Clean process to utilize the potassium-containing phosphorous rock with simultaneous HCl and KCl production via the steam-mediated reactions

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1. INTRODUCTION

Food security is paramount to the countries with a large population. The productivity of major crops relies on the sufficient supply of nutritional elements such as phosphate (P) and potassium (K). The common practice of obtaining P and K elements derives from metallurgical processes using different sources, that is, phosphorous rock (PR) and potash. In China, the sedimentary phosphorous deposits are one of the main resources for extracting the useful P derivate to produce phosphorous fertilizers and phosphoric acid (H₃PO₄). Most phosphate fertilizers and H₃PO₄ are produced through a so-called wet process (or the dihydrate process), in which the ternary acidic system of H₂O—H₂SO₄—H₃PO₄ is used as the leaching medium and its byproduct CaSO₄·2H₂O (gypsum) is largely produced. The inherent drawback of this conventional process lies in the continuous consumption of the acid (H₂SO₄) and accumulation of the byproduct gypsum. When the sulfate ion (SO₄²⁻) migrates from sulfuric acid (H₂SO₄) to gypsum (CaSO₄·2H₂O), it creates a great challenge for recovering the sulfuric acid in the aspect of energy consumption at high temperature. In other words, the decomposition of calcium sulfate (CaSO₄) back into the base and acid is an adverse thermodynamic barrier to overcome. To make the situation even worse, this conventional wet process also shows poor performance in effective utilization of the residues (i.e., K, Ca, Al, Mg, Si, etc.), more than 60 wt % becomes waste slag. For example, the waste slag often ends up with stockpiling as phosphorous gypsum in colossus amount, posing a great hazard to the local environment, such as the great potential hazard made by stockpiling of phosphorous gypsum at the Yangzi River region. Depending on the originality, the primary mineral matrix of the phosphorus rock (PR) varies significantly. Taking the PR from southwest China as an example, the primary mineral matrix in PR often includes the fluoroapatite groups (Ca₅(PO₄)₃F), dolomite (CaMg(CO₃)₂), microcline (KAl...
Si₃O₈), quartz (SiO₂), and so on. To overcome the problem that the conventional sulfuric acid-based wet process encountered during PR processing, a new process based on chloride ions (Cl⁻) is developed in this work. The benefits of using hydrochloride acid (HCl) lie in the following reasons: (i) different reactivities between the minerals and acid, generating the acid-dissolving products and insoluble residues (phosphorous-rock slag (PS)) to separate, (ii) simultaneously generating and removing KCl from PS during the steam-mediated reaction, and (iii) relative easiness in recovering HCl through the addition of calcium chloride salts (CaCl₂) to the PS in the presence of steam at higher temperature. With regard to the reactivity during acid leaching, the acid-dissolving minerals, that is, fluoroapatite groups (Ca₅(PO₄)₃F) and

**Figure 1.** Characterization of PR: (A) XRF analysis of the main compositions in PR within ±10% uncertainties, (B) XRD, and (C) photo image of PR.

**Figure 2.** Characterization of phosphorous-rock slag (PS): (A) XRF analysis of the main compositions in PS within ±10% uncertainties, (B) XRD, (C) photo image of PS, (D) and SEM–EDS mapping for the key elements in PS.
of PS (Figure 2B). The color of PS becomes dark gray (Figure 2C). The SEM–EDS mapping (Figure 2D) of PS also indicates its complexities with existence of various elements such as K, Fe, Al, Si, and so on.

The filtrated and dried PS was then impregnated with CaCl₂ (precipitated from the leachate during the step of crystallization) at different ratios. The mixture was put in a ceramic boat reactor to react with steam at different temperatures (600–1000 °C). The experimental rig for this reaction is shown in Figure 3. The steam is quantitatively counted using a peristatic pump, and the volume fed into the reaction zone is calculated using the ideal gas law (the experimental conditions were kept at high temperature and ambient pressure, where the use of the ideal gas law is appropriate)

\[ PV = nRT \]  

