Experiments on enhancing the particle charging performance in an electrostatic precipitator

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Abstract

Particle charging is an important process of particle removal in an electrostatic precipitator (ESP). A particle charge measurement system, which can adjust the flue gas temperature, was designed to study the effects of the flue gas parameters (i.e., temperature and humidity), main component of the particle, and discharge electrodes on particle charging. When the temperature changed from 300 K to 363 K, the particle charge increased with increasing temperature at the same applied electrical field strength. For the particles with a size of 0.73 µm, the average particle charge increased by 30% from 140 e to 183 e with increasing temperature from 300 K to 363 K. The particle charge was increased by 98% when the relative dielectric constant increased from 4.5 to 11.8 at −4.2 kV cm⁻¹. The increase in relative humidity significantly affected the acceleration of particle charging. For particles with sizes greater than 0.1 µm, the average charge increased by more than 50% with increasing relative humidity from 30% to 80%. Discharge electrode optimization also enhanced particle charging. When the electrode was changed from the wire electrode (d = 1 mm) to the ribbon electrode, the particle charge was increased by more than 75% for 0.7 µm particles at −4.2 kV cm⁻¹.

Keywords: Particle charging, Temperature, Humidity, Discharge electrode, Relative dielectric constant

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Introduction

The global consumption of coal by the power sector is estimated to increase over the next two decades (ExxonMobil, 2017). In China, nearly 50% of coal is consumed in coal-fired power plants (Wang et al., 2015). Many pollutants, such as NO\textsubscript{x} and SO\textsubscript{x}, are generated during the coal combustion process (Hu et al., 2017; Wang et al., 2017; Yang et al., 2018b; Yang et al., 2018c; Zheng et al., 2019). Among pollutants, particulate matter (PM) can result in potential visibility problems and adverse effects on human health (Crilley et al., 2014; Kim et al., 2015; Yang et al., 2017; Lin et al., 2018; Zheng et al., 2018; Zhou et al., 2018; Sun et al., 2019). Therefore, developing technology to control PM emission is essential.

The electrostatic precipitator (ESP) is a well-known, highly efficient device for cleaning exhaust gases from industrial processes (Wang et al., 2015; Sun et al., 2018; Yang et al., 2018a). The overall mass-based collection efficiencies of ESPs can exceed 99% (Xu et al., 2016a; Xu et al., 2016b). However, a decrease in collection efficiency was observed for different kinds of particles with the reduction in particle size to less than 0.1–1 µm (Zhuang et al., 2000). One important reason for the decrease is the low charging efficiency of fine particles (Lawless, 1996). Studying the charging mechanism of particles is useful for enhancing the charging efficiency. Different numerical techniques, theoretical models, and experiments have been proposed and developed to elucidate the charging mechanism of particles.

Numerical techniques have been adopted by researchers to analyze particle charging to model unipolar chargers. The Monte Carlo simulation was developed to capture the charging
characteristics of particles with complex shapes (Biskos et al., 2004). Apart from the Monte Carlo simulations of theoretical charging processes, numerical techniques were also developed to analyze the charging process in unipolar chargers. A 2D numerical model was developed to predict the charging process in real unipolar chargers with complex geometries, which contain the fully coupled calculation of the flow, electric potential, and ion concentration fields and the charging dynamics of particles (Chien et al., 2011; Chien and Tsai, 2013). In the aspect of theoretical models, a nonspherical particle charging model for particles of all sizes and a charge distribution model for particles sized 0.01–10 µm were developed in recent years (Gopalakrishnan et al., 2013; Domat et al., 2014). Many measurements of electric charges were developed to confirm the validity of the field charging model. Frank et al. (Frank et al., 2004) measured the electric charge on droplet particles sized 0.1–20 µm by using three different methods. The average charge were fairly consistent with the prediction of the combined charging model of Lawless (Lawless, 1996). Long (Long and Yao, 2010) analyzed different kinds of charging models in the continuum regime and compared their data with the experimental data (Hewitt, 1957), which indicated that the model established by Lawless should be the first choice for modeling the particle charging dynamics in ESPs. The effect of the dielectric constant of materials on the unipolar diffusion charging of nanoparticles was also investigated (Shin et al., 2009). Xu et al. (Xu et al., 2009) investigated the effect of power supply with different frequencies and different particle flow rates on particle charging. Park et al. (Park et al., 2011) used multichannel chargers to demonstrate that increasing ion concentration is an important
factor in increasing the average charge of the particles. Another new charger (Alonso and Huang, 2015) was also designed to enhance the nanoparticle charge.

