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Sulfation of Condensed Potassium Chloride by SO₂

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ABSTRACT: The interaction between alkali chloride and sulfur oxides has important implications for deposition and corrosion in combustion of biomass. In the present study, the sulfation of particulate KCl (90–125 μm) by SO₂ was studied in a fixed bed reactor in the temperature range 673–1023 K and with reactant concentrations of 500–3000 ppm SO₂, 1–20% O₂, and 4–15% H₂O. The degree of sulfation was monitored by measuring the formation of HCl. Analysis of the solid residue confirmed that the reaction proceeds according to a shrinking core model and showed the formation of an eutectic at higher temperatures. On the basis of the experimental results, a rate expression for the sulfation reaction was derived. The model compared well with literature data for sulfation of KCl and NaCl, and the results indicate that it may be applied at even higher SO₂ concentrations and temperatures than those of the present study. Simulations of sulfation of KCl particles with different size indicate that only for very small KCl particles, below 1 μm, a considerable in-flight sulfation is achievable at the short gas residence times typical of combustion systems.

INTRODUCTION

The sulfation of alkali chlorides has important implications for thermal conversion of biomass. During pyrolysis and combustion of biomass at temperatures above about 1073 K, potassium is partly released to the gas-phase,1−12 typically as potassium chloride. For agricultural residues, the KCl release can be significant,3,4,11,13,14 while woody biomass contains smaller quantities of K and Cl.8,9,13−16 Once released, gaseous KCl may be sulfated by reaction with SO₃17−22 or it may, during cooling, condense on particulates or heat transfer surfaces.18,23−26 If the particle loading in the system is low, KCl may form an aerosol by homogeneous nucleation.27

Potassium chloride has a number of adverse effects in combustion units. Once deposited on heat transfer surfaces, it may act to lower the melting temperature of the deposit, forming a sticky surface that promotes further deposition.26,28 Furthermore, its presence in the deposits may lead to severe corrosion by a mechanism where both potassium and chlorine are active in destroying the metal oxide protective layer.29 Another adverse effect of KCl is the ability to deactivate SCR catalysts;30−33 a phenomenon that limits the application of SCR for NOx control in biomass fired systems. The sulfation of condensed alkali chlorides proceeds as a gas-solid or gas-liquid reaction that may involve SO₃34 or SO₂:35

\[ 2\text{KCl}(s, l) + \text{SO}_2 + \frac{1}{2} \text{O}_2 + \text{H}_2\text{O} \rightarrow \text{K}_2\text{SO}_4(s, l) + 2\text{HCl} \]  

(1)

\[ 2\text{KCl}(s, l) + \text{SO}_3 + \text{H}_2\text{O} \rightarrow \text{K}_2\text{SO}_4(s, l) + 2\text{HCl} \]  

(2)

Here s and l denote solid and liquid species, respectively, while the remaining species are gaseous. These reactions have been reported to be slower than the homogeneous sulfation occurring in the gas-phase,17,36 leading to the assumption that the in-flight sulfation of particulate KCl is insignificant in a furnace where the residence time in suspension is of the order of a few seconds. However, for KCl condensed on deposits, the available reaction time for sulfation is orders of magnitude longer. Corrosion due to HCl release close to the metal surface is in discussion,29,37,38 but recent results indicate that the sulfation reaction serves to reduce the corrosion propensity.40

In addition to its relevance for biomass combustion, the sulfation of alkali chlorides has implications for industrial production of HCl,34 HCl emissions from municipal solid waste incinerators41,42 and kraft recovery boilers,43 as well as for corrosion of gas turbine engine components39 and cement kilns.44 Despite the importance, there are only few reported experiments on sulfation of solid alkali chlorides. Most previous studies investigated the sulfation of NaCl(s) by SO₂.34,36 In more recent work, Matsuda and co-workers45,46 compared the sulfation rates of NaCl(s), KCl(s), and CaCl₂(s). The reported experiments were typically conducted with a fixed bed of alkali chloride, varying temperature, and concentrations of SO₂, O₂, and H₂O. While these studies offer data for reaction rates and orders in reactants, results were mostly obtained for SO₂ concentrations of 3000 ppm and larger; i.e., considerably above the levels typically found in combustion systems. For this reason, rate coefficients may not extrapolate well to practical conditions.

