from Sintering Process and Electric Arc Furnace

Tang Wei Chen¹, Jyh Cherng Chen², Zhen Shu Liu³, Kai Hsien Chi⁴, Moo Been Chang*¹

¹ Graduate Institute of Environmental Engineering, National Central University, Taoyuan, Taiwan
² Department of Environmental Engineering and Science, Feng Chia University, Taichung, Taiwan
³ Department of Safety Health and Environmental Engineering, Ming Chi University of Technology, New Taipei, Taiwan
⁴ Institute of Environmental and Occupational Health Sciences, National Yang Ming Chiao Tung University, Taipei, Taiwan

ABSTRACT

Iron and steel industry is crucial for economic development. However, large exhaust volume and complex feeding of the steel-making process results in enormous emissions of air pollutants such as particulate matter (PM), sulfur oxides (SOₓ), nitrogen oxides (NOₓ), polycyclic aromatic hydrocarbon (PAHs) and carbon monoxide (CO). This study sampled and analyzed the concentrations of PM, PAHs, SOₓ, NOₓ and CO at the inlet and outlet of the air pollution control devices (APCDs) adopted in sintering process (SP) and electric arc furnaces (EAF), respectively, to evaluate the performance of APCDs and emission characteristics of these air pollutants. The results show that filterable PM (FPM) concentrations measured at stacks of SP and EAF are 8.2 ± 1.3 and 13.6 ± 2.2 mg Nm⁻³, respectively. The FPM₂₅ concentrations emitted from SP and EAF are 1.8 ± 0.5 and 3.2 ± 1.1 mg Nm⁻³, respectively, while the condensable PM (CPM) concentrations emitted from SP and EAF are measured as 37.7 ± 1.3 and 3.4 ± 0.8 mg Nm⁻³, respectively. Moreover, the flue gas desulfurization (FGD) and selective catalytic reduction (SCR) equipped in SP removed 82% and 58% of SOₓ and NOₓ, respectively, whereas SOₓ and NOₓ emitted from EAF are relatively low (< 1 ppm). The overall PAH removal efficiencies achieved with the APCDs adopted by SP and EAF are 72% and 26%, respectively. For total toxicity equivalency quantity (TEQ) PAH concentrations, similar results for SP (5.45 µg BaPₑq Nm⁻³) and EAF (4.74 µg BaPₑq Nm⁻³) are obtained. Moreover, the diagnosis ratios of FL/CHR, FL/(BaA + CHR), Pyr/S-MC and Pyr/(BaA + CHR) are suggested as the indicators for EAF. The emission factors calculated indicate that operating conditions of the APCDs in the SP investigated are well controlled. The concentrations of air pollutants emitted from SP investigated are lower than other studies. On the other hand, the concentrations of FPM and PM₂₅ emitted from EAF are higher than other studies. However, NOₓ, SOₓ and PAHs are well controlled. Moreover, the CO emission factors calculated from these two steel-making processes are higher than most reports. Overall, the operation of APCDs should be improved to reduce air pollutants emitted from these two steel-making processes.

Keywords: Iron and steel industry, Nitrogen oxides (NOₓ), Sulfur oxides (SOₓ), Particulate matter (PM), Polycyclic aromatic hydrocarbons (PAHs)

1 INTRODUCTION

Iron and steel industry is crucial for economic development. According to the statistics released by the World Steel Association, annual global crude steel production reached 1,869 million tonnes in 2019, and the output has been increasing since 1950 (Association 2020). Accounting
for 50% of global production, China is the largest iron and steel producing country. The crude steel production in Taiwan (22.4 million metric tons in 2017) ranked eleventh in the world, accounting for 1.33% of global output, of which crude steel produced by SP and EAF accounted for 61% and 39%, respectively. According to the statistical data released by the Taiwan EPA, PM, PM$_{2.5}$, SO$_x$, NO$_x$, and CO emitted from the steel-making process accounted for 15.9%, 17.5%, 16.7%, 13.0%, 8.31% and 82.1% of industrial emissions (TEDS 10.0, 2016) in Taiwan, respectively, confirming the importance of the steel-making process as a source of air pollutants. Previous studies also indicate that iron and steel industries are major contributors of PM and persistent organic pollutants (POPs) in the vicinity (Amadio et al., 2012; Van Thuong et al., 2014; Xiao et al., 2013).

Particulate matter (PM) including filterable and condensable PM is one of the most important pollutants emitted from steel-making processes. FPM comprises of the particles emitted directly from a source as a solid or liquid at release conditions and is captured on the filter of a stack sampling train. On the other hand, CPM consists of material in the vapor phase at stack conditions that condenses and reacts upon cooling and dilution in the ambient air to form solid on liquid PM after exiting the stack. CPM passes through the filter and can be captured by the impingers of the sampling train. PM$_{2.5}$ emissions would be greatly underestimated if CPM is ignored since CPM accounts for a significant portion of total PM$_{2.5}$ emissions (Yang et al., 2014; Yang et al., 2015).

Steel-making process can be classified into sintering process (SP) and electric arc furnace (EAF) process. The former primarily uses iron ore as raw material for making iron in a blast furnace and steel in a basic oxygen furnace; the latter uses scrap steel as the primary feeding material. Because the feeding of iron and steel-making process is complex and requires large volume of raw materials, air pollutants are inevitably produced. The concentration of air pollutants emitted from steel-making processes varies greatly, depending on operating conditions and APCDs adopted. In addition to PM, the iron and steel industry is also one of the primary sources of SO$_x$ and NO$_x$ emissions in China (Li et al., 2019a). Tang et al. (2020) investigated the emissions of air pollutants from iron and steel industry in China, indicating that compared with coke making process, basic oxygen and blast furnaces, the sintering process was the largest emitter among the iron and steel production processes in China, accounting for 69.52% of SO$_x$, 75.48% of NO$_x$, 45.81% of PM$_{10}$, and 47.99% of PM$_{2.5}$ emissions, respectively. Additionally, Yang et al. (2015) measured the concentrations of PM$_{2.5}$ and water-soluble ions emitted from steel plants, indicating that sintering and EAF has high FPM$_{2.5}$ and CPM emission concentrations, while SO$_x$ is the highest water-soluble ion emitted from sintering and EAF. Yu et al. (2018) investigated the correlation of CPM with SO$_x$ concentration and indicated that CPM tended to increase as SO$_x$ concentration increased. Li et al. (2019a) confirmed that SO$_x$ and NO$_x$ are primary precursors of fine PM in the environment. Thus, the removal efficiencies of SO$_x$ and NO$_x$ achieved with the APCDs not only affect the emissions of SO$_x$ and NO$_x$ but also CPM. Therefore, it is important to investigate the characteristics of air pollutants emitted from SP and EAF. Nevertheless, simultaneous sampling and analysis of FPM$_{2.5}$ and CPM emissions from the steel-making process are not well documented so far.

