New Novel Reactive Distillation—Pervaporation Coupled Process for Ethyl Acetate Production with Water Removal from Reboiler and Acetic Acid Recycle

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ABSTRACT: Ethyl acetate (EtAc) is an essential raw material widely used in the chemical industry. In this paper, a new reactive distillation (RD)—pervaporation (PV) coupled process for ethyl acetate (EtAc) production is proposed, in which the PV membrane is located in the bottom stream in order to selectively remove the water from the reboiler and recycle the acetic acid (HAc) into the feed. The main operating conditions of the RD process are discussed by experimental and simulation methods. The RD–PV coupled process was established by coupling the PVA/ceramic composite membrane with the RD reboiler. The effects of PV operating temperature and HAc/ethanol molar ratio on the RD performance are thoroughly discussed. It was found that the PVA/ceramic composite membrane exhibited good PV performance and stability with a total flux of 600 g·m⁻²·h⁻¹ and a separation factor of 14 in 90 wt % HAc/water solution at 70 °C. Owing to the water removal and HAc recycle from the reboiler via PV, both the ethanol conversion and EtAc purity are remarkably improved, from 82.4 to 85.6 wt % and from 81.3 to 84.8%, respectively. This new RD–PV coupled process can be expected to be an effective and energy-saving way for ethyl acetate production.

1. INTRODUCTION

Ethyl acetate (EtAc) is an essential raw material widely used in the chemical industry including in paints, coatings, perfumes, and adhesive agents.⁹–¹³ EtAc is mainly produced industrially via a classical esterification process in which acetic acid (HAc) and ethanol (EtOH) react in the presence of a strongly acidic catalyst in a reactor.⁴ In 2001, the international market demand for EtAc was 1.0 million tons, and the demand reached 2.5 million tons in 2008. In working toward the elimination of toxic solvents, the consumption of EtAc is expected to grow at a rapid pace because of its low health and environmental effects.⁵

For reversible reactions such as esterification and etherification, the equilibrium limitation is a key issue, motivating the use of reactive distillation (RD) to achieve high conversion and selectivity by shifting the chemical equilibrium boundaries, reducing byproduct, and overcoming azeotropic limitations. In addition, the RD process is motivated by benefits such as a reduction in the capital and operational costs due to combining the reaction and separation in a single unit.⁶⁻⁷ The RD process was first applied to the production of EtAc in 1932.⁷ Since the production of methyl acetate via a commercial RD process by Eastman Chemical Co. in the 1980s,⁸ it has become a new focus in both industry and academia.⁹⁻¹⁰

Pervaporation (PV) has been considered to be an effective technique to separate azeotropic mixtures and close-boiling-point mixtures because it is not limited by the volatility of the components.¹¹⁻¹³ As a low energy consumption, high selectivity, compact and modular design, and environmentally friendly process, PV has attracted intensive attention recently.¹⁴⁻¹⁷ However, PV is seldom used as a single process for the separation of species with high concentrations due to its relatively high capital cost and low capacity. Therefore, hybrid processes, coupling PV with other traditional separation technologies, e.g., a distillation process, are commonly used due to rising synergy effects and overcoming the disadvantages of each other.¹⁷⁻²¹ Lipnizki et al.¹⁷ made a detailed review of PV-based hybrid processes. Buchaly et al.¹⁸ described the modeling, simulation, and process analysis of hybrid separation processes combining RD with membrane separation for n-propyl propionate. Wang et al.¹⁹ proposed a new method for analyzing the RD–PV process for ethyl tert-butyl ether from the viewpoint of nonequilibrium thermodynamics and the phenomenological theory. Aïouache and Goto²⁰ placed a tubular NaA zeolite membrane in an RD column for the continuous removal of generated water. The total output of tert-amyl ethyl ether was 10% higher than that without water removal. Huang et al.²¹ introduced an innovative separation process, which combines distillation, mechanical vapor compression, and vapor permeation, to improve process energy efficiency greatly.