The objective of the present work is to study the sulfation of condensed KCl by SO₂ in the temperature range 673–1023 K and lower SO₂ concentrations than in previous work.
Experiments are conducted in a fixed bed reactor with reactant levels of 500–3000 ppm SO2, 1–20% O2, and 4–15% H2O, and the degree of sulfation monitored by measuring the formation of HCl. A rate expression for the sulfation reaction is derived and results are compared to data from literature.

**Experimental Section**

The experiments were conducted in a fixed bed quartz reactor with solid KCl placed on a porous quartz plate in an isothermal zone. A schematic of the reactor is shown in Figure 1. Reactant gases (SO2, O2, and H2O in N2) were introduced separately into the reactor and mixed just above the porous plate. The reactor temperatures were measured under inert conditions with a type K thermocouple (accuracy ±3 K). The temperature in the isothermal zone of the reactor was accurate within 10 K. The product gas was collected in the bottom part of the reactor and led to the online gas analyzers. Heated lines were used to avoid cold-spots where condensation could occur. The concentrations of HCl and SO2 were measured continuously. The HCl analyzer (Servomex Servotough 2510) had an accuracy of ±3%, but the possible adsorption of SO2 onto tube surfaces contributes to the uncertainty for this species. In selected experiments, we employed a Langmuir–Hinshelwood expression to describe the dependence on the SO2 concentration:

\[
\frac{dX}{dt} = k_d(1 - X)^{2/3} \frac{Y_{SO_2} Y_{O_2} Y_{H_2O}}{1 + K_{SO_2} Y_{SO_2}^n (Y_{H_2O})^n}
\]

In the second formulation, we employ a Langmuir–Hinshelwood expression to describe the dependence on the SO2 concentration:

\[
\frac{dX}{dt} = k_d(1 - X)^{2/3} \frac{Y_{SO_2}}{1 + K_{SO_2} Y_{SO_2}^n (Y_{H_2O})^n}
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\]

Here, the rate constant \( k_d \) (s\(^{-1}\)) in Arrhenius form, describes the temperature dependence and \( F \) describes the functional dependence on the reactant mole fractions (\( Y_1 \)).

Two formulations of the rate expression were used to analyze the experimental results. The first equation is similar to the one used by Matsuda et al.:

\[
\frac{dX}{dt} = k_d(1 - X)^{2/3} \frac{Y_{SO_2}^n (Y_{O_2})^n (Y_{H_2O})^n}{1 + K_{SO_2} Y_{SO_2}^n (Y_{H_2O})^n}
\]

In the second formulation, we employ a Langmuir–Hinshelwood expression to describe the dependence on the SO2 concentration:

\[
\frac{dX}{dt} = k_d(1 - X)^{2/3} \frac{Y_{SO_2} Y_{O_2} Y_{H_2O}}{1 + K_{SO_2} Y_{SO_2}^n (Y_{H_2O})^n}
\]
Equations 6 and 7 are only valid for the specific particle size \( R_0 \) used in the experiments. To predict the sulfation rate for other particle sizes, we derive a surface area based rate constant \( k'_s \) (kg m\(^{-2}\) s\(^{-1}\)), which is assumed to be universal for particles of difference size. Substitution of eq 3 for \( X \) into eq 5 yields

\[
\frac{dm}{df} = -k'_s m_0 \left( \frac{m}{m_0} \right)^{2/3} F(Y_{SO_3}, Y_{O_2}, Y_{H_2O})
\]  

(8)

For a spherical particle, the equation above can be converted to

\[
\frac{dm}{df} = -k'_s \left( \frac{1}{3} R \rho_{KCl} \right) \frac{4}{3} \pi r^2 F(Y_{SO_3}, Y_{O_2}, Y_{H_2O})
\]  