On the other hand, the rapid expansion of global steel-making industry inevitably puts tremendous pressure on the environment. For instance, PAHs are incomplete combustion substances generated in the iron and steel-making processes. PAHs are mainly formed via pyrolysis of hydrocarbons, cyclization of chain alkanes or single-ring aromatics, condensation and polymerization of low-carbon PAHs (Zhang et al., 2020). In the SP and EAF steel-making process, PAHs may be present in the scrap steel and are volatilized during production processes or they may form as a result of incomplete combustion of residual organic matter, fuels and additives. Wang et al. (2020) investigated PM$_{2.5}$-bound PAHs in ambient air in Anshan of China during 2014–2015, indicating that the concentrations of 16 PAHs in steel-making dust were much higher than that from other emission sources. Sintering process has long been recognized as a significant source of PAHs emissions due to the input of a large amount of coal and iron ore. According to the statistics, approximately 42 and 22 tons PAHs were emitted from sintering process per year in the USA and UK, respectively (Chen et al., 2012). In addition, previous study showed that coal combustion and steel-making dust contributed to over 86% of the total cancer risk linked to PAHs exposure (Wang et al., 2020). Hence, sintering process and electric arc furnaces have been identified as important organic pollutant sources (Odabasi et al., 2009; Amadio et al., 2013). Because PAHs are toxic and pose great adverse impact on human health and environment, numerous studies have been
conducted to investigate PAHs emissions from iron and steel processes (Odabasi et al., 2009; Amodio et al., 2013; Van Thuong et al., 2014; Khaparde et al., 2016). In fact, 16 PAHs have been listed as the priority pollutants by the U.S. Environmental Protection Agency (U.S. EPA) and the International Agency for Research on Cancer (IARC) and classified by the latter into four categories based on mutagenicity and carcinogenicity (Yang et al., 2002). Currently, most studies on PAHs focused on the 16 PAHs prioritized for regulation by the U.S. EPA, but few studies have focused on other PAHs. The TEF values for the 11 PAHs are relatively high, some are even higher than the 16 PAHs regulated by U.S. EPA. PAHs may have equal numbers of rings, but the configurations and arrangements of the rings bring about differences in properties such as chemical and physical properties (Cheruiyot et al., 2015). Therefore, the European Union (EU) and the EU Scientific Committee on Food (SCF) have listed 24 and 15 PAHs, respectively, as the priority targets for control with high hazards. This study investigates 27 PAHs, including U.S. EPA, EU and SCF priority control species to better understand the characteristics of PAHs emitted from steel-making processes. On the other hand, CO is often considered as an indicator of incomplete combustion. Previous studies have shown that when the concentration of CO increases, the concentration of PAHs increases as well, indicating CO concentration has a strong correlation with PAHs (Blumenstock et al., 2000; Oh et al., 2007). According to the Taiwan EPA, the CO emission from the steel industry reached 188,196 ton yr⁻¹. Based on the comprehensive emission inventory of air pollutants in China’s iron and steel industry, each ton of iron and steel produced by SP and EAF generates 22,000 and 9800 grams of CO, respectively (Wang et al., 2016).

Information on the emissions of FPM2.5, CPM, and PAHs from SP and EAF is rather limited. In this study, the concentrations of PM (FPM2.5 and CPM), PAHs, SOx, NOx and CO at the inlet and outlet of APCDs in SP and EAF, respectively, were simultaneously measured to evaluate the APCD performance and the emission characteristics of these pollutants. In addition, emission factors of these air pollutants are calculated and unique diagnostic ratios of PAHs emitted from the steel-making processes are identified.

2 METHODS

2.1 Description of the Sampling Locations

Concentrations of air pollutants at the inlet and outlet of APCDs in SP and EAF located in central Taiwan were measured to evaluate the performance of the APCDs adopted and the emission characteristics of the air pollutants. In the SP investigated, iron ore, flux, and recovery material are agglomerated to produce sintering ore at the capacity of 476 ton h⁻¹. The fuel in the SP contains coke and coking oven gas while that in EAF are natural gas and coking oven gas. The sampling locations and pollutants measured at SP and EAF are clearly marked in Fig 1. Dry electrostatic precipitator (DESP) is integrated with semi-dry FGD as well as SCR as APCDs for SP. The EAF is equipped with only baghouse (BH) as APCD. In this study, the sampling location of air pollutants (PAHs, NOx, SOx, CO, FPM, FPM2.5, and CPM) are clearly marked in Fig 1. On the other hand, the pollutant concentrations reported for the SP are corrected with the oxygen content of 15%, while oxygen content correction is not required for EAF.

2.2 Flue Gas Compositions Analysis

A portable flue gas analyzer (TESTO 350, Germany) was used to measure the concentrations of NOx, SO2, O2, CO, and CO2 at the inlet and outlet of the APCDs. The resolutions of O2, CO2, CO, NOx, and SO2 are 0.01 vol.%, 0.01 vol.%, 1.0 ppm, 0.1 ppm, and 0.1 ppm, respectively. The analyzer was calibrated with standard gases before sampling. The PM and water vapor were firstly removed through the filter and water vapor removal devices, respectively, and the flue gas was subsequently introduced into the gas analyzer for analysis. On the other hand, the data of NH3 concentrations emitted were provided by the plant based on the routine measurement.

2.3 FPM2.5 and CPM Sampling and Analysis

FPM2.5 was measured based on the U.S. EPA method 201A (Determination of PM10 and PM2.5 emissions from stationary sources) while U.S. EPA method 202 (the dry impingers method for
determining condensable particulate emissions from stationary sources) was applied for CPM measurement. Fig. 2(a) shows the sampling train for FPM$_{2.5}$ and CPM measurements. Before sampling, the flow rate should be estimated to meet the isokinetic principle; and appropriate front nozzle for sampling the flue gas must then be selected. The gas stream passed through a PM$_{2.5}$ cyclone to remove the PM with the aerodynamic diameter greater than 2.5 µm and PM$_{2.5}$ were collected on a 47 mm quartz filter (Tissuquartz, 2500 QAT-UP). The filter was conditioned at a relative humidity of 30–40% and temperature of 20–23°C for 24 h before and after sampling.