where \( P \) refers to the pressure of the operational condition (in this work, the operational pressure was kept at 1 atm), \( V \) is the volume of the gas (m³), \( n \) refers to the mole of the gas (mole), \( R \) refers to the gas constant, and \( T \) refers to the operational temperature (K). Once the water was pumped into the preheating section, it was heated up and evaporated into vapor at the designated temperature. Then, the preheated steam (at a flow rate of 3 mL/min) was fed into the tube furnace to react with the mixture (PS + CaCl₂) at different temperatures. The outlet of the reactor was ducted to the heat exchanger to separate the condensed KCl (below its dew point) and HCl vapor. The concentration of Cl⁻¹ in HCl vapor was determined by mercury nitrate titration at the outlet. The concentration of K⁺ is measured by inductively coupled plasma–optical emission spectrometry (ICP-OES). After reaction, the residue in the ceramic boat reactor was taken out of the tube furnace for elementary mass balance calculations and subsequent characterization.

The total process of utilizing PR with the main mass flows (based on 100 g of PR) for simultaneous HCl and KCl regeneration is depicted in Figure 4. Apart from producing phosphorous acid (H₃PO₄), more than 90 wt% of K, Ca, and Si can be effectively high-value-converted into useful products, let alone the relative easiness in regenerating the acid (HCl, the consumption of hydrochloric acid is substantially reduced to 90 wt% less compared with the conventional leaching process, in which the acid is not regenerated). By recovering the acid through the process, it significantly reduces the amounts of external demands of the acid consumed for dissolving the PR and the production of H₃PO₄. For the convenience of study, in this work, we simply focus on the CaCl₂ + PS mixture and its

2. EXPERIMENTS

2.1. Description of the Process with HCl Recovery and KCl Generation. Due to the complex mineral matrix of PR (obtained from the Xingfa group in Hubei Province, People’s Republic of China), this work focuses on the investigation of the main elements such as K, Ca, and Si and their corresponding associated reactions. All experiments were repeated with three replica, and the experimental uncertainties were kept within ±5%. The characterization of PR used in this work is shown in Figure 1. The type of sedimentary rock is featured with (i) a relative lower level of P in Ca₅(PO₄)₃F (Ca₅(PO₄)₃F content less than 50 wt% in PR) and (ii) a relative higher level of impurities, that is, K and Ca. The X-ray fluorescence (XRF) results indicate that the percentages of three main compositions, namely, Ca₅(PO₄)₃F, CaMg(CO₃)₂, and SiO₂, are over 85 wt% in the PR (Figure 1A). The crystallite quartz, fluorapatite (Ca₅(PO₄)₃F), and dolomite (CaMg(CO₃)₂) are the main compositions in the crystallite matrix of PR (Figure 1B). The mineral often presents a dark-greenish color (Figure 1C). The scanning electron microscopy (SEM)–energy-dispersive X-ray spectroscopy (EDS) mapping (Figure S1) of PR also indicates its complexities with existence of trace elements such as S, Fe, Al, Mg, and so on. Once the PR is leached by HCl, the majority of P, Ca, and Mg elements comes into the leachate, and the characterization of the remaining acid-insoluble residue [phosphorous-rock slag (PS)] is shown in Figure 2.

The XRF results indicate that the percentages of three main compositions, namely, KAlSi₃O₈, SiO₂, and Fe₂O₃, are over 90 wt% in the PS with trace amounts of elements, that is, Mg (0.6 wt%), P (0.3 wt%), and Ca (0.2 wt%), respectively (Figure 2A). The crystallite microcline (KAlSi₃O₈), quartz (SiO₂), and pyrite (Fe₂O₃) are the main composites in the crystallite matrix of PS (Figure 2B). The color of PS becomes dark gray (Figure 2C). The SEM–EDS mapping (Figure 2D) of PS also indicates its complexities with existence of various elements such as K, Fe, Al, Si, and so on. The total process of utilizing PR with the main mass flows (based on 100 g of PR) for simultaneous HCl and KCl regeneration is depicted in Figure 4. Apart from producing phosphorous acid (H₃PO₄), more than 90 wt% of K, Ca, and Si can be effectively high-value-converted into useful products, let alone the relative easiness in regenerating the acid (HCl, the consumption of hydrochloric acid is substantially reduced to 90 wt% less compared with the conventional leaching process, in which the acid is not regenerated). By recovering the acid through the process, it significantly reduces the amounts of external demands of the acid consumed for dissolving the PR and the production of H₃PO₄. For the convenience of study, in this work, we simply focus on the CaCl₂ + PS mixture and its

