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### Introduction

Organic acids are organic compounds composed of a weak acid group, ranging widely from carboxylic ( $pK_a = 3$ ) to phenolic groups  $(pK_a = 9)$ .<sup>1</sup> Due to their biodegradability, there has been an increase in the applications of organic acids in the last decade; for example, lactic acid has significant potential for applications in biodegradable plastics and bio-adaptable medical sutures;2-4 succinic acid plays an important role as a precursor molecule in the synthesis of biodegradable

recovery

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polyester resins, dyestuffs, and pharmaceuticals;5 citric acid is widely used as a flavoring agent for foods and beverages as well as a cleaning and polishing agent,<sup>6-8</sup> and formic acid is mainly used in the textile and paper industries.9,10

Electro-membrane processes for organic acid

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recovery with lower byproduct concentration and energy consumption.

With an increase in the organic acid requirement, the production of organic acids has been increased over the years. To achieve cost-effective production of organic acids, efficient recovery processes are needed. Electro-membrane processes, including electrodialysis (ED), electrometathesis (EMT), electro-ion substitution (EIS), electro-electrodialysis (EED), electrodialysis with bipolar membrane (EDBM), and electrodeionization (EDI), are promising technologies for the recovery of organic acids. In the electromembrane processes, organic acids are separated from water and other impurities based on the electro-

migration of ions through ion-exchange membranes. These processes can recover various types of

organic acids from the fermentation broth with high recovery yield and low energy consumption. In

addition, the integration of fermentation and the electro-membrane process can improve the acid

In general, there are two routes for the production of organic acids: chemical synthesis and carbohydrate fermentation. The chemical synthetic route is limited by its production capacity because it is associated with the byproduct of another process<sup>11</sup> and has high manufacturing costs; moreover, carbohydrate fermentation can utilize various raw materials and produce organic acids with a higher degree of safety; thus, it has mainly been chosen to produce organic acids in the recent years.<sup>12</sup> A wide variety of renewable resources, such as silage, grains, syrups, molasses, and cheese whey, can be used as raw materials in carbohydrate fermentation. However, the fermentation



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broth obtained from these resources usually consists of various ingredients, and thus, its separation and purification are necessary to achieve high purity of the organic acids.

About 50-70% of the cost of the production of organic acid is generated by the recovery of the fermentation product.13,14 Therefore, various technologies, such as precipitation, extraction, adsorption, ion-exchange system, membrane, etc.,15-23 have been developed for the efficient recovery of organic acids. However, some of these conventional technologies, especially extraction, adsorption, and ion-exchange, require further concentration steps, produce dangerous waste, require hazardous solvents, and consume high energy.12 Therefore, the recovery of organic acids has been shifted from conventional technologies to electro-membrane processes including electrodialysis (ED), electrometathesis (EMT), electro-ion substitution (EIS), electro-electrodialysis (EED), electrodialysis with bipolar membranes (EDBM), and electrodeionization (EDI). The electro-membrane process separates organic acids from water and other impurities based on the electro-migration of ions through ion-exchange membranes. Thus, it can provide high

quality product in a short time without salt introduction or discharge.<sup>3</sup>

Fig. 1 shows the number of publications on the recovery of organic acids since 1980. It can be seen that the number of publications on the recovery of organic acids has increased over years, with the highest number of publications in 2010. Based on technology, the publications on the recovery of organic acids via extraction was dominant with up to 51%. However, since 2000, there has been an increase in the number of publications on the recovery of organic acids via electro-membrane processes due to their ability to recover organic acids with high yield and without the requirement of solvent addition. In the earlier studies, ED was the only electro-membrane process applied for organic acid recovery. Compared to conventional processes, ED requires lower energy consumption and eliminates the use of solvent. However, the purity of the acid product from the ED process is relatively low. Therefore, many researchers have tried to develop the ED process via stack modification (i.e. EMT, EIS, and EED), membrane modification (i.e. EDBM), resin addition (i.e. EDI), and integration with other processes.



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Fig. 1 The number of reported studies related to the search term organic acid recovery, as indexed by Scopus (TITTLE-ABS-KEY (terms); September 2018).