For the production of EtAc by the RD process, the separation of products is difficult because of this four-component system in the formation of several heterogeneous and homogeneous azeotropes. Most of the researchers focused on the effective separation of the distillate.²²⁻²⁵ In our previous work, a polyvinyl alcohol (PVA)/ceramic composite membrane was applied for the PV dehydration of the EtAc/H₂O binary and EtAc/EtOH/H₂O ternary systems.²⁶⁻²⁸ Since the water content in the stoichiometry composition is 17.0% while in the
azeotrope composition it is 8.7%, the distillate contains less water in comparison to the water produced by chemical reaction. Accordingly, most of the water will coexist with the unreacted HAc in the reboiler, which significantly increases the energy consumption of the reboiler. The removal of water from HAc is therefore necessary, and the recycle of a large amount of unreacted HAc can be realized simultaneously. However, the difference in the volatilities of water and HAc is so small, especially at high HAc concentrations, that the energy-intensive azeotropic distillation or extractive distillation is always used to separate water from HAc.

In this work a new RD−PV coupled process was proposed for the EtAc production, in which the RD process was used for esterification and the PV process was simultaneously applied to remove the water from the reboiler and recycle the HAc into the feed. The water removal is beneficial for the esterification reaction, and moreover, the large amount of unreacted HAc in the reboiler can be reused. Therefore, this coupled process is not a simple integration of RD and PV, but the PV process could have feedback to the RD process. To the best of our knowledge, although many processes are proposed for EtAc,7,29−34 this coupled process has not been reported to date. The synthesis of EtAc in an RD column was investigated via experiments and simulation using the Aspen Plus program. Meanwhile, we adopted the PVA/ceramic composite membrane for PV of model bottom solutions (HAc/H2O and HAc/EtAc/EtOH/H2O mixtures). Subsequently, the RD−PV coupled process was established and the effects of coupled temperature and HAc/EtOH molar ratio on the reaction conversion and EtAc in the distillate were systematically studied.

2. EXPERIMENTAL SECTION

2.1. Materials. Polyvinyl alcohol (PVA) was purchased from Shanghai RichJoint Chemical Reagents Co., Ltd., China. HAc, EtOH, maleic anhydride, and sulfuric acid of analytical reagent grade were obtained from Sinopharm Chemical Reagent Co., Ltd., China. The preparation of the PVA/ceramic composite membrane followed the method reported in our previous work.26

2.2. RD Experiment Setup, Packing, and Operation. Figure 1a shows the experimental apparatus. The pilot plant scale experiments were carried out in a glass column with an inner diameter of 30 mm and total height of 1.5 m at atmospheric pressure. The RD column consists of a 0.6 m long reactive section packed with homemade catalyst bales such as the catalyst licensed by CR&L containing 0.1 mm diameter Purolite CT275 (a strong macroporous cation exchange resin) with a packing density of approximately 185 kg/m3. The ion exchange capacity of CT275 is 5.2 H+ equiv/kg, and the maximum temperature limit is 145 °C. The stripping section is 0.3 m and the rectifying section is 0.6 m, and are packed with a Dixon ring (3 mm i.d.).

The column is equipped with an electronic reflux splitter to control the reflux ratio, a total condenser, and a reboiler with a four-neck round-bottom flask of a low liquid holdup (approximately 1 L). The liquid in the reboiler was heated by a heating jacket. The value of the reboiler heat duty was calculated from the electric power to the mantle heater. The column and the reboiler were insulated by mineral wool to ensure adiabatic operation.

A sample of 200 g of EtOH was placed inside the bottom flask and heated to 100 °C. When the distillate appeared at the top, the column was operated under total reflux until the temperature was maintained almost constant. Then HAc and EtOH were fed simultaneously by two pumps to the distillation column at the desired flow rates. The HAc feed port is located between the rectifying and reactive sections, and the EtOH feed port is located between the reactive and stripping sections. The excess liquid in the reboiler was withdrawn intermittently by another peristaltic pump to maintain the constant liquid level. The reflux splitter was adjusted to the desired values. Subsequently, the continuous operation was started and the experiment was conducted until steady state was achieved (usually 2.5 h). The distillate, bottom withdrawal liquids, and permeate were periodically collected and analyzed. The pressure drop along the column is 540 Pa/m. The mass balance in the steady-state operation is about 98% closure.