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corresponding reactions in the presence of the steam at high

2.2. Characterization. The characterization of samples includes X-ray diffraction (XRD) using an Empyrean PANalytical B.V. with 50 kV, SEM, and EDS using a ZEISS Sigma300 VP, a photoelectron spectrometer (XPS), Thermo-Fisher Scientific ESCALAB 250Xi, with parameters set at Al Kα (source), 200 W (power), 500 μm (spot size), and 2.0 \times 10^{-9} \text{ mbar} (vacuum) with carbon (1s) as a calibration peak. The metal elements in solution were determined by ICP-OES (Optima 5300 DV, PerkinElmer, USA).

2.3. Thermodynamic Analysis of the Potential Reactions. In this work, the HSC chemistry 6 software (Outokumpu Technology) was used to perform a preliminary thermodynamic analysis of the potential reactions. The chemical potentials of these reactions were estimated at different temperatures at ambient pressure. To narrow down the scope of reaction selection, this work focuses on those existing mineral compositions in the PS matrix (Figure 2) and the main products after reaction (in both the solid residue and outlet vapor such as KCl and HCl). Therefore, the main possible reactions during HCl and KCl generation are illustrated below

\[
\begin{align*}
\text{CaCl}_2 + 2\text{KAlSi}_2\text{O}_8 & \rightarrow 2\text{KCl} + \text{CaAl}_2\text{SiO}_6 + 5\text{SiO}_2 & (2) \\
3\text{CaCl}_2 + 3\text{H}_2\text{O} + 2\text{SiO}_2 & \rightarrow \text{Ca}_3\text{Si}_2\text{O}_7 + 6\text{HCl} & (3) \\
4\text{CaCl}_2 + 3\text{H}_2\text{O} + 2\text{KAlSi}_3\text{O}_9 & \rightarrow \text{Ca}_3\text{Si}_2\text{O}_7 + 2\text{KCl} + 6\text{HCl} + \text{CaAl}_2\text{SiO}_6 + 3\text{SiO}_2 & (4)
\end{align*}
\]

Among these reactions, reaction 2 refers to the roasting the mixture (PS + CaCl₂) in the absence of steam at high temperature, while reactions 3 and 4 refer to the reactions between the PS + CaCl₂ mixture and steam. The corresponding Gibbs free energies of these reactions were plotted and are shown in Figure 5. Among these three different reactions 2–4 at the investigated temperature range, the chemical potentials of reactions 3 and 4 fall into the spontaneous zone. This indicates that the absence of steam is thermodynamically unfavorable to the formation of KCl and HCl at high temperature. With regard to the temperature range, the lowest temperature for reaction 4 to occur is around 655 °C, while it is around 964 °C for reaction 3. This indicates that reaction 4 might predominately occur at the investigated experimental conditions, while reaction 3 will step in once the system temperature reaches 964 °C. In addition, because of the feature of the experimental setup, it offers a transport advantage of continuous removal of the generated products, that is, HCl and KCl, out of the system by the influx of the steam, which will make reaction balance further skewing toward the generation of HCl and KCl (reactions 3 and 4). Therefore, from the above preliminary thermodynamic analysis, it is found that the introduction of steam to (PS + CaCl₂) plays a critical role during the formation of HCl and KCl. From the values of entropy and enthalpy of reactions 3 and 4, the Gibbs free energy is calculated as follows

\[
\Delta G = \Delta H - T\Delta S
\]

where \(G\) refers to the Gibbs free energy (kJ mol\(^{-1}\)), \(H\) refers to the enthalpy (kJ mol\(^{-1}\)), and \(S\) refers to the entropy (kJ mol\(^{-1}\)). As reaction approaches equilibrium, the equilibrium constant is expressed as follows
\[ \ln K = -\frac{\Delta G}{RT} \]  

\[ K = K_0 e^{E_a/RT} \]  

where \( K \) refers to the reaction constant, \( K_0 \) refers to the reaction constant under standard conditions, \( R \) refers to the gas constant (8.314 J mol\(^{-1}\) K\(^{-1}\)), and \( E_a \) refers to the activation energy (kJ mol\(^{-1}\)). The calculation of activation energies for reactions 3 and 4 is shown in Figure S2, with 150 (kJ mol\(^{-1}\)) for B and 208 (kJ mol\(^{-1}\)), respectively. These values generally fall into the range of activation energies of chemical reactions such as MnCl\(_2\) + steam reactions.  