(9)

where \( R \) (m) is the initial radius of the particle, \( \rho_{KCl} \) (kg m\(^{-3}\)) is the density of the KCl particle, and \( r \) (m) is the radius of the remaining unconverted particle. From eq 9, the surface area based rate constant \( k'_s \) can be obtained:

\[
k'_s = k'_s \left( \frac{1}{3} \right) \rho_{KCl}
\]  

(10)

Since eq 10 is assumed valid for all particle sizes, the conversion-based rate constant \( k_1 \) (s\(^{-1}\)) for a particle with an initial radius of \( R_1 \) can be calculated as

\[
k_1 = k'_s \frac{R_0}{R_1}
\]  

(11)

In the present study, the diameters of the KCl particles were in the range 90–125 \( \mu \)m. By assuming an equal mass distribution (with an interval of 1 \( \mu \)m) of the particles in this size range, we calculate a harmonic mean radius \( R_0 \) of 53.2 \( \mu \)m.

**RESULTS AND DISCUSSION**

Experiments on sulfation of solid KCl were conducted, varying temperature (673–1023 K) and reactant concentrations (500–3000 ppm SO\(_2\), 1–20% O\(_2\), 4–15% H\(_2\)O). All reported concentrations are volume based and on a wet basis. In the following, the effect of temperature and gas composition on the sulfation reaction rate is investigated. On the basis of these data, a model for the rate of sulfation is established and compared with results from the literature.

**Effect of Temperature.** For the baseline gas composition (959–965 ppm SO\(_2\), 4.8% O\(_2\), 3.9–4.6% H\(_2\)O, balance N\(_2\)), the temperature was varied in the range 673–1023 K. Figure 2 shows the measured HCl profiles for the different temperatures.

At the lowest temperatures, the amount of HCl generated is comparatively small and the reaction rate derived is more uncertain. Above 900 K, the formation of HCl increases rapidly with temperature. At 1023 K, a second peak in HCl is observed after approximately 2500 s reaction time, indicating a sudden increase in the reaction rate. We attribute this phenomenon to a phase transition from a solid to a melt, likely due to the formation of an eutectic mixture of KCl and K\(_2\)SO\(_4\). The melting is supported by the detection of an increased pressure in the reactor, caused by blocking of the pores in the quartz plate by the molten phase.

To confirm that a KCl–K\(_2\)SO\(_4\) eutectic was formed during the experiments at 1023 K, the collected residual particles were sectioned and analyzed by SEM-EDS. Typical results are presented in Figure 3, where a mixture of KCl–K\(_2\)SO\(_4\) with a lamellar structure is observed. Such a regular lamellar structure is a characteristic microstructure of an eutectic, strongly supporting the presence of a KCl–K\(_2\)SO\(_4\) eutectic under the given experimental conditions. The formation of a eutectic has been reported earlier by Fielder et al. in sulfation of NaCl(s) by SO\(_3\) above 900 K. Matsuda et al. reported eutectic formation in sulfation of CaCl\(_2\) but not for KCl or NaCl. The SEM-EDS analysis also shows that the core of the KCl particles is unreacted and that the sulfation reaction takes place on the surface. These results supports the assumption of a shrinking core model.

Figure 4 shows an Arrhenius plot of the sulfation reaction. It is evident that a change in reaction mechanism takes place between 823 and 873 K, in agreement with observations in the literature for KCl, NaCl, and CaCl\(_2\). In the present study, the activation energy for the high-temperature mechanism is determined to be 157 kJ mol\(^{-1}\), in good agreement with the values of 137 (KCl) and 144 kJ mol\(^{-1}\) (NaCl) reported by Matsuda and co-workers, but somewhat higher than the 115 kJ mol\(^{-1}\) derived for NaCl by Henrikkson and Warnqvist. Due to the small number of data points in the 623–873 K range and the limited experimental accuracy at low HCl
concentrations, we cannot reliably determine the activation energy for the low-temperature mechanism, but our results are consistent with values of 3.4−23 kJ mol\(^{-1}\) reported for KCl and NaCl.

