Formation and Dispersion of Nitric Acid Vapor from Stack Flue Gas

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ABSTRACT

Extreme recovery of the thermal energy from the combustion of flue gas may bring about early gas condensation resulting in the increased formation of nitric acid vapor. The behavior of the nitric acid formed inside the stack and in the atmosphere was investigated through a computer-aided simulation in this study. Low temperatures led to high conversion rates of the nitrogen oxide to nitric acid, according to the Arrhenius relationship. Larger acid plumes could be formed with the cooled flue gas at 40°C than the present exiting gas at 115°C. The acid vapor plume of 0.1 ppm extended to 25 m wide and 200 m high. The wind, which had a seasonal local average of 3 m/s, expanded the influencing area to 170 m along the ground level. Its tail stretched 50 m longer at 40°C than at 115°C. The emission concentration of the acid vapor in the summer season was a little lower than in the winter. However, a warm atmosphere facilitated the Brownian motion of the discharged flue gas, finally leading to more vigorous dispersion.

Key words: Heat recovery, Nitric acid, Numerical simulation, Atmospheric reaction, Flue gas

1. INTRODUCTION

Currently, urban air pollution limits human activities outdoors and increases the cost of health care (Toronto Public Health, 2007). Industrial stacks are the primary emission sources that deteriorate air quality and waste the residual energy by releasing hot fluid. Therefore, the effective utilization and stringent management of energy sources are of great interest in Korea with regards to global energy issues. A combined heated and power plant attempts to take the thermal energy from the extreme heat of the exhausting flue gas (Wu et al., 2014). A low-temperature discharge can bring about less dispersion of the plume and pollutants.

Researchers have previously investigated the factors of flue gas dispersion, including the initial conditions of the exhaust gas, weather conditions, and environmental topography (Zhou et al., 2009). Table 1 briefly summarizes the emission conditions and practical models based on recent investigations. Most of the studies dealt with the NOx flow in the atmosphere, but a few of them took into consideration the nitric acid formation based on the chemical reactions of NO2. Once in the wet air, the production of nitric acid from NO2 should also be considered. Acid rain arises from the oxidation of NO2 in the troposphere that forms nitric acids, which are subsequently deposited on the earth’s surface, either as precipitation or dry deposition. Nitric acid can easily be dissolved in the precipitation and contribute to local acid rain, which may cause the corrosion of structures as well as human health issues.

The test field in this study was a combined heated and power plant for electricity and steam generation in Ilsan, Korea. The plant uses liquefied natural gas (LNG) as its primary fuel. The exhausting stack gas temperature is 115°C at 20.5 m/s and contains an average of 60 ppm NO. Prior to installing the practical heat recovery system, which reduced the flue gas temperature down to 40°C, we estimated the nitric acid concentration in the vicinity of the test area and its dispersion through numerical simulation in this study.

2. CHEMICAL CONVERSION TO NITRIC ACID

Nitric oxide and nitrogen dioxide, produced from the high temperature combustion of liquefied natural gas in a thermal power station, participate in a complex series of reactions in the atmosphere. One of the main reactions is the conversion of nitric oxide to nitrogen dioxide, as follows:

\[ 2\text{NO} + \text{O}_2 \rightarrow 2\text{NO}_2 \] (1)

This overall reaction has been widely studied and con-
firmed under various environmental conditions (Morrion et al., 1996). According to a kinetic study, the conversion of NO to NO$_2$ is inversely proportional to the absolute temperature due to the negative activation energy (Baulch et al., 1973).