### 2.4. Kinetic Study of the Process

The reaction was carried out in a boat reactor heated using the tube furnace. By carefully controlling the amount of water using the peristatic pump, one can quantitatively estimate the amount of steam at the designated operational temperature. The reaction zone was maintained at different temperatures from 700 to 900 °C with the duration from 10 to 150 min. The obtained solid residue in the ceramic boat was grinded into powder, followed by mixed acid [hydrofluoric acid (HF) and nitric acid (HNO\(_3\))] by 1:1 volume ratio leaching at a liquid-to-solid ratio of 10:1 for 30 min. The resultant slurry was finally filtered and oven-dried (105 °C). The K\(^+\) content in the filtrate was measured by ICP-OES. The corresponding conversion (\( x \)) of KCl is calculated as follows

\[ x = \frac{m_1\alpha - m_2\beta}{m_1\alpha} \times 100 \]  

where \( m_1 \) is the mass in the reactants, \( \alpha \) is the K content in the PS (wt %), \( m_2 \) is the mass in the solid residue, and \( \beta \) is the K content in the solid residue (wt %).  

### 2.5. Process Optimization Using the Supervised Machine Learning Approach

The preliminary experiment indicates that \( X_t \), reaction temperature; \( X_d \), reaction duration; and \( X_i \), impregnate ratio (PS over CaCl\(_2\)) were critical during reaction (KCl conversion). Therefore, these three operational parameters were investigated to find out quantitative correlations with conversion (based on KCl conversion). In this work, 17 experimental conditions were used as training data set (Table S1). For neuron network construction, the activation functions in the neuron network architecture are one of the most important ones, \( 1,2,20 \) which are used to transform input (using the summed weighted approach) into the activation of the node. In this work, the rectified linear activation function (ReLU) was used in the Tensorflow during supervised machine learning (SML) process to minimize the errors caused by the vanishing gradient.\(^{11} \) The outcome of SML for the optimization are shown in Figure 6. With multiple objective functions by maximizing KCl conversion, the optimal conditions were achieved as \( X_t\sim\, 850 \, ^\circ C, X_d\sim\, 40 \, min, \) and \( X_i\sim\, 2.5 \), respectively. The subsequent validation experiment conducted at this achieved optimal condition (\( X_t\sim\, 850 \, ^\circ C, X_d\sim\, 40 \, min, X_i\sim\, 2.5 \)) shows \( \pm 10\% \) uncertainties, indicating a relatively good prediction from this SML algorithm.  

### 2.6. Molecular Simulation Study

The total energy is calculated and expressed by assuming all ground-state properties to be functionals of the charge density\(^{23} \)

\[ E_1[\rho] = T[\rho] + U[\rho] + E_{XC}[\rho] \]  

where \( E_1[\rho] \) is the total energy, \( T[\rho] \) is the kinetic energy (non-interacting particles with density \( \rho \)), \( U[\rho] \) refers to the classical electrostatic energy contributed from the Coulombic interactions, and \( E_{XC}[\rho] \) refers to the exchange and correlation energies that contribute to the total energies. For the density function theory (DFT) calculation, the semi-empirical dispersion–correction schemes were used. The simulated annealing is deployed to locate a good approximation with the global minimum of a given function in a large search space.\(^{24} \) The Metropolis Monte Carlo method is deployed to estimate the total energy of water steam adsorption on the model composites at ambient pressure and 1097 K. In this method, the sorbate structure is treated as rigid. Only rigid body translations and reorientations are incorporated during calculations.  