In this study, the FPM$_{2.5}$ and CPM sampling devices were connected in series for simultaneous sampling of FPM$_{2.5}$ and CPM. After the collection of FPM$_{2.5}$ with a 47 mm quartz filter, the gas stream passed through a condenser; and CPM was subsequently collected in the water dropout impinger, the modified Greenburg Smith impinger and CPM filter (Advantec, J020A090C PTFE). After sampling, the impinger and CPM filter were immediately purged with ultra-high purity nitrogen for 1 h to remove the dissolved sulfur dioxide. Deionized water and n-hexane were used to rinse the whole samples for obtaining inorganic (water soluble) and organic (organic solvent soluble) fractions of CPM, which were dried and weighed in the laboratory. The summation of these two fractions results in CPM concentration.

2.4 PAH Sampling and Analysis

The U.S. EPA Method 23A was applied for sampling PAHs and Fig. 2(b) shows the PAH sampling train which includes a front nozzle, a stainless probe, a filter holder, a condenser, an XAD-2 adsorbent and an impinger. The flue gas containing PAHs was sampled and passed through a
Fig. 2. Sampling train of (a) FPM$_{2.5}$ and CPM (U.S. EPA Method 202 A), and (b) PAHs (U.S. EPA Method 23A).

heated filter holder packed with a 90 mm quartz filter (Tissuquartz, 2500 QAT-UP) to collect FPM and solid-phase PAHs, and through a condenser to capture the remaining gas-phase PAHs with XAD-2. Next, the filter containing solid-phase PAHs and the XAD-2 resin containing gas-phase PAHs were extracted with dichloromethane (Fisher, 99.9%) for 22 ± 2 h using the Soxhlet extraction process. Before the extraction process, an isotope-spiked solution of PAHs was added into the samples to quantify the PAH concentration. The extraction solution was then concentrated to 2 mL using a rotary evaporator at 45°C and then replaced by 8 mL hexane, which was purified with a silica gel cleanup procedure. These samples were later analyzed using a gas chromatography/mass spectrometry (GC/MS) (Thermo Fisher Scientific, Double Focusing Sector Mass Spectrometer) with a fused silica capillary column DB-5 MS (60 m × 0.25 mm × 0.25 µm) under selected ion monitoring (SIM) mode conditions. The 27 PAHs containing 16 U.S. EPA PAHs, 15 EU PAHs, and three additional PAHs are analyzed in this study. Detailed information regarding the physical and chemical properties and the toxic equivalency factors (TEFs) of 27 PAHs analyzed are presented in Table S1. The TEFs are used to calculate the toxic equivalent (TEQ) concentration of 27 PAHs and assess the carcinogenic risk of PAHs. The TEF is based on the toxicity of BaP. The calculation is described as follows:

\[
BaP_{eq} = \sum_{i}^{n} PAH_i \times TEF_i
\]

where \(PAH_i\) is the concentration of the individual PAH in each sample, and \(TEF_i\) is the TEF of the individual PAH.
The PAH concentrations were calculated using a group of six concentration levels ranging from 0.01 to 1 µg mL⁻¹. The calibration curves were linear in the concentration ranges, with the correlation coefficient (R²) of 0.995. The method detection limits of the 27 PAHs analyzed were determined from the lowest standard in the calibration curve. The mean recovery rate of the 27 PAHs analyzed ranged from 66% to 77%.

2.5 Emission Factor

The emission factor (EF) is used to estimate the pollutant emissions and is defined as the pollutants emitted to the atmosphere by industrial activity of the emission source. The calculation is described as follows,

\[
\text{Emission factor (g ton}^{-1}) = C \text{ (g Nm}^{-3}) \times Q \text{ (Nm}^3 \text{ hr}^{-1})/P \text{ (ton hr}^{-1})
\]  

C: Concentration of pollutant in flue gas (g Nm⁻³).
Q: Flow rate of flue gas (Nm³ hr⁻¹).
P: Production rate (ton hr⁻¹).

3 RESULTS AND DISCUSSION

3.1 Flue Gas Compositions

Table 1 shows the flue gas compositions measured at the inlet of APCD (DESP + FGD + SCR) and the stack of SP, respectively. The moisture, oxygen, and CO₂ contents measured at the SP stack are 20%, 13.9%, and 7.2%, respectively, which are close to that measured at the APCD inlet. Moreover, the CO concentrations measured at the APCD inlet and stack are 5,857 and 5,208 ppm, respectively, confirming that CO removal achieved with existing APCDs is insignificant. The flue gas compositions in the baghouse inlet and stack of EAF are also provided in Table 1. The concentrations of CO₂ and CO in the EAF stack are measured as 1.0% and 191 ppm, respectively. Sintering process agglomerates iron ore material with other recycle material such as ESP ash at high temperatures. Materials containing oily reductants (aliphatic hydrocarbons and aromatic hydrocarbons) and carbonaceous by-products are mixed, and varieties of pollutants may be generated during sintering process. The feeding material of electric arc furnace was mainly scrap steel while graphite electrode was applied to provide the heat to melt scrap steel. Although the EAF steel-making is a high temperature process, liquid oxygen was applied in the furnace and NOₓ generation was insignificant. Moreover, even though the scrap steel contained small amounts of chlorine and sulfur, which were neutralized by the limestone added in the furnace. The quench tower adopted by EAF also helped remove SOₓ and HCl. Thus, NOₓ, SOₓ, and HCl concentrations measured at the BH inlet and stack of the EAF were relatively low compared to the sintering process. However, SP and EAF used petroleum coke and clean natural gas as fuel, respectively.

Table 1. Flue gas compositions of the SP and EAF.

| Locations          | SP         |          | EAF       |          |
|--------------------|------------|----------|-----------|----------|
|                    | *APCD inlet| Stack    | BH inlet  | Stack    |
| Flue gas temperature (°C) | 105       | 156      | 85        | 65       |
| H₂O (%)            | 18         |          | 4.0       | 3.0      |
| O₂ (%)             | 12.7       | 13.9     | 20.4      | 20.5     |
| CO (ppm)           | 5857       | 5208     | 195       | 191      |
| CO₂ (%)            | 7.4        | 7.2      | 1.0       | 1.0      |
| NOₓ (ppm)          | 102        | 43       | < 1       | < 1      |
| SO₂ (ppm)          | 96         | 17       | < 1       | < 1      |
| HCl (ppm)          | < 0.1      | < 0.1    | < 0.1     | < 0.1    |
| NH₃ (ppm)          | *          | 2.4      | *         | *        |

*APCD: air pollution control device (DESP + FGD + SCR).
*: Not sampled.
resulting in much higher CO₂ and CO emissions from SP than that from EAF. Moreover, EAF is equipped with a CO converter at the back end of the combustion outlet to reduce CO emissions. Hence, additional control device for reducing CO emitted from SP should be considered.