Although many experiments were conducted, most of them mainly focused on the effects of the charge devices, particle sizes, and ion concentrations on the particle. How to adjust the actual operating conditions of the ESP and the main component to enhance the particle charge is unclear (Zheng et al., 2016). In this research, a particle charge measurement system was designed to study the effects of particle properties, flue gas parameters, and discharge electrodes on particle charge. The temperature varied from 293 K to 383 K, and the range of relative humidity was 0%–100%, which were aimed at how to improve the efficiency of ESP under real operating conditions. The effects of different discharge electrodes, including ribbon electrode, sawtooth electrode, screw electrode, and two wire electrodes (d = 1, 3 mm), on particle charge were compared. Six types of particles, namely, power plant fly ash (ash A), coal pyrolysis furnace fly ash (ash B), glass bead (particle C), silica (particle D), aluminum oxide (particle E), and calcium oxide (particle F), were investigated. This research mainly aimed to study how the main particle component, flue gas parameters, and discharge electrodes influence the charging mechanism of particles and to obtain the method of enhancing the particle charge.

**Experimental setup and methods**
A schematic of the experimental system is shown in Fig. 1. The system consisted of five parts, namely, a gas-controlled system, particle feeders, a particle charging device, a high-voltage power supply, and a particle charge measurement system.

The gas-controlled system consisted of a fan, a buffer vessel, an electric heater and a rotameter. The main stream of simulated flue gas was produced by a draught fan from the air with a maximum flow rate of 70 L min$^{-1}$ during the experiments. The buffer vessel was a cylindrical metal barrel, with a diameter of 20 cm and a height of 40 cm, connected to the middle of the pipeline linked to the feeder and the charging system. This configuration prevented excess particles from depositing in and blocking the pipes and aimed to adjust the flow rate. The relative humidity of the gas, which ranged from 0% to 100%, was measured by an electronic hygrometer (CX309; Anseny, Ltd., China). The gas temperature was controlled in the range of 293 K to 383 K by an electric heater. The temperature of these experiments were generally 313 K, apart from the experiments aimed at exploring particle charging characteristics at different temperatures. All pipes were covered with thermal insulation material to reduce undesired heat loss. In addition, the designed pipelines in were as straight and short as possible to reduce the mass loss. Then, the gas entered the charging system. The charging time varies in the range of 0.06–0.1 s by changing the gas flow rate using the rotameter. Afterward, the particle charge and concentration were measured at the outlet of the particle charging system. The measurement was taken when each experiment was adjusted to a stable condition as the experiment required. The time of each experiment was very short, usually within 2 minutes.
Six types of particles, namely, power plant fly ash (ash A), coal pyrolysis furnace fly ash (ash B), glass bead (particle C), silica (particle D), aluminum oxide (particle E), and calcium oxide (particle F), were used to study the charging characteristic of different particles. The experimental particle mass concentration and the number concentration were approximately 120 mg m$^{-3}$, 190000 cm$^{-3}$, respectively. The experimental particles of these experiments were generally power plant fly ash, apart from the experiments aimed at exploring effect of the particle main component on particle charging. The particles were injected into the simulated flue gas by an electromagnetic vibrating feeder, which was self-designed and consisted of four springs and an electromagnet system. When the electromagnet was magnetized, the springs would be extruded and the base of the spring moved downward. The particles moved forward with the vibration of the base, then entered the inlet of the pipes through the connecting tube. The feeding rate could be adjusted in accordance with the vibration frequency of the vibration controller, of which the vibration frequency range was 0-100Hz. The particle feeder was replaced by an aerosol generator (TSI 3475; TSI, USA) to study the particle charging characteristics at different temperatures. It can produce polydisperse aerosol particles with an average particle size of 0.1–8 µm, which are almost electrically neutral. The material it generated was bis (2-ethylhexyl) sebacate (DEHS) aerosol particles, which were generated by the DEHS vapor condensing on the sodium chloride crystal nucleus. In this research, the total mass and number concentration of the generated aerosol particles was approximately 120 mg m$^{-3}$, 233843 cm$^{-3}$, respectively, which were controlled by adjusting the nitrogen pressure, saturator flow valve, temperature, and reheater temperature.
The particle charging device was a wire–plate discharge configuration with a negative direct current power supply (TRC2020; Teslaman Electronics Co., Ltd., China). The wire–plate negative corona discharge device, which was made of Plexiglass (600 mm long, 200 mm wide, and 100 mm high), consisted of a wire discharge electrode and two grounded plates. The plates were cleaned after finishing several experiments to avoid the influence of precipitated aerosol. Ribbon electrode, sawtooth electrode, screw electrode, and two wire electrodes with diameters of 1 and 3 mm were used as the discharge electrodes (as shown in Fig. 1). The ribbon electrode were generally used as the discharge electrode, apart from the experiments aimed at exploring the effect of electrodes on particle charge. The discharge electrodes were 200 mm long, and the discharge gap was 50 mm. The length of the corona discharge region was 60 mm. The charging time was generally 0.1 s, while the charging time of the experiments aimed at effect of dielectric constant was 0.06s. The charging time set can minimize the deposition of particles while ensure a sufficient charging time(Long and Yao, 2010). The negative high voltage generated by the negative power supply with a range of 0–30 kV was applied to the discharge electrode.