Nitrogen dioxide and water vapor react in the gas phase to form nitric acid vapor and nitric oxide, as follows:

$$3\text{NO}_2 + \text{H}_2\text{O} \rightarrow \text{NO} + 2\text{HNO}_3$$

(2)

The following reactions are proposed to describe the above reaction (England and Corcoran, 1974):

$$2\text{NO}_2 + \text{H}_2\text{O} \rightarrow \text{HNO}_2 + 2\text{HNO}_3 \text{ (slow)}$$

(3)

$$2\text{HNO}_2 \rightarrow \text{H}_2\text{O} + \text{NO} + \text{NO}_2 \text{ (fast)}$$

(4)

The gas-phase reaction of nitrogen dioxide with water vapor is a homogeneous reaction that initially follows third-order kinetics, according to the following equation (5):

$$\text{R}_{\text{HNO}_3} = -k \left[\text{NO}_2\right]^2(\text{H}_2\text{O})$$

(5)

where $R_{\text{HNO}_3}$ represents the rate of production of nitric acid, and $k$ is the Arrhenius constant dependent on the activation energy and operating temperature. The activation energy of the reaction was found to be approximately $-978$ cal (England and Corcoran, 1974). The final obtained value of the reaction rate, $k$, was $5.5 \times 10^4 \text{L}^2\text{mol}^{-2}\text{sec}^{-1}$ at $25^\circ\text{C}$. The reaction of the part-per-million concentrations of nitric oxide with large excesses of both oxygen and water vapor were examined. As a result, the concentrations of the latter two compounds did not change significantly during the reactions (England and Corcoran, 1975). In addition, the decomposition of nitric acid vapor at low pressures was found to be very slow at ordinary temperatures and, therefore, it was not considered to be an influencing factor in the system (Johnston et al., 1951).

### 3. NUMERICAL CALCULATION

#### 3.1 Computational Model

In this study, we examined a combined heated and power plant for the generation of both electricity and steam in Ilsan, Korea. The site is in a highly populated residential area that includes apartments and office buildings near the plant, as shown in the map in Fig. 1(a). A 41-m-high hill is situated upstream of the stack and a subway station is located 450 m to the southeast. The model shown in Fig. 1(b) was created by Gambit Software and matches the map. The model domain was designed to be 2,000 m by 1,000 m in a plane and 300 m in height. A triangular cell was chosen for the ground face, hexahedral-based mesh was used for the cylinder shape, and tetrahedral-based mesh was adopted for the domain. We hypothesized that a total of 1.2 million gradient meshes were sufficient for the composition of the ground.

The standard $\kappa$–$\varepsilon$ turbulent model based on the eddy viscosity hypothesis was used for the pollutant dispersion (Konig and Mokhtarzadeh-Dehghan, 2002). It is a semi-empirical model composed of transport equations for the turbulence kinetic energy ($\kappa$) and its dissipation rate ($\varepsilon$). The $\kappa$–$\varepsilon$ model was derived with the assumptions that the flow is fully turbulent and the effects of the molecular viscosity are negligible. The governing equations of the fluid flow (mass, momentum, and energy equations) represent mathematical statements of the conservation laws of physics: the law of mass conservation, Newton’s second law, and the first law of thermodynamics. The Navier-Stokes equation solves the viscous stress term, which is unknown within the governing equations (Versteeg and Malalasekera, 1995). In this study, the chemical conversion of nitric oxide to nitric acid was simulated on the basis of volume. The solver, SIMPLE, suggested by Patankar and Spalding (1972), discretized the equations of momentum and continuity.
3.2 Meteorological Conditions

The annual wind rose on the test field from 2010 to 2012 is presented in Fig. 2(a). The prevailing wind direction was eastward with an average wind speed of 3 m/s. The velocity profile of the wind followed the principle of increasing with altitude, as defined by Deacon’s equation, Eq. (6) (Peterson and Hennesse, 1977),

\[ U_z = U_{ref} \cdot \left( \frac{Z}{Z_{ref}} \right)^\alpha \]

where the wind speed, \( U_{ref} \), corresponds to \( Z_{ref} \), which was 3 m/s at 10-m-high, and \( \alpha \) indicates the constant of the atmospheric stability, which was 0.2 in this study based on the weak instability of the atmospheric situation in the urban area (Irwin, 1979). The vertical temperature gradient was applied in order to estimate the plume rise according to the atmospheric conditions, as shown in the following Eq. (7),

\[ T = T_0 - 6 \times H \]

where \( T_0 \) is the ground temperature (°C) and \( H \) is the altitude (km). Ilsan has four distinctive seasons in the air temperature. Table 2 lists the average temperature of the day and night in 2012. The maximum and minimum air temperature at the field site reached 35.7°C in the summer and −17.4°C in the winter, respectively. From the present meteorological background, the gradient profiles of the air temperature and wind speed were calculated and are described in Fig. 2(b).