Based on the concentrations of NOₓ and SO₂ measured (Table 1), the removal efficiencies of NOₓ and SO₂ achieved with the APCDs (DESP + FGD + SCR) of SP are 58% and 82%, respectively. Surprisingly, the NOₓ removal achieved with SCR is much lower than the original design value of 80% (Busca et al., 1998; Heck et al., 2016). Overall, the SCR performance depends on the operating parameters including temperature, NH₃/NOₓ ratio, and catalyst activity. If the operating temperature of SCR is not appropriate, NH₃ would escape from the reactor or even start to oxidize, resulting in lower NOₓ removal efficiency. In addition, inappropriate NH₃/NOₓ ratio also affects NOx removal efficiency. On the other hand, SO₂ reacts with water vapor present in flue gas and with ammonia injected to form sulfuric acid and ammonium sulfate, respectively. Ammonium sulfate may deposit and accumulate onto the catalyst, reducing the catalytic activity. Table 1 verifies that the concentrations of NOₓ and SO₂ measured at the baghouse inlet of EAF are lower than 1 ppm. Although SO₂ might be also formed by the reaction of sulfur from the scrap steel and oxygen from the air; SO₂ generated could be captured by the limestone which is one of the raw materials in the EAF. Consequently, with no additional desulfurization and de-NOₓ devices, the emissions of NOₓ and SO₂ from EAFs meet the emission standards.

Heck et al. (2016) investigated the correlation among NOₓ conversion, operating temperature, NH₃/NOₓ ratio, and ammonia slip of an SCR, indicating that more than 95% NOₓ conversion was obtained when operated at 360°C and [NH₃]/[NO] of 1.0, and the ammonia slip decreases with increasing operating temperature. The NOₓ removal efficiency achieved with SCR of the SP was 58%, indicating the SCR performance is not as expected and can be improved. The NH₃ slip at the SP stack was measured as 2.4 ppm, which is lower than the original design value of 10 ppm. However, the amount of NH₃ emitted is significant about 1.5 kg hr⁻¹ due to the large flow rate of flue gas (805,626 Nm³ hr⁻¹). To minimize NH₃ slip, [NH₃]/[NO] ratio is typically controlled at the range of 0.9–1 to provide satisfactory NOₓ conversions (Crolla, 2015). The operating conditions of the SCR adopted by SP should be optimized since NOₓ and NH₃ are primary precursors of PM₂.₅ and CPM.

### 3.2 Mass Concentrations of FPM, FPM₂.₅, and CPM in Stack Exhaust

As shown in Fig 3, the FPM concentrations measured at the stacks of SP and EAF are 8.2 ± 1.3 mg Nm⁻³ and 13.6 ± 2.2 mg Nm⁻³, respectively, indicating that both ESP and baghouses are in good operating conditions for controlling FPM generated in steel-making processes.

Additionally, FPM₂.₅ emitted from the SP is measured as 1.8 ± 0.5 mg Nm⁻³, which is lower than that emitted from EAF (3.2 ± 1.1 mg Nm⁻³). Emission of FPM₂.₅ depends on the type and operating conditions of the APCDs adopted. SCR is the last air pollution control device in the SP. In order to

![Fig. 3. Concentrations of FPM, FPM₂.₅, and CPM emitted from two steel-making processes.](https://example.com/fig3.png)
avoid plugging of the catalyst, additional BH is installed at downstream of the semi-dry FGD to capture PM. It is also the main reason why PM emissions from SP are lower than that of EAF.

Fig 3 indicates that CPM concentrations emitted from SP and EAF are 37.7 ± 1.3 mg Nm\(^{-3}\) and 3.4 ± 0.8 mg Nm\(^{-3}\), respectively. The CPM emitted is related to the type and operating conditions of APCDs adopted, exhaust temperature, and gas-phase pollutants including SO\(_2\), NO\(_x\), and NH\(_3\) (Yang et al., 2015; Li et al., 2019a; Yu et al., 2018; Gong et al., 2016). Li et al. (2019b) indicate that low-low temperature electrostatic precipitator (LLT-ESP) provides a good removal efficiency of CPM (> 77%). For SP, the SCR needs to be reheated to 280°C to remove NO\(_x\), resulting in a stack exhaust temperature above 156°C (Table 1), which is unfavorable for removing CPM. Moreover, SP emits much higher concentrations of SO\(_2\), NO\(_x\), and NH\(_3\) if compared with EAF (Table 1), resulting in higher CPM emissions. Therefore, the SO\(_2\), NO\(_x\), and NH\(_3\) concentrations emitted from the SP stack and the temperature of the stack exhaust should be lowered to reduce CPM emissions.

### 3.3 Emission Characteristics of PAHs from Two Steel-making Processes

#### 3.3.1 Mass concentrations of PAHs

This study sampled and analyzed gas-phase and solid-phase concentrations of 27 PAHs at the ESP inlet and stack of the SP and at the baghouse inlet and stack of the EAF, respectively, to explore PAH formation potential and emission characteristics from these two steel-making processes. The total PAH concentrations in the flue gas of the SP and EAF are shown in Table 2. The results indicate that the concentration of PAHs at ESP inlet of the SP is 226 µg Nm\(^{-3}\), with the baghouse reveals a high removal efficiency for gas-phase PAHs but is not effective in removing solid-phase PAHs achieved with the APCDs is 72% and 87%, respectively (Table 2). The overall removal efficiency achieved with the APCDs is 72%. Additionally, the results show that removal efficiencies of gas-phase and solid-phase PAHs achieved with the APCDs are 72% and 87%, respectively. The PAHs concentration measured at the baghouse inlet of the EAF is 182 µg Nm\(^{-3}\) and is reduced to 136 µg Nm\(^{-3}\) at the stack. The overall removal efficiency of PAHs achieved with baghouse (26%) is relatively low. To be specific, the removal efficiencies of gas-phase and solid-phase PAHs achieved are 23% and 81%, respectively (Table 2). The baghouse reveals a high removal efficiency for solid-phase PAHs but is not effective in removing gas-phase PAHs.

Moreover, Yang et al. (2002) studied 21 PAHs concentration emitted from steel-making industry (including SP and EAF) and the results showed that the PAHs emitted from the SP and EAF are 778–1,360 µg Nm\(^{-3}\) and 910 µg Nm\(^{-3}\), respectively. The results indicated that in addition to the 16 PAH priority species regulated by the U.S. EPA, emission of other 11 PAH species also deserves further discussion.