### 3. RESULTS AND DISCUSSION

### 3.1. Kinetic Study

The kinetic profiles of reaction with and without steam at 800 °C are shown in Figure 7A. Obviously, the reaction in the absence of steam yields a poor performance in KCl conversion at the exit of the reactor. This agrees with the preliminary thermodynamic analysis of reaction 2. With the introduced steam, the reactions (reactions 3 and 4) toward the formation of products, either HCl or KCl or both, occur as a certain temperature threshold is reached. The kinetic profile of reaction with the steam and the morphology of solid residues and formed KCl are depicted in Figure 7B. Starting from PS, the color of the solid residue gradually changes from dark to gray as the reaction (eq 4) proceeds. In
addition, the KCl carried by the steam flux presents a very similar appearance with an increase of duration. More than 75% of K⁺ ions are converted within the first 10 min, indicating that the reaction at 800 °C is kinetically favorable. Once the temperature reaches 800 °C, the chemical potential for reaction (eq 4) becomes favorable to the direction toward the formation of KCl and HCl. Besides, the continuous removal of the products, that is, KCl and HCl, out of the reaction system will enhance the skewing reaction direction toward the formation of KCl and HCl. Therefore, both thermodynamic evidence and kinetic evidence suggest the feasibility and practicality of this proposed model for simultaneous HCl and KCl production in the presence of steam.

In this work, the shrinking core model (SCM) is used to estimate the conversion (on the KCl basis). The reaction time (t) and the KCl conversion (x) can be expressed in the following form, which includes all the controlling steps

$$t = \frac{C_{SO}}{3k_r C_{A0}} x + \frac{C_{SO}}{3k_c C_{A0}} [1 - (1 - x)^{1/3}]$$

$$+ \frac{C_{SO}^2}{6D_r C_{A0}} [1 - 3(1 - x)^{2/3} + 2(1 - x)]$$

(10)

where t is the reaction time (min), C_{SO} is the initial solid concentration (mol/L), C_{A0} is the initial concentration (mol/L), k_r is the gaseous diffusion constant factor (m²/s), k_c is the surface reaction constant factor (m²/s), and D_r is the effective diffusivity (this refers to diffusion of vapor KCl in steam at a higher temperature, m/s). The modified correlation between the conversion and kinetic constant factor for the reaction is depicted as follows

$$x = \left[1 - \frac{k_c}{d_{0.6}t}\right]^{1/3}$$

(11)

$$k_c = k_{c0} e^{E_r/RT}$$

(12)

$$k_s = \frac{\rho_k k_s^{0.4}}{M^{2/6}}$$

(13)

where k_s is the modified reaction rate constant (m^{-0.6}/s), d is the average diameter of the particle (in this work, the averaged particle size is around 23 μm and the particle size distribution of PS + CaCl₂ is shown in Figure S3), E_r is the activation energy (kJ/mol), R is the ideal gas constant (8.31 J/mol/K), T is the temperature of reaction (K), ρ_k is the density of the PS particle (kg/m³), M is the molecular mass of the reagent (kg/mol), and r is the average particle radius (m) with the following correlation

$$r = \frac{\rho_k k_s^{0.4}}{M^{2/6}}$$

(14)

$$k_s = \frac{3}{\rho_k S}$$

(15)

where S refers to the surface area of the reagent (PS + CaCl₂).

In this work, all particles in the ceramic boat reactor are assumed to be around 1.5 m²/g. The conversion expression can be further reduced to a simpler expression if the overall rate is controlled by the surface reaction as follows

$$\frac{t}{\tau} = [1 - (1 - x)^{1/3}]$$

(16)

$$\tau = \frac{C_{SO} r}{k_s C_{A0}}$$

(17)

Then its derivative can be cast into the following

$$\frac{dx}{dt} = \frac{3(1 - x)^{2/3}}{t}$$

(18)

The conversion expression can be simplified into the following if the reaction is controlled by the diffusivity (transport control)

$$\frac{t}{\tau} = [1 - 3(1 - x)^{2/3} + 2(1 - x)]$$

(19)

$$\tau = \frac{C_{SO} r^2}{6D_r C_{A0}}$$

(20)

and its derivative can be achieved as

$$\frac{dx}{dt} = \frac{1}{2} \frac{(1 - x)^{1/3}}{\tau (1 - x)^{1/3}}$$

(21)