The particle measurement system consisted of an electrostatic low-pressure impactor (ELPI⁺; Dekati, Ltd., Finland), a vacuum pump (SV25B; Ernst Leybold, Ltd., Germany), and a sampling probe. The inside of the sampling probe was technically coated to prevent electrostatic adsorption, which can minimize the loss of small particles. Using the vacuum pump, the flue gas was sampled through the sampling probe at a flow rate of 10 L min⁻¹. The particle charge and number concentration of particles of different sizes were obtained by the ELPI⁺. The measured size range
of the particles was 0.006–10 µm, and these particles were classified into 14 stages. The particle number concentration of each stage was obtained by detecting the signal of the induced current.

The particle number concentration was calculated using the fractional current resulting from the continuous deposition of charged particles on each aluminum/polycarbonate foil. As expressed in Eq. (1), the ELPI+ will automatically convert the measured current into the fractional concentration of the particles:

\[ I = PneQN \]  

where \( I \) is the fractional current, \( N \) is the fractional number concentration of particles, \( P \) is the fractional efficiency of particle charging, \( n \) is the number of elementary charges carried by the particle, \( e \) is the elementary charge (1.602 × 10\(^{-19}\) C), and \( Q \) is the sampling flow rate (10 L min\(^{-1}\)). The power plant fly ash size distribution measured by the ELPI+ is shown in Fig. 2.

After closing the charger of the ELPI+, the charge of a particle was calculated using the signal of the fractional induced current of the ELPI+. For each experiment, \( n \) is calculated using Eq. (2):

\[ n_1 = \frac{I_1}{Q_1 \times N \times e} \]  

where \( n_1 \) is the number of elementary charges carried by a particle, \( I_1 \) is the signal of the fractional induced current of the ELPI+ (A), \( Q_1 \) is the sampling flow rate (cm\(^3\) s\(^{-1}\)), \( N \) is the particle number concentration (cm\(^{-3}\)), and \( e \) is the elementary charge (1.602 × 10\(^{-19}\) C).