The reason for the observed change in mechanism is still unresolved. Results from simultaneous thermal analysis (STA) for KCl indicated a phase transition of KCl around 1058 K, but no change in the crystal form of KCl was observed in the temperature range 823−923 K. Potassium sulfate exhibited a reversible change in crystal form at around 860 K, in agreement with literature reports, but the enthalpy of transformation is only about 5.8 kJ mol\(^{-1}\) and cannot in itself explain the change in activation energy of the sulfation reaction.

A plausible explanation to the observed change of mechanism is that the sulfation at low temperatures involves a condensed-phase intermediate, perhaps a potassium sulfate, with reduced thermal stability at temperatures above 900 K. Another possibility is that the low-temperature reaction involves SO\(_3\) rather than SO\(_2\), as reactant. The reaction between KCl\((s,l)\) and SO\(_3\) has been shown to proceed essentially without an activation energy. Even though quartz is not very effective as a catalyst for SO\(_2\) oxidation, its presence (wool bed, reactor surface) may facilitate formation of SO\(_3\) in quantities sufficient to explain the observed (small) conversion. A mechanism that involves surface oxidation of SO\(_3\) would also help to explain the large scatter in reported sulfation rates in the 600−900 K range (see below).

It should be noted that the sulfation products at low temperatures (e.g., 673 K) may be dominated by potassium pyrosulfate (K\(_2\)S\(_2\)O\(_7\)), which is thermodynamically favored compared to K\(_2\)SO\(_4\) under these conditions. For both SO\(_2\) and SO\(_3\) as reactants, the formation of K\(_2\)S\(_2\)O\(_7\) is exothermic. When the temperature is increased to a range of 823−873 K, decomposition of K\(_2\)S\(_2\)O\(_7\) to SO\(_3\) + K\(_2\)SO\(_4\) \((\alpha\ or\ \beta\ phase)\) becomes important. This reaction is endothermic, with calculated reaction enthalpies of 65−75 kJ mol\(^{-1}\).

**Effect of Reactant Concentrations.** A range of experiments were conducted at 923 K to determine the dependence of the sulfation rate on the concentrations of SO\(_2\), O\(_2\), and H\(_2\)O. Figure 5 shows results for variation of the SO\(_2\) concentration, maintaining the concentrations of O\(_2\) and H\(_2\)O at constant levels. A regression for the range 500−1500 ppm SO\(_2\) is consistent with a \((Y_{SO2})^{0.78}\) dependence (shown as the long-dashed line). As the SO\(_2\) concentration is increased further (3000 ppm), the reaction order in SO\(_2\) appears to decrease, possibly due to a saturation in SO\(_2\) when adsorbing on the surface. Over the range 500−3000 ppm, the best fit is obtained with \((Y_{SO2})^{0.59}\) (short-dashed line). These values can be compared with a reaction order of 0.5, obtained for the range 3000−13000 ppm SO\(_2\) by Matsuda et al. An alternative to a power fit is the Langmuir−Hinshelwood type expression (eq 7) of the type \(Y_{SO2}/(1 + KY_{SO2})\) (shown as a solid line for \(K = 708\), which may provide a more accurate extrapolation to higher concentrations of SO\(_2\).

Figure 6 shows results for variation of O\(_2\). These data, obtained over a wide range of O\(_2\) concentrations (1−20%), are consistent with a reaction order of 0.37, in good agreement with the value of 0.4, obtained for the range 2.5−15% O\(_2\) by Matsuda et al. The water vapor level was varied between 4.3 and 15% (Figure 7). The water concentration was more difficult to control and to quantify than the other reactant concentrations, but our results show only little sensitivity of the sulfation rate toward H\(_2\)O in the investigated range. A best fit to the data yields a reaction order in H\(_2\)O of 0.15, but within the uncertainty of the data, the sulfation rate can be assumed independent of [H\(_2\)O]. The results obtained by Matsuda et al. indicate that above 923 K the water vapor level (5−20%) has a larger impact on the sulfation rate, but this cannot be verified from the present work.