On the other hand, the concentrations of PAHs emitted from steel-making processes are significantly higher than those emitted from coal-fired boilers (Li et al., 2016; Lu et al., 2019). Oh et al. (2007) investigated the characteristics of air pollutants emitted from municipal solid waste incinerator, and indicate that high CO concentration during bad operating conditions is closely related with PAHs concentrations (R > 0.77). Blumenstock et al. (2000) reported the increase of PAHs during incinerator malfunctions and that PAH formation was well correlated with CO. Hence, more attention should be given to the emissions of CO from steel-making processes, which affects PAHs emissions.

### Table 2. Concentrations and removal efficiencies of gas- and solid-phase PAHs in the SP and EAF

| PAHs       | ESP inlet (µg Nm\(^{-3}\)) | Stack (µg Nm\(^{-3}\)) | Removal efficiency | Baghouse inlet (µg Nm\(^{-3}\)) | Stack (µg Nm\(^{-3}\)) | Removal efficiency |
|------------|-----------------------------|------------------------|--------------------|---------------------------------|------------------------|--------------------|
| **Mass conc. (%)** | **TEQ conc. (%)** | **Mass conc. (%)** | **TEQ conc. (%)** | **Mass conc. (%)** | **TEQ conc. (%)** | **Mass conc. (%)** | **TEQ conc. (%)** |
| Gas-phase  | 218 (96%)                   | 62 (98%)              | 72                 | 80                             | 173 (95%)            | 134 (99%)           | 23                 | 65                |
| Solid-phase| 8 (4%)                      | 1 (2%)                | 87                 | 38                             | 9 (5%)               | 2 (1%)              | 81                 | 90                |
| Total      | 226 (100%)                  | 63 (100%)             | 72                 | 76                             | 182 (100%)           | 136 (100%)          | 25                 | 68                |

conc.: concentration.
* PAHs contribution of gas and particulate PAHs to total PAHs (%).
3.3.2 Distributions of gas-phase and solid-phase PAHs

This study also explores the distributions of gas-phase and solid-phase PAHs in the flue gases of two steel-making processes. As shown in Table 2, the distribution of gas-phase PAHs increased from 96% to 98% as flue gas passed through the APCDs of SP. The gas-phase PAHs concentration is much higher than the solid-phase PAHs concentration. The results also indicate that the APCDs adopted by the SP had a higher removal efficiency for solid-phase PAHs (87%) compared with gas-phase PAHs (72%). Fig. 4(a) shows the distributions of gas-phase and solid-phase PAHs with different numbers of ring at the ESP inlet and stack of the SP, respectively. The results indicate that gas-phase PAHs at the ESP inlet are primarily composed of LMW (light-molecular-weight, 2–3 ring) PAHs (95%); PAHs with MMW (medium-molecular-weight, 4-ring) and HMW (high-molecular-weight, 5–6 ring) accounts for 4% and 1%, respectively. On the other hand, the solid-phase PAHs are primarily composed of MMW and HMW PAHs (81%), accounting for 14% and 67%, respectively. The 2-ring and 3-ring accounts only for 6% and 13%, respectively. As the flue gas passed through the APCDs, distribution of LMW PAHs decreased to 93% and the MMW PAHs increased to 6%. However, the LMW PAHs remained as the dominant species. In addition, the contribution of HMW PAHs in solid phase is decreased significantly from 67% to 20%. The 2-ring PAHs decreased from 6% to 1%, and the major increases are 3-ring and 4-ring PAHs, which increase from 13% to 22% and 14% to 58%, respectively. Li et al. (2015) studied the impact of SCR on PM$_{2.5}$ emissions from coal-fired power plants and the results showed that PAHs formed within the SCR may adsorb on the particles. The overall results showed that existing APCD cannot effectively remove gas-phase PAHs, particularly the 2–4 ring gas-phase PAHs. For solid-phase PAHs, after passing through the SCR, the percentage of 3–4 ring PAHs in solid phase increased significantly in the SP stack.

Table 2 shows the distributions of gas-phase and solid-phase PAHs at the baghouse inlet and stack of the EAF. The gas-phase and solid-phase PAHs at the baghouse inlet of EAF accounted for 95% and 5%, respectively. As the flue gas passed through the baghouse, distributions of gas-phase PAHs increased to 99%. The concentration of gas-phase at the baghouse inlet and stack are much higher than that of solid-phase PAHs. In addition, the EAF treats the flue gas with baghouse only. Thus, the baghouse had a good removal efficiency for solid-phase PAHs but did not effectively remove gas-phase PAHs. Fig. 4(b) shows the distributions of gas-phase and solid-phase PAHs with different numbers of rings at the baghouse inlet and stack of the EAF, respectively. The results show that most (89%) PAHs at the baghouse inlet are 2–3 ring while the 4–6 ring PAHs account for 11%. The solid-phase PAHs are primarily composed of 4–6 ring PAHs (91%) and the 2–3 ring PAHs contribute only 9%. As the flue gas passes through the baghouse, the 4–5 ring solid-phase

![Fig. 4. Distribution of solid and gas phase PAHs species with different numbers of aromatic rings in the flue gas of (a) SP and (b) EAF.](image-url)
PAHs are effectively captured and removed by the baghouse, and the contribution decreases from 68% to 23%. On the other hand, 6-ring PAHs is significantly increased from 23% to 70%. Wang et al. (2018) noted that some gas-phase PAHs may be transformed into solid-phase PAHs within the baghouse. Distribution of gas-phase and solid-phase PAHs is not in equilibrium because the solid-phase PAHs with 4–6 ring can be removed with particles, whereas gas-phase PAHs with 2–3 ring are of high concentrations and are difficult to remove by baghouse (Yang et al., 2002). Zhao et al. (2014) studies the fine particle emission from a coal-fired boiler equipped with a baghouse, and the result shows that the size-classified removal efficiency of PM$_2.5$ through the baghouse is of the lowest removal efficiency at 0.1 to 1 µm. In addition, the pulse blowing of the baghouse can reduce the thickness of the cake on the baghouse, hence, diminish the effectiveness of the baghouse in removing fine particles. Zhang et al. (2020) studies the size distribution of particulate PAHs in combustion source and ambient air, and the results show that 3-4 ring PAHs preferentially accumulate in coarse particles, while 5-6 ring congeners tend to be strongly adsorbed on fine particles. These studies indicate that the baghouse may increase the distribution of 6-ring PAHs.