The results of kinetics at three different temperatures (from 700 to 900 °C) are shown in Figure 8. Among different controlling steps (diffusion control, r² = 88%; reaction control, r² = 86%; and diffusion + reaction control, r² = 99%), the best fit curve of SCM (judging from the r² square value) is achieved by considering all controlling steps (diffusion and chemical reaction). This indicates that the intrinsic kinetics and
transport are both influential at the investigated experimental conditions. The corresponding activation energy ($E_a$) is around 14.63 (kJ/mol). In general, the magnitude of the activation energy for chemical reactions will be around 40−200 kJ/mol for roasting reactions between CaCl$_2$ and potassium sources such as K-feldspar in the absence of steam, $^{27,28}$ which generally agrees with the values calculated from theoretical thermodynamic analysis for the activation energies of the proposed potential reactions 3 and 4 (reaction 3, $E_a = 150$ kJ/mol; reaction 4, $E_a = 208$ kJ/mol). However, according to the definition of the SCM, this type of model is used to describe the situation in which solid particles are consumed by either dissolution or reaction; as a result, the amount of material being consumed shrinks. Therefore, the activation energy obtained from SCM simply based on KCl conversion might not be able to completely represent the intrinsic kinetics of the reactions (in reactions 3 and 4, respectively). This indicates that the use of SCM might underestimate the activation energy of the reactions during the contact of the mixture (K-feldspar + CaCl$_2$) with steam at the investigated temperature range. The

**Figure 8.** Kinetics of KCl conversion using SCM at different temperatures: (A) curve fitting using the diffusion control model, (B) curve fitting using the chemical control model, (C) curve fitting using the diffusion + chemical control model, and (D) activation energy estimation, where $E_a$ refers to the activation energy and $A$ refers to the pre-factor.
The merit of using SCM lies in the quantitative identification of controlling dominant steps (diffusion, chemical reaction, or the combination) during reactions. For process intensification, some experimental conditions such as the relative smaller particle size of PS + CaCl$_2$ (Figure S3) and the involvement of high-temperature steam (continuously removing the produced products, i.e., HCl and KCl) might all contribute to further reduction of the value of the actual apparent activation energy of the reaction.

The improvement of these factors, that is, reducing particle size, increasing flows, and so on, will be helpful to the process intensification during the step of process scale-up.

3.2. Characterization of Solid Residues at Different Temperatures. Although various parameters affect the reaction, the reaction temperature is one of the most critical parameters that influence the reaction. The characterization of the solid residues at different temperatures is shown in Figure 9. With an increase of reaction temperature, no substantial morphological differences are observed among all residues after reaction (Figure 9A). The XPS of Ca 2p$_{1/2}$ and Ca 2p$_{3/2}$, O 1s, and Si 2p is shown in Figure 9B–D, respectively. It is found that the binding energy of Ca 2p$_{1/2}$ and Ca 2p$_{3/2}$ (especially Ca 2p$_{3/2}$) of the solid residue tends to shift toward relatively lower binding energies with an increase of reaction temperature. This is associated with the broken ionic bond between Ca$^{2+}$ and Cl$^-$ (CaCl$_2$) and the formation of calcium aluminum silicate (CaAl$_2$SiO$_6$). The binding energies of both O 1s and Si 2p tend to shift to a relatively higher energy with an increase of reaction temperature. The change from microcline (KAlSi$_3$O$_8$) to calcium aluminum silicate (CaAl$_2$SiO$_6$) and the formation of the Si-contained composites (in the form of Si–O connection) are both associated with the rearrangement of bindings between the oxygen atom and other atoms (i.e., Al, Ca) in the mineral matrix.

A similar shifting pattern from Si 2p is observed, suggesting that more lattice silica atoms are found on the surface of the solid residue with an increase of temperature. The XRD spectra are shown in Figure 9E. The combined sharp and small peaks in each sample indicate the complexity of crystallite minerals in the solid residue. With an increase of temperature, the intensity of the peaks from quartz becomes smaller. This indicates that the original crystallite structure of quartz in PS might be consumed because of reactions (i.e., reactions 3 and 4). The characteristic peaks of KAlSi$_3$O$_8$ experience a decrease, indicating the consumption of microcline as reaction proceeds. This orchestrates with the increased

![Figure 9. Characterization of solid residues at different temperatures: (A) SEM morphology, (B) XPS for Ca$^{2+}$, (C) XPS for O 1s, (D) XPS for Si 2p, and (E) XRD. All other parameters were kept at the optimal condition (duration = 30 min, impregnate ratio = 2).](https://doi.org/10.1021/acsomega.2c02362)
The intensity of the peaks for calcium aluminum silicate (CaAl$_2$SiO$_6$) with an increase of temperature. Other impurities such as pyrite persistently exist in the solid residue with different intensities.