**Rate Expression for KCl Sulfation.** On the basis of the experimental data, we have determined the sulfation rate as a function of temperature (873−1023 K) and reactant concentrations as follows. For SO\(_2\) concentrations of 500−1500 ppm, the eq 6 format works well and we get
For a wider range of SO₂ concentrations, we recommend the eq format with the following parameters:

\[
\frac{dX}{dt} = 3.7 \times 10^6 \exp(-18900/T)(1 - X)^{2/3}(Y_{SO_2})^{0.37}
\]

\[
(Y_{O_2})^{0.37}
\]

For a wider range of SO₂ concentrations, we recommend the eq format with the following parameters:

\[
\frac{dX}{dt} = 2.7 \times 10^7 \exp(-18900/T)(1 - X)^{2/3} \frac{Y_{SO_2}}{1 + 708Y_{SO_2}} Y_{O_2}
\]

\[
(Y_{O_2})^{0.37}
\]

The Arrhenius plot in Figure 8 compares the present results with data from the work of Boonsongsup et al. ⁴⁶ for NaCl and Matsuda et al. ⁴⁶ for NaCl and KCl. The comparison is complicated by the fact that the measurements were conducted in different SO₂ concentration ranges. The data shown in the figure by Boonsongsup et al. and by Matsuda et al. were obtained at 9000 ppm SO₂, while in the present work the SO₂ range was 500–3000 ppm. We extrapolate our results to the higher SO₂ level using the Langmuir–Hinshelwood expression (high temperature) and to \((P_{SO_2})^{0.2}\) (low temperature⁴⁶), respectively, while no corrections were made for O₂ and H₂O.

The data for the sulfation rate of KCl and NaCl obtained by Matsuda et al. ⁴⁶ above 873 K are in good agreement with the results of the present work. Our rate constant for KCl is slightly higher than that reported by Matsuda et al., perhaps because their value is derived for longer reaction times, where product layer diffusion may have affected the sulfation rate. For the low temperature regime, the scatter in the data is significantly larger.

Our results are in good agreement with those reported by Boonsongsup et al. and in between data reported by Matsuda et al. for KCl and NaCl, respectively.

**Extrapolation to Higher Temperatures.** In order to evaluate how well the model extrapolates to higher temperatures, predictions were compared to data reported by Lisa et al. They studied sulfation of KCl particles (65–125 μm) at temperatures of 1173–1373 K in an entrained flow reactor. At these temperatures, a significant fraction of the KCl evaporates to the gas phase, and three mechanisms of sulfation may be active:

(A) Sulfation of particulate KCl by reaction with SO₂
(B) Sulfation of particulate KCl by reaction with SO₃
(C) Sulfation of gaseous KCl by reaction with SO₃

**Figure 8.** Arrhenius plot for the rate constant of sulfation of KCl and NaCl with SO₂. Nominal conditions for the comparison: SO₂ 9000 ppm; O₂ 5.0%; H₂O 10%; N₂ balance. Literature data is from Boonsongsup et al. ⁴⁶ for NaCl and Matsuda et al. ⁴⁶ for NaCl and KCl. The present data for KCl were obtained at conditions with 959–965 ppm SO₂, 4.8% O₂, and 3.9–4.6% H₂O. To extrapolate to the conditions of the figure, SO₂ was corrected according to the Langmuir–Hinshelwood expression (high temperature) and to \((P_{SO_2})^{0.2}\) (low temperature⁴⁶), respectively, while no corrections were made for O₂ and H₂O.
Part of the inlet SO₂ may be oxidized homogeneously to SO₃, and the presence of SO₃ increases the sulfation rate of both particulate KCl and gaseous KCl (mechanism C) considerably. Iisa et al. analyzed both the coarse particles collected in the cyclone and the fine particles captured in the filter, from these data they determined the degree of sulfation of both particulate and gaseous KCl.