4. RESULTS AND DISCUSSION

In the present study, the formation and dispersion of nitric acid were investigated by numerical simula-
formation as summarized in Table 3, and discussed mainly in terms of the gas temperature, the emission concentration of NO, and the wind speed.

### 4. Formation of Nitric Acid from Flue Gas

In practice, the discharged NO\textsubscript{2} may form acid mist in a humid atmosphere. A typical gas phase reaction of NO\textsubscript{2} with the water vapor from flue gas or atmospheric moisture is homogeneous and initially follows third-order kinetics (Goyer, 1963). The intermediates of the serial reactions are dinitrogen tetroxide (N\textsubscript{2}O\textsubscript{4}) and nitrous acid (HNO\textsubscript{2}), which have very short lifetimes. The presence of oxygen does not change the initial reaction rate, but instead promotes the equilibrium of the reaction. The reaction extent is critically dependent on the temperature: the lower the temperature, the greater the production of nitric acid (England and Corcoran, 1974).

#### 4.1 Effects of the Flue Gas Temperature

The conversion of NO\textsubscript{x} to nitric acid within the stack varies with the temperature inside. Fig. 3 shows the concentration profiles of the nitric acid mist at the downflow of the stack with the flue gas temperature. The inner figures (a2, b2) indicate the formation along the center of the stack. The stack inlet concentration of the nitric oxide was assumed to be 60 ppm, which was the average value released from the test power plant. Obviously, a larger plume of acid mist was found at a flue gas temperature of 40°C rather than 115°C. The greatest amount of nitric acid was found at the stack exit, and the concentration contour for 0.1 ppm rose up to 200 m in height and 50 m in width at 40°C, while it reached about 180-m-high and 40-m-wide for a flue gas temperature of 115°C. The dilution rate

| Spring/°C | Summer/°C | Autumn/°C | Winter/°C |
|-----------|-----------|-----------|-----------|
| Day (max) | 22.9      | 35.7      | 19.9      | -9.1      |
| Night (min) | 6.6      | 26.5      | 8.9       | -17.4     |
| Average   | 12.1      | 30.9      | 13.5      | -13.5     |

Table 2. Day-night temperature variation and average of the four seasons in 2012 in Ilsan.
should be more significant for the formation of nitric acid contours in the atmosphere than the reaction rate. As can be seen in the inside figure (a2), the acid concentration approaches 3.6 ppm at the upper side of the stack. Therefore, it is possible that the acid vapor is condensed as soon as it contacts the cold air outside. Then, it can seriously impact the stack structure as well as deteriorate the visual appearance. Therefore, the use of corrosion-resistant materials is essential for the stack inner shell when the stack gas is cooled.

4.1.2 Effects of the Initial Concentration of Nitric Oxide

The initial amount of NO in the flue gas is directly related to the production of nitric acid. As presented above, nitric oxide converts to nitric dioxide in an oxygen-sufficient atmosphere, and some of the nitric dioxide then becomes nitric acid with the presence of the water moisture. Fig. 4 shows the nitric acid concentration contours with initial NO concentrations of 10 ppm, 50 ppm, 60 ppm, and 100 ppm, which are the target concentrations of the power plant by the new facility, the new national emission standard, the present emission average, and the standard for old facilities, respectively.