2.3. PV and RD−PV Coupled Process. The PV performance of the membrane was conducted on a homemade PV apparatus,26 and the RD−PV coupled process is shown in Figure 1b. The tubular PVA/ceramic composite membrane was
sealed in a so-called “pencil module,” with an effective membrane area of 110 cm². The PV feed was heated by the oil-bath heater and circulated by a liquid pump between the membrane module and the liquid tank with a circulation flow of 23.4 L/h. The vapor permeated through the membrane was collected in a liquid nitrogen trap, and the permeate pressure was kept at 300–400 Pa during collections. After the RD column reached a steady state, the membrane was coupled with the reboiler and the RD–PV coupled process was started. The RD process was kept going while the generated water was removed from the reboiler simultaneously via the pervaporation process.

The PV performance of a membrane is usually expressed in terms of the permeation flux \( J_i \) and separation factor \( \alpha \).

\[
J_i = \frac{W_i}{Ar} \tag{1}
\]

\[
\alpha = \frac{x_i}{y_i} \tag{2}
\]

where \( W_i \) is the weight of component \( i \) in the permeate (kg), \( A \) is the effective area of the membrane (m²), and \( t \) is the permeation time interval for the PV (h); \( x_i \) and \( y_i \) represent the weight fractions of the components \( i \) and \( j \) in the permeate and feed, respectively.

The selectivity of the membrane \( \beta \) is defined as

\[
\beta = \frac{Q_i}{Q_j} \tag{4}
\]

where \( Q_i \) and \( Q_j \) are the permeances of components \( i \) and \( j \), respectively.

The apparent activation energy \( (E_{\text{app}}) \) is calculated using the Arrhenius equation:

\[
J_i = K \exp\left(\frac{-E_{\text{app}}}{RT}\right) \tag{5}
\]

where \( J_i \) is the flux of component \( i \), \( K \) is the permeation rate constant, \( T \) is the feed absolute temperature, and \( R \) is the molar gas constant.

2.4. Analysis. A gas chromatograph (GC-2014, SHIMADZU, Japan) equipped with a thermal conductivity detector (TCD) was used to analyze the mixture of EtOH, HAc, EtAc, and water. The analysis process was carried out in a Porapak-Q column with helium as a carrier gas at a flow rate of 25 mL/min. The oven temperature was varied from 120 to 180 °C.

3. RESULTS AND DISCUSSION

In order to establish the RD–PV coupled process, the main influencing factors of the individual RD and PV process were investigated respectively, because the performance of the RD–PV coupled process was to some extent determined by these two individual processes. For the RD process, three conventional factors are chosen: reflux ratio, HAc/EtOH molar ratio, and total feed rate; for the PV process, the operating temperature and feed concentration are usually the main factors.

3.1. RD Experiment and Simulation. 3.1.1. Effect of Reflux Ratio. The reflux ratio is a very important parameter in an RD process. The influences of the reflux ratio on the EtAc purity in the distillate, the water concentration in the reboiler, and the EtOH conversion were investigated. The molar ratio of HAc to EtOH was set at 1.1, the flow rate of EtOH was 42 g/h, and the mass ratio of distillate to feed (D/F) was set at 0.71.

As shown in Figure 2, both the EtAc purity in the distillate and the water in the bottom are highest at the reflux ratio of 3.