**Results and discussion**
Particle charging characteristics at different temperatures

Fig. 3 shows the effect of temperature on particle charge at $-4.4 \text{ kV cm}^{-1}$. The particle charge increased with increasing temperature and particle size. For the particles of 0.73 µm in size, the average particle charge was increased by 30% from $140e$ to $183e$ when the temperature increased from 300 K to 363 K. Particle charge increased with increasing temperature (Fig. 4). With high applied electrical field strength, the enhancement of particle charge with rising temperature was rapid. When the average applied electrical field strength was $-6.0\text{ kV cm}^{-1}$, the particle charge increased from $1.8 \times 10^4 e$ to $4.3 \times 10^4 e$, with increasing temperature from 300 K to 363 K. By contrast, when the average applied electrical field strength was $-6.4\text{ kV cm}^{-1}$, the particle charge increased from $1.8 \times 10^4 e$ to $6.4 \times 10^4 e$, with the temperature increasing from 300 K to 363 K, the increment was more than that at the average applied electrical field strength of $-6.0\text{ kV cm}^{-1}$. High temperature led to an increase in the mean free path of the surrounding gas molecules. A larger mean free path leads to more energy gained by an electron when it is accelerated in the applied electric field before colliding with a background molecule. The electron energy and probability of the gas molecule to be ionized increased with increasing mean free path (Hamou et al., 2013), such that the number of ions and the discharge current increased (as shown in Fig. 5). Therefore, the high temperature accelerated the particle charge at the same applied voltage.

Fig. 6 shows the effect of temperature on particle charge at maximum applied voltages. For corona discharges above 323 K, the particle charge decreased with increasing temperature. The
maximum applied voltage (Ghaleb et al., 2008; Ghaleb and Belasri, 2012), electric field strength, and ion concentration in the corona discharge region decreased with increasing temperature. Hence, the movement of ions to the surface of the particles and the collision frequency decreased, thereby resulting in a corresponding decrease in the particle charge. For 293 K to 323 K, the particle charge at 323 K was slightly higher than that at 293 K, because the decrease in the maximum operating voltage was small. However, the accelerating effect of temperature on corona discharge was obvious. Therefore, the discharge current at 323 K was slightly higher than that at 293 K. The ion concentration also increased. For particles smaller than 0.5 µm, diffusion charging was stronger than field charging (Jaworek et al., 2007). With the increase of temperature, the diffusion charging will be much stronger. Therefore, the charge of particles less than 0.5 µm at 323K was higher than that at 293K.

**Effect of the particle main component on particle charging**

In this research, six types of particles, namely, power plant fly ash (ash A), coal pyrolysis furnace fly ash (ash B), glass bead (particle C), silica (particle D), aluminum oxide (particle E), and calcium oxide (particle F), were used to study the charging characteristic of different particles. Among these particles, ashes A and B are mainly consisted of silica, aluminum oxide, iron oxide, calcium oxide, and magnesium oxide. The main components of particles D, E, and F were silica, aluminum oxide, and calcium oxide, respectively. The mass concentration of particle F, E, A, B, C, D was approximately 139, 394, 196, 91, 48, 149 mg cm$^{-3}$, respectively. The number concentration of particle F, E, A, B, C, D was approximately 233843, 210959, 182489, 215488, 208578, 155549 cm$^{-3}$, respectively. While during the experiment the particle mass and number...
concentration vary slightly around the set value. The particle charging characteristics of different main components under 0.1 s charging time and 30% relative humidity are shown in Fig. 7. The particle charge under the same applied voltage was ranked in the order of F > E > A > B > D > C. As indicated by other researchers (Chełkowski, 1980), the relative dielectric constant of particles F, E, D, and C are 11.8, 9, 4.5, and 4.3–5, respectively. In addition, X-ray fluorescence is used to detect the elemental compositions of ashes A and B. For ashes A and B, the relative dielectric constant is calculated using Eq. (3) (Rao et al., 2000):

\[ \ln \varepsilon = \sum x_i \varepsilon_i \]  

(3)

where \( x_i \) is the component i taking up the percentage of the particle, and \( \varepsilon_i \) is the relative dielectric constant of component i. The relative dielectric constant of all experimental ashes is shown in Table 1. The polarization capability of the particle with high relative dielectric constant is much stronger. Then, much more ions collided with the particle. Thus, the particle charging was enhanced. The particle charge under different applied voltages was investigated for clarification.