Increased amounts of nitric acid were formed with 100 ppm of NO, while a very slender plume body was generated with 10 ppm of NO. From Eq. (5), the production rate of HNO3 is directly proportional to the NO2 concentration in the second order, while the NO2 formation also depends on the NO concentration in the second order (Morrion et al., 1996). In particular, the NO concentration, rather than those of the oxygen and water vapor, directly determines the reaction rate for part-per-million level reactions. As observed in Fig. 4, the inner contours, indicating a high level of nitric acid close to the stack outlet, are more clearly growing with NO concentration of the flue gas. However, the outside contour of 0.1 ppm has not been expanded very much, which implies the significance of atmospheric dilution. The mixing with a large volume of air instantaneously reduces the acid concentration to lower than 0.1 ppm.

Fig. 5 summarizes the HNO3 concentration variations along the stack center with four initial concentrations. The reaction was initiated even from the stack bottom and then increased exponentially as it rose up through the stack. Upon reaching the maximum level at the top of the stack, the flue gas began to dilute with fresh air, showing a steep decrease in the acid level. At a height greater than 250 m, an obvious acid level could no longer be seen. The predictions in Fig. 4 and Fig. 5 were calculated based on the conditions of no wind and no background nitric acid. In such a case, downwash of the flue gas including nitric acid vapor was not observed. Therefore, it can be concluded that there was no direct impact of the acidic atmosphere on the ground near the emission source.

4.2 Dispersion of Acid Vapor in the Atmosphere

The vertical momentum and horizontal wind cause the discharged flue gas to rise up and to disperse into the air. The flue gas penetrates into a large body of air more quickly on windy days, leading to the efficient dilution of pollutants. The distribution pattern of the pollutants and its influencing area were estimated in the area adjacent to the test power plant.

4.2.1 Nitric Acid Concentration Distribution in the Atmosphere

The discharged flue gas is dispersed into the ami-
ent air in three dimensions: the downwind (x-axis), crosswise (y-axis), and longitudinal (z-axis) directions as described in Fig. 6. The wind flows to the east (from left to right in Fig. 6(a)) at 1 m/s. In Fig. 6(a), the acid concentrations reach their peak values at certain distances from the emission source stack. For example, the maximum concentration at an 80 m height from the ground was found at a distance of 180 m from the stack. In this study, we focused on the low level altitude in order to investigate the human effects on the ground. Despite some difference in the acid concentration near the stack with altitude, it approached cer-

Fig. 3. Vertical concentration profiles of the nitric acid mist with the flue gas temperature: (a) 40°C; (b) 115°C (supplementary figures (a2, b2) show the vertical distribution of the nitric acid concentration along the stack center-line).
tain levels over 400 m or 450 m; approximately 6 ppb for 40°C and below 4 ppb for 115°C. In other words, a thick band of higher acid levels at 40°C propagates a long distance. The altitude indicating the ground level, 1.5 m, could have an acid impact by flue gas emissions in quite a large area, particularly with 7 ppb at 40°C and 3.5 ppb at 115°C. These values are quite noticeable comparing to the field measurement which was detected at less than 1.2 ppb.

Fig. 6(b) represents the nitric acid concentration in the crosswise direction, namely, the y-axis direction, with different altitudes at x=200 m. The profiles demonstrate that the distribution from and to the rear of the stack is a Gaussian pattern, which indicates that there is not a significant influence due to wind shifting, but that there are differences according to altitude. This implies that there are only effects from natural diffusion and dilution without any direct wind effects.

In the longitudinal direction (the same as the z-axis, shown in Fig. 6(c)), the distribution trends also followed a Gaussian distribution with the height. The
Fig. 6. 3-D concentration profiles for (1) 40°C and (2) 115°C flue gas with wind speed of 1 m/s: (a) downwind-x-axis; (b) crosswise-y-axis; and (c) longitudinal-z-axis.
maximum peaks were found at 150 m and 140 m high for 40°C and 115°C, respectively. As shown in Fig. 6(a), the concentration peaks were at approximately 150 m to 140 m away from the stack. Buoyancy due to warm flue flow promotes greater lifting up of the acid plume body and, therefore, touches the peak at approximately 150 m high rather than 80 m from the stack outlet. Buoyancy also facilitates the formation reaction of nitric acid as it is floating up.