The results reported by Iisa et al. indicate an accelerated sulfation rate of the particulate KCl at longer residence times in the reactor. We attribute this mostly to interference from mechanisms B and C. Potassium sulfate, formed in the gas-phase (C), may condense on the coarse KCl particles, and also reaction of KCl(s) with SO₃ (B) may have contributed to the observed degree of sulfation. The B and C sulfation mechanisms are both controlled by the rate of SO₂ formation in the gas-phase. If they were important for the observed sulfation degree of solid KCl particles, the homogeneous and heterogeneous sulfation rates should be similar. Indeed, this is in agreement with the observations of Iisa et al. for longer reactor residence times (>1 s).

Iisa et al. reported the sulfation degree of particulate KCl as a function of temperature and reactor residence time. Figure 9 compares dX/dt values derived from these data with predictions using eq 13. Unfortunately, data for short residence times were only reported for a temperature of 1373 K (0.6 s). The data shown for 1173 K were obtained for a residence time of 1.0 s, and they are more prone to be influenced by mechanisms B and C, since SO₃ has had more time to form. Considering the uncertainty, the level of agreement between the entrained flow reactor data and the model is satisfactory.

**In-Flight Sulfation of Particulate KCl.** To assess the extent of in-flight sulfation of particulate KCl, the conversion has been calculated for different particle sizes and temperatures. The calculations were based on eq 13, correcting for the effect of particle radius following eq 11:

\[
\frac{dX}{dt} = 2.7 \times 10^7 \exp(-18900/T)(1-X)^{12/3} \frac{Y_{SO_3}}{1 + 708Y_{SO_2}}
\]

(14)

Here, \( R_0 \) is 53.2 \( \mu \)m; the harmonic mean diameter of the particle size range (90–125 \( \mu \)m) used in deriving the kinetic parameters.

The results are shown in Figure 10, which are obtained based on a gas mixture with 1000 ppm SO₂ and 5% O₂ and a residence time of 1 s. It can be seen that only for very small KCl particles (0.1–1 \( \mu \)m), a considerable in-flight conversion (in the range 1–100%) can be achieved. However, KCl aerosols, from homogeneous nucleation (at low particle loadings) or from heterogeneous condensation on K₂SO₄ nuclei, are unlikely to form at temperatures above 1100 K. At temperatures below this value, the conversion of the submicrometer particles is limited to below 20%. For particles with diameters larger than 10 \( \mu \)m, the conversion is only a few percent even at very high temperatures. These results indicate that in-flight sulfation of particulate KCl in combustion systems is limited, in agreement with earlier findings.

**CONCLUSIONS**

The sulfation of condensed KCl by SO₂ was studied in the temperature range 673–1023 K and with reactant concentrations of 500–3000 ppm SO₂, 1–20% O₂, and 4–15% H₂O. Experiments were conducted in a fixed bed reactor, and the degree of sulfation monitored by measuring the formation of HCl and analysis of the solid residue. On the basis of the experimental results, the rate of the sulfation reaction can be described by the expression \( \frac{dX}{dt} = 2.7 \times 10^7 \exp(-18900/T)(1-X)^{12/3} \frac{Y_{SO_3}}{1 + 708Y_{SO_2}}(Y_{O_2})^{0.37} \) for particles in the range 90–125 \( \mu \)m. The model compares well with literature data for sulfation of KCl and NaCl, and the results indicate that it may be applied even at higher SO₂ concentrations and temperatures than those of the present study. Simulations of...
the condensed phase sulfation of KCl particles with different size indicate that a considerable in-flight sulfation may be achieved only for KCl particles below 1 μm in combustion systems.

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Notes
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■ REFERENCES

(53) Elliott, R. Eutectic solidification processing: crystalline and glassy alloys, Butterworths, 1983.