### 3.3.3 Species and TEQ of PAHs

Fig. 5(a) shows the distribution of 27 PAHs species measured at the ESP inlet and stack of the SP.
SP, respectively. The results indicate that the PAHs measured at the ESP inlet and stack are primarily composed of 2–4 ring PAHs, and naphthalene is of the highest concentration. Specifically, Nap concentrations measured at the ESP inlet and stack are 81.7 and 19.7 µg Nm⁻³, respectively. The second and third highest contributing PAH species measured at the ESP inlet and stack are 2-MN (66.6 and 22.6 µg Nm⁻³, respectively) and PA (21.5 and 7.65 µg Nm⁻³, respectively). The results are similar to the data reported by Yang et al. (2002). After passing through APCDs, distribution of PAH species did not show significant variation because dominant 2–3 ring PAHs mainly exist in gas-phase and they are difficult to remove with existing APCDs. However, some 6-ring PAHs (DBaP, DBP, DBaIP, and DBahP) species reveal increasing trend.

Additionally, the TEFs of different PAH species are used to calculate PAH TEQ concentrations ($\text{BaP}_{\text{eq}}$) (Fig. 6(b)). The total TEQ concentrations of PAHs measured at the ESP inlet and stack exhaust of the SP are 23.0 and 5.45 µg-$\text{BaP}_{\text{eq}}$ Nm⁻³, respectively. The overall removal efficiency of the $\text{BaP}_{\text{eq}}$ achieved with the APCDs (DESP + FGD + SCR) is 76%. The primary contributing species for the $\text{BaP}_{\text{eq}}$ concentrations at the ESP inlet and stack are BcFE (21 and 5.1 µg-$\text{BaP}_{\text{eq}}$ Nm⁻³, respectively) and BaP (0.70 and 0.11 µg-$\text{BaP}_{\text{eq}}$ Nm⁻³, respectively).

Fig. 6(a) shows the distributions of 27 PAHs species at the baghouse inlet and stack of the EAF, respectively. The results are similar to those of the SP and 2–4 ring PAH species dominate in the
gas streams of EAF. The concentrations of Nap (89.7 and 68.2 µg Nm\(^{-3}\)) are the highest, followed by AcPy (21.9 and 19.8 µg Nm\(^{-3}\)), and PA (21.0 and 17.9 µg Nm\(^{-3}\)) measured at the baghouse inlet and stack, respectively. After passing through the baghouse, the concentrations of 5–6 ring PAHs are significantly reduced since most 5–6 ring PAHs are present in the solid phase and could be effectively captured by baghouse, while the removal efficiency of gas-phase PAHs is not significant. Nevertheless, the pulse blowing effect caused by the baghouse clearing reduces the removal efficiency of fine particles (Zhao et al., 2014), resulting to the an increase in the distribution of HMW PAHs (BghiP, DBalP, DBP, DBaIP, and DBahP) in the flue gas stream after baghouse.

Fig. 6(b) shows that the BaP\(_{eq}\) concentrations measured at the baghouse inlet and stack of the EAF are 14.9 and 4.74 µg-BaP\(_{eq}\) Nm\(^{-3}\), respectively. The overall removal efficiency of TEQ achieved with the baghouse is 68%. The primary contributing species for the BaP\(_{eq}\) at the baghouse inlet and stack exhaust are BcFE (13.7 and 4.41 µg-BaP\(_{eq}\) Nm\(^{-3}\), respectively) and BaP (0.21 and 0.02 µg-BaP\(_{eq}\) Nm\(^{-3}\), respectively). The contribution of BcFE is significantly higher than other species due to its high TEF value (30).

Table 2 compares the removal efficiencies of PAHs based on mass and TEQ concentrations of the SP and EAF, respectively. The results show that the removal efficiencies of solid-phase PAHs concentrations and PAH TEQ achieved with the existing APCDs (DESP + FGD + SCR) of SP are 87% and 38%, respectively. On the other hand, the removal efficiencies of solid-phase PAH concentrations and PAH TEQ achieved with the baghouse of the EAF are 81% and 90%, respectively. The comparison shows that the APCDs adopted by SP have a higher removal efficiency of solid-phase PAHs concentration than that of EAF, but the removal efficiency of solid-phase BaP\(_{eq}\) is much lower than that of EAF. Due to varying combinations of APCDs in the two steelmaking procedures, the removal efficiencies of pollutants achieved are different. Chen et al. (2007) studied the influence of SCR on the emission characteristics of PAHs and showed that adsorptive characteristics of SCR may facilitate the formation of PAHs and increase Nap, BaA, and BbF concentrations. Li et al. (2015) studied the impact of SCR on fine particle emissions from coal-fired power plants. The results showed that as flue gas passed through SCR, generation of new particles from insoluble inorganic/organic compounds may occur. This results in the increase in PM\(_{2.5}\) emissions and PAHs adherence to the new particles. On the other hand, most solid-phase PAHs are composed of 4–6 ring with high TEFs (the TEFs are 100–20,000 times higher than that of 2–3 ring PAHs); the concentrations and overall toxicity contribution of PAHs with 4–6 ring are much higher than those of 2–3 ring PAHs. Therefore, particle removal should be optimized to reduce the emission of solid-phase PAHs, especially for the fine particles containing high concentrations of 4–6 ring PAHs.

Table 2 shows the concentrations of gas-phase and solid-phase PAHs and Fig. 4 shows PAH species with different rings in the flue gas of the steel-making processes. The results show that the solid-phase 5–6 ring PAHs of proportion (76%) in EAF flue gas is higher than SP (20%). In terms of TEQ, the baghouse effectively captured particles and removed 4–6 ring PAHs which are of high toxicity. The baghouse removed the coarser particles efficiently, and improves the overall removal efficiency. However, the pulse back blowing of baghouse would result in the finer particles being emitted after the baghouse and lead to a high proportion of 5–6 ring PAHs increase. The low removal efficiency (38%) of the TEQ of solid-phase PAHs in the SP is primarily attributed to the low removal efficiency of fine particles achieved with the APCDs (i.e., DESP + FGD + SCR). Although DESP can capture the particles from SP, capture of fine particles is relatively inefficient. The FGD is a semi-dry scrubber in which alkaline slurry is sprayed during operation and generates multiple particles. The decrease of the flue gas temperature and formation of alkaline particles in the FGD process also contribute to the condensation of gas-phase PAHs. No additional particle removal device such as WESP or baghouses is installed after FGD; therefore, alkaline and fine particles containing 4–6 ring PAH are not effectively captured and lead to the formation of some highly toxic 4–6 ring PAHs.