To investigate the detailed binding energies of the formed oxidative minerals, the broad O 1s peak is split into two parts; the binding energy centers around 529 eV are associated with Ca−O, and the binding energy centers around 531 eV are associated with Al−O. The main bands by deconvolution are shown in Figure 10. With an increase of reaction temperature, the binding energy of Al−O tends to gradually increase, while Ca−O experiences a decrease from 600 to 900 °C, indicating the gradual consumption of CaCl$_2$ and the formation of calcium aluminum silicate (CaAl$_2$SiO$_6$) in the solid residue after reaction. As the reaction temperature reaches 1000 °C, the portion between 531 eV (Al−O) and 529 eV (Ca−O) experiences an appreciable transitional change. A relative increased portion of 529 eV (Ca−O) up to 24% and a decrease to 76% of 531 eV (Al−O) are observed. One of the reasons might be due to the start of other side reactions that lead to the formation of the new calcium and aluminum oxides. According to the preliminary thermodynamic analysis, reaction 3 will start to step into the spontaneous zone once the reaction temperature is over 946 °C (Figure 5), where new products such as rankinite (Ca$_3$Si$_2$O$_7$) start to form and the corresponding O 1s binding energy will change accordingly (Figure 11).

The XPS spectra of Si were also recorded to ascertain the compositions associated with silica in the solid residues. Although spectra displayed the broad Si band, they also revealed the presence of valence around the Si atom arising from bonding with oxygen. Through the deconvolution, two
bands at 102.5 eV (Si−O) and 103.4 eV (O−Si−O) are achieved, which correspond to Si 2p$_{1/2}$ and Si 2p$_{3/2}$ respectively. With an increase of temperature over 600 °C, the majority of the valence around the Si atom is in the form of Si 2p$_{1/2}$ (Si−O) with some valence around Si existing in the form of O−Si−O and the substoichiometric SiOx. This also agrees with O 1s spectra; that is, the binding energy of the oxygen atom for O−Si−O locating at 532.5 eV is not appreciably observed. Therefore, the valence that the Si atom binds with other atoms is the most through Si−O. In other words, the Si atom exists mostly in the complex composite’s clusters (i.e., CaAl$_2$SiO$_6$, Ca$_3$Si$_2$O$_6$, etc.) rather than in the pure quartz (SiO$_2$) crystal structure which agrees reasonably well with the postulated reactions (reactions B and C). This result also agrees with the XRD spectra (Figure 9E) analysis that the existence of Si 2p signals will contribute the postulated reactions (reactions B and C). This result also agrees with the XRD spectra (Figure 9E) analysis that the intensity of quartz in the solid residue at a higher temperature does not appreciably increase.

### 3.3. Preliminary Reaction Mechanism Discussion.

As steam plays a critical role during reaction, it is necessary to investigate the interactions between the steam molecule and the key minerals in the PS + CaCl$_2$ matrix. Since the adsorption is the first step prior to the reaction, the total energies of the adsorption between the steam molecule and the microcline, silica, and CaCl$_2$ composites might shed some insightful light on the reaction mechanism. As mineralogy of PS is quite complicated, we only focus on the main mineral composites (shown from PS characterization data in Figure 2) such as microcline (the crystal 004 plane was chosen for adsorption using the lattice parameters $a = 24.97$, $b = 9.31$, $c = 8.57$ Å and angles $\alpha = 90$, $\beta = 81.1$, $\gamma = 90$), quartz (the crystal −100 plane was chosen for adsorption using the lattice parameters $a = 4.913$, $b = 4.913$, $c = 5.4052$ Å and angles: $\alpha = 90$, $\beta = 90$, $\gamma = 120$), and CaCl$_2$ (the crystal 010 plane was chosen for adsorption using the lattice parameters $a = 15.25$, $b = 18.65$, $c = 3.64$ Å and angles: $\alpha = 90$, $\beta = 90$, $\gamma = 90$).