![Figure 2. Effect of reflux ratio on (a) EtAc purity in the distillate and water concentration in the reboiler and (b) EtOH conversion (HAc/EtOH molar ratio = 1.1, EtOH feed rate = 42 g/h and D/F = 0.71).](image)

The conversion of the EtOH is 81.3 wt %. The increase of the reflux ratio from 1 to 3 overcomes the chemical equilibrium limitation, enhances the conversion beyond the chemical equilibrium conversion, and improves the EtAc purity in the distillate and EtOH conversion. When the reflux ratio increased from 3 to 6, the high reflux ratio led to high EtAc concentration in the reactive zone, reducing the residence time and suppressing the shift of the positive reaction; thus the EtAc concentration in the distillate and the EtOH conversion changed slightly. Considering the energy consumption, the reflux ratio of 3 is appropriate.

3.1.2. Effect of HAc/EtOH Molar Ratio. It is well-known that increasing one of the reactants in an equilibrium limited reaction can lead to increasing the conversion of the other reactants. For EtAc production, excessive HAc is usually used. The effect of the HAc/EtOH molar ratio on EtAc purity in the
distillate and water concentration in the reboiler was studied with the EtOH feed rate and reflux ratio of 42 g/h and 3, respectively. The results are shown in Figure 3.

EtAc purity and EtOH conversion increase with increasing HAc/EtOH molar ratio. EtAc purity in the distillate is 84.4 wt% at the ratio of 1.7. Because of the low EtOH concentration, the distillate can be separated into organic phase and water phase spontaneously. It is noted that the EtAc purity increases rapidly as the HAc/EtOH molar ratio increases from 1.7 to 2.0, whereas the water concentration has not been changed dramatically. At the HAc/EtOH molar ratio of 2.0, the water concentration is about 10 wt% in the reboiler, which is half that at the molar ratio of 1.1. Considering the economy, the appropriate molar ratio is 2.0.

3.1.3. Effect of Total Feed Rate. The total feed rate has a significant influence on the residence time of the reactants in the reactive zone. Experiments were carried out to study the effect of total feed rate on the performance of RD with the reflux ratio and HAc/EtOH molar ratio of 3 and 2, respectively. As shown in Figure 4, when the feed rate increases, the residence time of the reactants decreases, causing the drop in both EtAc purity and EtOH conversion. Considering the column capacity and EtOH conversion, a total feed rate of 152 g/h is appropriate.

Furthermore, the Aspen Plus program was applied to check the optimum results of the RD experiment (see Supporting Information). The simulation was conducted under optimized RD parameters. The results and the corresponding experimental data are listed in Table S1 in the Supporting Information. It can be found that the theoretical results agree well with the experimental data for both distillate and bottom concentrations.

3.2. PV Dehydration of Bottom Solutions. 3.2.1. PV Dehydration of HAc/H2O Solution. The PV dehydration of RD bottom solutions was investigated first based on the model HAc/H2O mixtures. The effects of operating temperature and feedwater content on the PV performance were investigated. It should be noted that the temperature of the RD reboiler was 100 °C and the maximum water content was less than 30 wt%. See Figure 5.

When the operation temperature increased, the partial pressure of the water and HAc in the feed side increased while the vapor pressure at the permeate side was not affected. As a consequence, the driving force for mass transfer across the membrane increased with the feed temperature. In addition, the diffusion of the water and HAc molecules was enhanced due to the high temperature, which favored the mass transfer of the process. Furthermore, high temperature would increase the mobility of the polymer chains and the expansion of the free volume. All these factors favored the increase of the water flux and the HAc flux, and the increase of the HAc flux was higher than that of water. Therefore, increasing temperature led to an increase in flux and an decrease in separation factor.

As water in the feed increases, the permeation flux of the membrane increases, whereas the separation factor significantly decreases (Figure 5). Increasing the water content in the feed facilitates the water sorption into the PVA membrane; thus the hydrophilic PVA layer is more swollen. This makes the diffusion of both water and HAc molecules through the membrane easy. As a result, the total permeation flux increases and the separation factor decreases.
Comparison of the PV performance of this work with other reports for HAc/H2O mixtures is summarized in Table 1. Considering the different operating conditions of the PV experiments, the permeance and selectivity that reflect the intrinsic properties of the membranes were also calculated by eqs 3 and 4. It can be seen that the PVA/ceramic composite membrane exhibits good performance for the dehydration of HAc/H2O mixtures, and might be a promising PV membrane for the RD−PV coupled process.