The removal efficiency of the gas-phase TEQ achieved with the APCDs (DESP + FGD + SCR) of SP and the baghouse of EAF are 80% and 65%, respectively. Because the FGD adopted by the SP is a low-temperature process and alkaline droplets are sprayed, thereby increasing the removal efficiency of gas-phase PAHs. Within the low-temperature FGD, condensation of gas-phase PAHs is promoted. Overall, removal efficiencies of PAHs based on the mass concentration and TEQ depend on the APCD combination, indicating that the two steel-making processes investigated still have the room for improvement regarding the removal of PAHs.
3.3.4 Diagnostic ratios of PAHs

PAHs are mostly emitted as mixtures and the relative molecular concentration ratio is regarded as a characteristic of specific emission source. PAHs vary significantly in their compositions and concentrations for different high-temperature sources. Because the existing differences between the sampling technique selected and analytical procedures adopted, a great inconsistency exists among the data reported for PAHs source. Thus, exploring the impact of specific emission sources is essential for risk assessment and diagnostic ratio of PAHs may serve as an important tool for identifying the possible sources of pollution. Most diagnostic ratios involve the pairs of PAHs with the same molar mass and similar physicochemical properties. As a basis, the distributions of PAHs congeners in the EAF are analyzed for the diagnostic ratios (Yang et al., 2002; Mackay et al., 2006; Ravindra et al., 2008; Tobiszewski and Namieśnik., 2012).

Fig. 7 shows the congener ratios of PAHs emitted from EAF. The results show that the diagnostic ratios of FL/CHR, FL/(BaA + CHR), Pyr/5-MC and Pyr/(BaA + CHR) are significantly different from other stationary sources. In this study, the diagnostic ratios of FL/CHR, FL/(BaA + CHR), Pyr/5-MC and Pyr/(BaA + CHR) of EAF are 16.6, 12.1, 12.8 and 9.4, respectively. Yang et al. (2002) studied 21 PAHs emitted from EAF, and the results show that the ratios of FL/CHR, FL/(BaA + CHR), Pyr/5-MC and Pyr/(BaA + CHR) are 25.3, 10.4, 19.7 and 8.09, respectively. The diagnostic ratios obtained in this study are close and can serve as diagnostic ratios of PAHs emitted from EAF. On the other hand, this study collected coal-fired boiler diagnostic ratios for comparison. The results show that the average diagnostic ratios of FL/CHR, FL/(BaA + CHR), Pyr/5-MC and Pyr/(BaA + CHR) are 1.03, 2.06, 1.12 and 0.73, respectively, with the average standard deviation of 0.36 (Lu et al., 2019; Lu et al., 2020). The overall results indicate that the diagnostic ratios listed can be applied to help identify the different stationary sources of PAHs emissions.

Fig. 7. Congener ratios of PAHs congeners from EAF and different studies coal-fired boiler emission sources.
In addition, this study uses T-TEST to calculate the p-value to determine whether the source is different. If the p-value is less than 0.05, it means that the averages of the two populations are significantly distinct. The diagnostic ratios of FL/CHR in EAF and coal-fired boiler generate a relatively low p-value (p < 0.001). Furthermore, the p-values of FL/CHR, FL/(BaA + CHR), Pyr/5-MC and Pyr/(BaA + CHR) in EAF and coal-fired boilers are 0.0044, 0.0003, 0.0050 and 0.0003, respectively, which are far less than 0.05. The results indicate that possible source of combustion such as mobile source, coal and oil-burning, and steel-making processes can be identified by specific diagnosis ratios.

3.5 Emission Factors of Air Pollutants in Two Steel-making Processes

This study also calculates the pollutant emission factors based on the results obtained and the operating parameters. Additionally, emission factors of pollutants from steel-making process in various countries are collected and compared (Table 3). Regarding the SP, the emission factors of FPM, FPM$_{2.5}$, and CPM are 13.3, 2.95, and 61.2 g ton$^{-1}$, respectively, which are lower than those reported for SP and EAF.

| Studies          | APCDs          | PM  | PM$_{2.5}$ | NO$_x$ | SO$_x$ | PAHs         | CO             |
|------------------|----------------|-----|------------|--------|--------|--------------|----------------|
| This study (SP)  | DESP + FGD + SCR | 13.3 | 2.95 (FPM$_{2.5}$) | 93.5 | 78.8 | 0.102 (27 PAHs) | 10565          |
| Taiwan EPA (2020)| ESP + FGD + SCR | 6.28 | 181.7 | 107.7 | 337.1 |
| U.S. EPA (2009) | DESP + SCR     | 800  | 2400–2800 | 0.8–107.7 | 0.03–337.1 |
| EU (2010)        | ESP or BH      | 40.7–559.4 | 302.1–1031.2 | 219.9–973.3 | 8783–37000 |

| Studies          | APCDs          | PM  | PM$_{2.5}$ | NO$_x$ | SO$_x$ | PAHs         | CO             |
|------------------|----------------|-----|------------|--------|--------|--------------|----------------|
| This study (EAF) | CO-con.** + BH | 170.6 | 40.2 (FPM$_{2.5}$) | 16.8 | 35.9 | 1.701 (27 PAHs) | 2996          |
| Taiwan EPA (2020)| CO-con.** + CYC | 23.88 | 96.1 | 79.8 | 129.5 |
| Taiwan THS (2019)| CO-con.** + BH | 3.62–101.7 | 7.4–217 | 23–285 | 335.4 |
| Taiwan JSS (2019)| CO-con.** + BH | 9.52–483.35 | 60–326 | 35–496 | 1247–1521 |
| U.S. EPA (2009) | BH             | 19000 | 99.79 | 4.54–13.15 (CPM$_{2.5}$) | 816.5 |
| China MEE. (2017)| BH             | 86–105 |          |        | 456–1190 |
| Lai (1998)       | CYC            | BH   |          |        |        |
| Yang et al. (2002)| BH           | 1.48–18.5 (21 PAHs) | 50–4500 |
| EU (2010)        | -              | BH   | 1.13–1900 (PM$_{2.5}$) | 2070 (PM$_{10}$) | 50–4500 |

* Venturi scrubber. ** CO converter.
reported in the US, EU, and China. The emission factors of NO\(_x\) (93.5 g ton\(^{-1}\)) and SO\(_x\) (78.8 g ton\(^{-1}\)) in the SP of this study are lower than those of China and EU. The results indicate that the raw material and operating conditions of the APCDs in this SP are well controlled.

As for EAF, the results show that the emission factors of FPM, FPM\(_{2.5}\), and CPM are 170.6, 40.2, and 43.0 g ton\(^{-1}\), respectively. Overall, the emission factors of these pollutants are lower than those of the EAF in China but slightly higher than those reported in the United States (US) and European Union (EU). Therefore, the pollution control of the EAF should be further improved. Moreover, the emission factors of NO\(_x\) (16.8 g ton\(^{-1}\)) from the EAF is lower than those reported for most domestic and foreign EAFs, while that of SO\(_x\) (35.9 g ton\(^{-1}\)) which is slightly higher than that the reported in some EU countries and China.