At the same applied voltage, particle charge increased with the dielectric constant. In addition, the particle charge at the same size was ranked in the order of F > E > A > B > D > C, indicating that the difference in dielectric constant was the main reason for the evident change in different types of particles' charge. Therefore, the particle charge increased with the dielectric constant
(Fig. 8). The particle charge was 377 e when the relative dielectric constant was 4.5 and increased to 746 e with increasing relative dielectric constant up to 11.8 at −4.2 kV cm⁻¹.

**Effect of the humidity in flue gas on particle charging**

Fig. 9 reveals the effect of humidity on particle charge when the relative humidity is 10%, 30%, 60%, and 80%. The average applied electrical field strengths are −4.2 and −5.0 kV cm⁻¹; the charging time was 0.1 s; and a ribbon electrode was used. The fly ash from a power plant was used as the experimental particle. For particles at the range of 0.1–10 μm, the particle charge increases with relative humidity under the same discharge voltage conditions. In addition, the enhancement of particle charge increased rapidly with the particle size (Fig. 9). Fig. 10 illustrates the effect of relative humidity on the charge of particle with three sizes. For the same particle, its charge increases with relative humidity. Increasing the relative humidity of the flue gas has a remarkable effect on the promotion of particle charge. For particles with sizes larger than 0.1 μm, the average charge was increased by more than 50% as the relative humidity increased from 30% to 80%. On the one hand, the ionization coefficient increased with relative humidity in the air, thereby leading to drastic ionization process of neutral molecules in corona discharge. Thus, the ion density was high, the number of negative ions that move to the low electrical field strength area of ESP is more, and the probability of collision with particle increases. Then the current increased (as shown in Fig. 11). Finally, a high relative humidity promotes particle charge. On the other hand, particle absorbed water vapor in the air in a stable high-humidity environment. The vapor condensed on the surface of the particle, thereby increasing the water content of the particle. The relative dielectric constant of water, which is 80, is higher than that of the fly ash from the...
power plant (4–12). Therefore, compared with the completely dry particle, the polarization
capability of the particle content containing more water was very strong in the electric field,
thereby allowing more electric field line to reach the surface of the particle. The collision
between particles and ions was frequent. Thus, the particle charge was enhanced.

**Optimization selection of a discharge electrode**

In this section, expect the effect of electrodes on particle charge, the effect of current density
on particle charge was studied. The corona current of different electrodes under the same applied
voltage can be transformed into average current density by calculation. The current can be
obtained by a current detecting device inside the power supply. The current density was
calculated using Eq. (4) at the grounded plates:

\[ J = \frac{I}{A} \]  

where \( J \) is the current density (mA m\(^{-2}\)), \( I \) is the current (mA), and \( A \) is the area of each grounded
plate (m\(^2\)).

**Fig. 12** reveals the effect of electrodes on particle charge when the relative humidity is 80%,
the average applied electrical field strength is \(-4.2 \text{ kV cm}^{-1}\), and the charging time is 0.1 s. In this
research, five types of discharge electrodes, including ribbon, sawtooth, screw, and wire (d=1, 3
mm), were examined to determine the performance of particle charging. The maximum particle
charge for different electrodes is ranked in the same order as the maximum current density is
ranked for different electrodes (**Fig. 12**). At the same applied voltage, the particle charge of the
ribbon electrode was larger than that of the other four electrodes. When the electrode changed
from wire electrode (d=1 mm) to ribbon electrode, the particle charge significantly rised to more than 75% for a particle size with 0.7 μm. At the same voltage, optimizing the electrode form greatly promoted the current density in the precipitator (as shown in Fig. 13) and then increased ion density and enhance particle charge. The effect of current density on particle charging is shown in Fig. 14. For a particle with the same size, the increase of the particle charge was positively correlated with the increase of current density. The ion density increased with the current density. The frequency of the collision between particle and ion increased, thereby resulting in the promotion of particle charge.