3.2.2. Long-Term Stability of the PVA/Ceramic Composite Membrane. The lifetime or durability of the membrane is one of the important factors that should be considered for practical application. The long-term stability of PVA/ceramic composite membrane was investigated in the HAc/H2O system at 70 °C, as shown in Figure 6. During the 100 h testing period, the permeation flux and separation factor are steadily kept at around 0.6 kg·m⁻²·h⁻¹ and 14, respectively. Because of the sufficient mechanical, chemical, and thermal stabilities of the PVA/ceramic composite membrane, it can be adopted for the dehydration of HAc/H₂O solution at relatively high temperature. The three-dimensional swelling of the organic layer is restricted by the ceramic supports, which is beneficial to the long-term stability of the PVA/ceramic composite membrane.41

3.2.3. PV Dehydration for HAc/EtAc/EtOH/H₂O Quaternary Solutions. The PVA/ceramic composite membrane shows good dehydration performance for the HAc/H₂O binary solution. In most of the RD process, however, the bottom products are HAc/H₂O/EtAc/EtOH quaternary solutions. Therefore, the PV dehydration performance of the PVA/ceramic composite membrane in a typical HAc/H₂O/EtAc/EtOH (80/12/6/2 wt %) quaternary solution was investigated at different temperatures, as shown in Figure 7. As temperature increases from 50 to 100 °C, the total flux increases from 0.35 to 1.27 kg·m⁻²·h⁻¹ while the water concentration in the permeate decreases from 67.7 to 63.3 wt %. The permeation fluxes and concentrations of EtOH and EtAc are very low. Furthermore, the Arrhenius plots of the PV performance of the PVA/ceramic composite membrane are shown in Figure 8. Good linearity relationships between the permeation fluxes and the reciprocal of the temperature (1/T) can be observed. The activation energy values for each component through the composite membrane were calculated from the slope. The activation energy values for water, HAc, EtOH, and EtAc are 25.1, 29.4, 28.7, and 16.6 kJ·mol⁻¹, respectively.

3.3. RD−PV Coupled Process. As discussed above, the operating temperature and water concentration have significant effects on the permeate flux and composition of the PV process. Thus, to investigate the effect of the PV process on the RD−PV coupled process, two critical factors were chosen: the PV operating temperature and the HAc/EtOH molar ratio of RD that determines the water concentration of the bottom solution. The experiments were carried out at a constant permeate flow rate of 0.6 kg·m⁻²·h⁻¹.
pressure of 350 Pa and a retentate reflux to HAc feed of 30 mL/h.

3.3.1. Effect of PV Operating Temperature. At the optimal conditions of the RD process (reflux ratio = 3, HAc/EtOH molar ratio = 2, total feed rate = 152 g/h), the effect of PV operating temperature on the performance of the RD–PV coupled process was investigated. As shown in Figure 9, the higher PV operating temperature is beneficial for the RD–PV coupled process. The EtAc purity in the distillate and the EtOH conversion reach 91.0 wt % and 90.2%, respectively, with the PV process operated at 100 °C.

The “coupling effect” of the RD–PV coupled process can be stated as follows: since water was removed from the bottom, the esterification was shifted toward the formation of more EtAc. In addition, due to the increase in the permeation flux and HAc content in the retentate, the actual HAc/EtOH molar ratio increased in the RD column, thus improving both the EtAc purity and the EtOH conversion simultaneously. Furthermore, the permeation fluxes of EtOH and EtAc are very low (as displayed in Figure 7); thus the unreacted EtOH (ca. 4 wt %) in the reboiler was reflowed to the reaction zone to react with HAc, improving the EtAc purity and EtOH conversion as well. Meanwhile, the EtAc in the reboiler was also reflowed to the reaction zone, which is beneficial for higher EtAc purity in the distillate.