4.2.2 Effects of Wind Speed

Wind is one of the key factors causing atmospheric turbulence. The most frequently occurring wind speed in the test area was 3 m/s at the top of the stack. As seen in Fig. 7, faster winds caused flatter plumes that extended further, particularly at low temperatures. The influencing area of 0.1 ppm HNO₃ with a 3 m/s wind was an ellipsoid with 170 m length, 40 m width, and 40 m thickness at a flue gas temperature of 40°C (b1), which was 50 m longer than that at 115°C (b2). Low speed winds formed larger and higher acid plumes (a1, a2), but influenced a shorter distance (120-140 m) windward. The plume width as seen from the top view (50 m) was also wider at 40°C than at 115°C (35 m) for the wind speed of 1 m/s. Therefore, low temperature flue gas in a light wind may more significantly affect the environment in the vicinity of the emission source.

4.2.3 Seasonal Variation of the Acid Concentration

As summarized in Table 2, the test area, Ilsan, had distinctive weather conditions throughout the seasonal variation, for the air temperature in particular. Fig. 8 presents the HNO₃ concentration curves along the stack center as a function of the outside air temperature without wind: −15°C, 0°C, 15°C, and 30°C. The concentration of the stack inside up to a height of 80 m was almost overlapped due to no direct contact with the atmosphere. As was the case close to the stack...
topside, external conditions may affect the reaction even inside of the stack. The acid level dropped steeply directly after exiting the stack, probably due to dilution. The effects of the air temperature were more significant at high temperatures, as clearly indicated by the data from approximately 160-m-high. Consequently, the seasonal variation of the acid plume formation would not be very large, at least in the range of $-15^\circ$C to $30^\circ$C. The absolute value of the concentration depends on the flue gas temperature instead.

On the other hand, the concentration profiles with a wind speed of 1 m/s are displayed in Fig. 9. A typical summer day at $30^\circ$C shows little further dispersion of the acid plume. Hot weather may lead to more active Brownian motion of the gas molecules, resulting in wide scattering. However, although it predicted an obvious tendency with the air temperature, the difference between the seasons was not as significant as the plume size growth by flue gas cooling to $40^\circ$C.

### 5. CONCLUSIONS

From the point of energy savings, the recovery of the internal heat of stack flue gas using high efficiency
heat exchangers has been attempted. However, extreme recovery of thermal energy may reduce the exiting temperature, possibly resulting in low buoyancy and less dilution effects. In particular, cooled flue gas includes high moisture content and may produce higher amounts of nitric acid depending on the environmental conditions. Therefore, we estimated the quantitative formation of the acid plume and its dispersion by mathematical computations in this study. As a result of the calculations, the following evaluations could be obtained:

1. NO₂ formed from the flue gas produces a nitric acid mist with the aid of the moisture in the stack and the humid air outside. The acid plume size was larger for the cooled flue gas (40°C), whose size could be expanded to 25 m in width and 200 m in height for a 0.1-ppm HNO₃ level. It was obviously larger in winter.

2. The direct influencing distance as well as the absolute concentration of nitric acid were more significant at 40°C than at 115°C due to less Brownian motion.

3. At the distance of 800 m from the stack, the acid still presented at approximately 6 ppb for 40°C and 3.5 ppb for 115°C. The Gaussian distribution of the HNO₃ concentration was presented in the crosswise and longitudinal directions.

4. While the influencing area of the acid vapor at low wind speeds could be limited to near the stack, high-speed winds stretched the acid plume to long distances, depending on the flue gas temperature.

5. Consequently, plume size would be determined by the flue gas temperature rather than by the outside air temperature.

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