To obtain some detailed information about the adsorption of water on the model compounds, the density functional theory (DFT) approach is employed and the results of water adsorption on quartz (SiO$_2$) and microcline (KAlSi$_3$O$_8$) are summarized in Figure 12. For water adsorption on quartz (SiO$_2$) and microcline (KAlSi$_3$O$_8$), the adsorption energy: $E_{\text{ads}} = -0.13$ eV. The total energy of water adsorbed on different composites using the Monte Carlo approach is shown in Figure 13. At the simulated condition ($T = 1073$ K and $P = 1$ atm), the energy barrier for steam molecules to approach the surface of microcline is much

![Figure 12. Molecular simulation for water adsorption on microcline: (a1) crystal structure of quartz, (a2) water adsorption configuration on quartz with the oxygen bond distance, (a3) water adsorption configuration on quartz with the hydrogen bond distance, (b1) crystallite structure of microcline, (b2) water adsorption configuration on microcline with the hydrogen bond distance, and (b3) water adsorption configuration on microcline with a larger bond distance.](image-url)
higher than that of CaCl$_2$ and SiO$_2$. The energy barrier of SiO$_2$ is the lowest, indicating that it is more favorable for the steam molecule to adsorb on the surface of SiO$_2$. In addition, the preliminary experiment and thermodynamic analysis also show that the reaction between the CaCl$_2$ and H$_2$O cannot proceed if it is in the absence of SiO$_2$, even when the temperature is over 1000 °C ($\Delta G = 6$ kcal/mol). With existence of SiO$_2$ (one of the acid-insoluble residues in the PS residue) and the presence of steam, CaCl$_2$ becomes reactive and the formation of HCl and KCl begins to proceed at the experimental condition. Therefore, apart from introducing the steam, the existence of SiO$_2$ also significantly facilitates changing PS into KCl and HCl, respectively.

With all the above-mentioned information, the reaction route might be postulated as follows: (i) starting from the adsorption of steam molecules on the surface of SiO$_2$ with defects, followed by the spill over the polarized H—OH water molecule toward the surface of CaCl$_2$; (ii) the energized water will participate in breakdown of the crystallite structure of CaCl$_2$ (also indicated from Ca 2p$_{3/2}$ shifting to a lower binding energy shown in Figure 9B), leaving the chloride ions (Cl$^-\text{I}$) out of the CaCl$_2$ matrix, which results in the formation of volatile HCl; iii) replacing the K$^+$ with Ca$^{2+}$ cations in the microcline matrix with the assistance of the HCl at high temperature (as indicated from Figure 12_b1, the crystallite structure of microcline, K is much more loosely coalited with the Si—Al—O complex structure), resulting in the formation of KCl; and (iv) finally, the formed Ca—Si—Al—O composite will rearrange into the composite with the minimum chemical potential as it cools down (the products suggested by reactions 3 and 4 from thermodynamic analysis). The schematic diagram of the possible reaction pathway is summarized in Figure 14. Obviously, this postulated route is a quite simplified reaction pathway; it is believed that there will be more possible side reactions in the real scenario.

4. CONCLUSIONS

A clean process based on the steam-mediated reactions for the simultaneous HCl and KCl production using potassium (K)-containing phosphorous-rock slag (PS) as a precursor is proposed. The generation of both liquid products (H$_2$PO$_4$ and CaCl$_2$) and solid residues (PS rich in elements, i.e., K, Al, Si, etc.) in the form of microcline-KAlSi$_3$O$_8$ and quartz-SiO$_2$ was achieved. The optimal conditions were achieved (X$_1$ -850 °C, X$_2$ -40 min, X$_3$ -2.5) using the SML approach. Thermodynamic analysis and molecular simulation indicate that the presence of both steam and SiO$_2$ in the (PS + CaCl$_2$) matrix plays critical roles during the formation of HCl and KCl at high temperature. The SCM suggests that both the intrinsic kinetics and transport (diffusion) are influential at the investigated conditions, and the corresponding activation energy of $E_a = 14.63$ kJ/mol is obtained. The potential reaction pathway based upon experimental characterization and theoretical simulation is also postulated.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsomega.2c02362.

Characterization of PR using SEM—EDS elementary mapping, activation energy estimation, particle size distribution of the PS + CaCl$_2$ mixture, and training data set for optimization using the SML approach (PDF)

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Notes
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