Conclusion

In this research, a particle charge measurement system was designed to study the effects of particle main component, and flue gas parameters, on particle charging. The results of this research obtain a series of methods to enhance particle charging. The following conclusions are drawn:

(1) Under the same applied electrical field strength, increasing temperature can enhance particle charge. In addition, when the applied electrical field strength is high, the promotion of particle charge with the increase of temperature will be rapid.

(2) Particle charge can be enhanced by adjusting its main component. Particle main component makes a great difference to particle charging, which mainly attributes to the difference of relative dielectric constant. Particle charge increases with relative dielectric
constant. The particle charge when the relative dielectric constant is 4.5 is 377 e and increases to 746 e as the relative dielectric constant increases to 11.8 at −4.2 kV cm\(^{-1}\).

(3) Under the same condition, high relative humidity indicates high particle charge. The average charge of a particle with size above 0.1 μm was increased by more than 50% as the relative humidity increases from 30% to 80%.

(4) Under the same applied voltage, optimizing the form of the discharge can promote particle charge. The ribbon electrode, which is most beneficial for particle charging, has the largest discharge current followed by sawtooth, screw electrode, and wire electrodes (d=1 mm or 3 mm). When the particle size is 0.7 μm, the particle charge of the ribbon electrode at −4.2 kV cm\(^{-1}\) is 70 e, which is significantly higher than that of any other electrodes.

Acknowledgments

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References:


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Figure captions

Fig. 1. Schematic of the experimental system.

Fig. 2. Power plant fly ash size distribution.

Fig. 3. Effect of temperature on particle.

Fig. 4. Effect of temperature on particle charge at different applied voltages.

Fig. 5. V–I characteristics curve in different temperatures.

Fig. 6. Effect of temperature on particle charge at maximum applied voltages.

Fig. 7. Effect of particle main component on particle charge at different applied electrical field strengths (a −4.2 kV cm⁻¹, b −5.0 kV cm⁻¹).

Fig. 8. Effect of dielectric constant on particle charge at different applied electrical field strengths.

Fig. 9. Effect of relative humidity on particle charge at different applied electrical field strengths (a −4.2 kV cm⁻¹, b −5.0 kV cm⁻¹).

Fig. 10. Effect of relative humidity on particle charge with three different sizes.

Fig. 11. V–I characteristics curve at different humidities.

Fig. 12. Effect of electrodes on particle charge.

Fig. 13. V–I characteristics curve at different electrodes.

Fig. 14. Effect of current density on particle charge with different sizes.
(a) The particle charge measurement system

(b) Geometry of discharge electrodes (a: ribbon electrode; b: sawtooth electrode; c: screw electrode; d: wire electrode [3 mm]; e: wire electrode [1 mm])

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Fig. 7 Effect of particle main component on particle charge at different applied electrical field strengths (a – 4.2 kV cm\(^{-1}\), b – 5.0 kV cm\(^{-1}\))
Fig. 8 Effect of dielectric constant on particle charge at different applied electrical field strengths
Fig. 9 Effect of relative humidity on particle charge at different applied electrical field strengths

(a −4.2 kV cm\(^{-1}\), b −5.0 kV cm\(^{-1}\))
Fig. 10 Effect of relative humidity on particle charge with three different sizes
Fig. 11 V–I characteristics curve at different humidities
Fig. 12 Effect of electrodes on particle charge
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Fig. 14 Effect of current density on particle charge with different sizes