Additionally, the emission factors of PAHs in the SP and EAF of this study are calculated as 0.102 and 1.701 g ton\(^{-1}\), respectively, which are significantly higher than those reported for a coal-fired power plant by Yin et al. (2013) in China (0.0182 g ton\(^{-1}\) of coal for 16 PAHs) and Hsu et al. (2016) in Taiwan (0.00156 g ton\(^{-1}\) of coal for 16 PAHs). Moreover, the emission factor of PAHs emitted from EAF is significantly higher than that of SP, and the difference in PAHs emission concentration is nearly 2 times. EAF is only equipped with a baghouse, so that gas-phase PAHs cannot be effectively trapped. In addition, EAF (2,263,794 Nm\(^3\) hr\(^{-1}\)) exhaust gas emissions are 2.8 times that of SP (805,626 Nm\(^3\) hr\(^{-1}\)). On the other hand, regarding the emission factors of SP, it is significantly lower than that of EU and close to the 21 PAHs emission factors studied by Yang et al. (2002). Furthermore, the emission factors of PAHs in the EAF investigated is lower than that reported by Yang et al. (2002) and Lai (1998).

CO is used as an indicator of combustion efficiency and pollution emissions. Regarding the SP and EAF investigated, the emission factors of CO are 10,565 and 2,996 g ton\(^{-1}\), respectively. The emission factor of CO from SP is significantly higher than that of EAF. Compared with EU statistics, this factor is within the EU range of 8,783–37,000 g ton\(^{-1}\), which is lower than that reported by Wang et al. (2016). However, the CO emission factor of this study are much higher than those reported by Lo et al. (2017) and Taiwan EPA (2020). The CO emission factor from EAF of this study is significantly higher than most reports but lower than the EU. There are many possible reasons for the difference in emission factor reported, such as 1. the boiler type, feed material, properties and compositions (e.g., type, size, shape, moisture content, and bulk density), 2. flue gas flow rate affects emissions (SP and EAF are 805,626 and 2,263,794 Nm\(^3\) hr\(^{-1}\), respectively), 3. operating temperature and oxygen supply, and 4. combustion method, removal performance, and operating conditions (e.g., fuel/air mixing ratio) (Wu et al., 2018; Wu et al., 2021).

**CONCLUSIONS**

In this study, representative SP and EAF are selected for simultaneous samplings and measurements of various pollutants (FPM, FPM\(_{2.5}\), CPM, SO\(_x\), NO\(_x\), PAHs and CO) at the APCD inlet and stack, respectively. Additionally, the removal efficiencies of pollutants achieved with existing APCDs and the emission characteristics/intensities of the two steel-making processes are investigated. The results can be summarized as follows:

1. The concentrations of NO\(_x\), SO\(_x\), HCl, and NH\(_3\) emitted from the SP are 43, 17, < 0.1, and 2.4 ppm, respectively; while the concentrations of all four gaseous pollutants emitted from the EAF are < 1 ppm. The NH\(_3\) concentration emitted from the SP is 2.4 ppm, and the operating conditions of SCR should be optimized to reduce NO\(_x\) and NH\(_3\) emissions and to suppress CPM formation.

2. The FPM concentrations measured at the stacks of the SP and EAF are 8.2 ± 1.3 and 13.6 ± 2.2mg Nm\(^{-3}\), respectively, while FPM\(_{2.5}\) emitted are 1.8 ± 0.5 and 3.2 ± 1.1 mg Nm\(^{-3}\), respectively.

3. The concentration of CPM emitted from the SP and EAF are 37.7 ± 1.3 and 34 ± 0.8 mg Nm\(^{-3}\), respectively. The concentration of CPM emitted from the SP is relatively high and merits further investigation for reduction.

4. The total concentration of 27 PAHs species emitted from SP is reduced to 62.6 μg Nm\(^{-3}\) with the use of APCDs (DESP + FGD + SCR), and the overall removal efficiency of PAHs achieved with the APCDs reaches 72%. On the other hand, the removal efficiency of 27 PAHs species
achieved with the baghouse of EAF is not significant (26%), indicating that baghouse did not efficiently reduce PAHs emissions.

5. As the flue gas passed through the APCDs, the gas- and solid-phase PAHs in the exhaust of SP are primarily 2–3 ring remained as the dominant species. HMW PAHs are effectively removed by APCD, but LMW PAHs are difficult to remove with existing APCD; As the EAF flue gas passes through the baghouse, the 4–5 ring gas- and solid-phase PAHs are effectively captured and removed, but solid-phase 6-ring PAHs is significantly increased trend.

6. The PAH TEQ concentrations measured at the stacks of SP and EAF are 5.45 and 4.74 µg Nm⁻³, respectively, which are much higher than those reported for coal-burning processes. The formation characteristics and control strategy for PAHs emitted from the steel-making industry require further study.

7. The diagnosis ratios of FL/CHR, FL/(BaA + CHR), Pyr/5-MC and Pyr/(BaA + CHR) are suggested as indicators for EAF.

8. The emission factors of FPM, FPM_{2.5} and CPM in the SP in this study are lower than those reported in China, EU and US, but the emission factors of FPM, FPM_{2.5}, and CPM in the EAF are slightly higher than those reported in the US and EU. However, emission factors of PAHs from SP and EAF are significantly higher than coal-fired power plant. Therefore, it is necessary to continue to pay attention to PAHs emissions from steel plants.

9. The emission factors of CO in SP and EAF are 10,565 and 2,996 g ton⁻¹, respectively. The CO emission factors are significantly higher than most studies and more effort are needed to reduce the emission from these two steel-making processes.

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DISCLAIMER

Ethics Approval and Consent to Participate
Not applicable

Consent for Publication
Not applicable

Availability of Data and Materials
All data generated or analyzed during this study are included in this published article [and its supplementary information files].

Competing Interests
Not applicable

Authors Contributions
Tang-Wei Chen developed and designed the methodology of this experiment and prepared the original draft. Jyh Cherng Chen and Zhen Shu Liu reviewed and edit the published work by those from the original research group, and management and coordination responsibility for the research activity planning and execution. Tang-Wei Chen and Kai-Hsien Chi conducted a research and investigation process, specifically performing the experiments, or data/evidence collection. Moo Been Chang supervised the project and had the oversight and leadership responsibility for the research activity planning and execution, including mentorship external to the core team. All authors read and approved the final manuscript.
SUPPLEMENTARY MATERIAL

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