3.3.2. Effect of HAc/EtOH Molar Ratio. In the RD process, as the HAc/EtOH molar ratio increases, a large amount HAc coexists with the water at the column bottom, which leads to the increase of the separation burden of the reboiler compounds. Therefore, we attempted to study the RD–PV coupled process at low HAc/EtOH molar ratios, in order to relieve the burden of reboiler. The performance of the RD–PV coupled process at various HAc/EtOH molar ratios at 70 °C is shown in Figure 10. It was found that by coupling with the PV process the performance of the RD process could be improved under all HAc/EtOH molar ratios. Moreover, the performance improvement of the RD process via coupled with PV at a low HAc/EtOH molar ratio was much higher than that of the RD–PV coupled process operating at a high HAc/EtOH molar ratio. The reason is that, at the molar ratio of 1.1, the high water concentration in the reboiler leads to a high permeation flux of
the PV process (as displayed in Figure 5a), resulting a significant “coupling effect” as discussed in section 3.3.1.

We also found that, when coupled with PV, the EtAc purity and EtOH conversion at the molar ratio of 1.1 was higher than those of the stand-alone RD at the molar ratio of 1.7; the performance of the RD–PV coupled process at the molar ratio of 1.7 was nearly up to that of the stand-alone RD at the molar ratio of 2.0, as well. This result suggests that the coupling of the PV process could improve the RD performance at a low HAc/EtOH molar ratio, meanwhile keeping a high product quality and reaction conversion. It favors the subsequent treatment of the bottom solution. Furthermore, the distillate can be separated into the organic phase and water phase due to the low EtOH concentration in this case, thus making the subsequent separation much easier.

To further investigate the effect of the PV process in this coupled process, we also carried out the RD process only with a recycle stream into the HAc feed without the water removal by the PV process. The results comparison is also shown in Figure 10. The recycle of the reboiler has two effects on the performance of the reaction. On one hand, the improved molar ratio of HAc to EtOH will facilitate the shift of positive reaction; on the other hand, the water in the recycle will also suppress the shift of positive reaction. When the molar ratio was 1.1 and 1.7, the domination of former effect resulted in the increase of the conversion and EtAc purity in the distillate. As the molar ratio increased to 2.0, the latter effect became dominant, leading to the drop of both the conversion and EtAc purity. Nevertheless, in the RD–PV coupled process, due to the water removal from the reboiler by the PV membrane, both the conversion and the EtAc purity were higher than those with the RD process only with a recycle stream. Therefore, it was further proved that the improvement of EtOH conversion and EtAc purity in the distillate were due to the water removal from the reboiler and the HAc recycle to feed by the coupling of the PV process. Especially, the pervaporation removal of water would play a dominant factor under high feed HAc/EtOH molar ratios.

4. CONCLUSIONS

A new RD–PV coupled process with PV selectively removing water from the reboiler and recycling HAc into the feed for EtAc production is presented. An optimal RD operating condition was obtained via experiments and Aspen Plus simulation. The PVA/ceramic composite membrane showed good performance and stability with a total flux of 600 g-m⁻²-h⁻¹ and separation factor of 14 in 90 wt % HAc/water solution at 70 °C. As a result, the removal of water and recycle of bottom withdrawal by PV coupling significantly improved both the product purity and the reaction conversion of the RD process, in which the ethyl acetate purity and ethanol conversion simultaneously increased from 82.4 to 85.6 wt % and from 81.3 to 84.8%, respectively, at low the HAc/EtOH molar ratio of 1.1. Our work demonstrated that this novel RD–PV coupled process based on the PVA/ceramic composite membrane could be a promising method to produce ethyl acetate. Further work could focus on the development of a higher-selective membrane for the HAc/H₂O system, which will solve the problem of high HAc contents in the PV permeate.

ASSOCIATED CONTENT

Supporting Information

Application of Aspen Plus program to check the optimum results of the RD experiment. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.

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