A review on the recovery of organic acids *via* electromembrane processes was published by Huang *et al.* in 2007.<sup>12</sup> However, their review mainly focused on the ED configuration and its economic evaluation. Thus, for a better understanding of this process, we aim to provide a more comprehensive review of the recovery of organic acids using electro-membrane processes. The conventional processes for the recovery of organic acids are summarized to give a better overview on the state of the art of organic acid recovery. Meanwhile, the discussion on electro-membrane processes includes not only their configuration, but also their performances and design development on the recovery of various organic acids. In addition, the issues with electro-membrane processes and their possible solutions are discussed.

# Conventional processes for organic acid recovery

Separation and purification of fermentation broth are the primary determinants for the cost-effective production of organic acids. Calcium precipitation using Ca(OH)<sub>2</sub> or CaO is one of the most used methods to recover organic acids from fermentation broth. This method has been used for recovery of citric acid,25,26 lactic acid,27 and succinic acid.5,28,29 Calcium precipitation offers the advantage of being directly performed on existing equipment, technology and infrastructure, and thus it has very low technological barriers and risks.24 Beside calcium precipitation, the recovery of succinic acids also has been conducted by ammonia precipitation. Succinic acid was recovered up to 93.3% using ammonia precipitation.30 Generally, precipitation produces a low amount of waste by-products since the used reagent can be recycled. However, this method requires high energy consumption and equipment erosion due to the low pH and high temperature in the reagent recycle process.

Liquid–liquid extraction has also been widely used for the recovery of organic acids. The liquid–liquid extraction system consists of two separate liquid phases, where the organic acid transfers from one phase to the other based on the solubility differences between the phases.<sup>31</sup> In liquid–liquid extraction, the selection of an appropriate solvent is fundamental to obtain a high organic acid recovery rate. There are seven important criteria for solvent selection: (a) good distribution coefficient for organic acid uptake, (b) high separation capacity, (c) high selectivity, (d) easy product backwashing with water, (e) low tendency to emulsify, (f) low cost, and (g) non-toxic.<sup>32</sup> Several organic solvents have been widely used, such as butyl alcohol, acetone and tributyl phosphate or with certain amines.<sup>23,33–36</sup> Among them, amine extraction is one of the most prospective methods for the separation of carboxylic or hydroxyl-carboxylic acid from aqueous solution.<sup>32,37,38</sup>

In several cases, the distribution of organic acids in the extraction phase seems to be unsatisfactory, and thus large quantities of extraction agent are required. Consequently, some researchers developed reactive extraction to solve this problem. The product from fermentation is firstly converted into a compound without carboxyl groups and then recovered by liquid–liquid extraction.<sup>24</sup> This method has been used to recover succinic acid in several studies.<sup>5,39,40</sup> The results showed that more than 95% succinic acid was recovered by reactive extraction. This method also has been applied for lactic acid,<sup>20,41-44</sup> acetic acid,<sup>45-48</sup> carboxylic acid,<sup>49,50</sup> and propionic acid<sup>21,51,52</sup> since these organic acids are poorly extractable by common organic solvents due to their hydrophilic nature.

The other alternative for the recovery of organic acids is the use of solid sorbents that are selective for organic acids. To obtain high recovery yield, the sorbent must have a high separation capacity for the acid and specificity for the product.24,41 Alumina, activated carbon, silica, and zeolite molecular sieves are examples of sorbents widely used to recover organic acids from fermentation broth. To adsorb succinic acid, alumina53 and a high-silica zeolite  $(SiO_2/Al_2O_3 = 218)^{54}$  were chosen as sorbents. Meanwhile, for lactic acid recovery, adsorption by silicalite molecular sieves15 and activated carbon55 was used. In addition, activated carbon also has been utilized to recover acetic acid, butyric acid, fumaric acid, and propionic acid.18,19 Several researchers also studied the use of ion exchange resins to adsorb organic acids. Various types of ion exchange resins such as alkaline-type anion exchange resins,56,57 weak base anion-exchange resins (Amberlite IRA-92,58 IRA-400,17,59-61 and IRA-900 (ref. 62)) and poly(4-vinylpyridine) (PVP) resins<sup>55,63-65</sup> have been used. The studies showed that a weakly basic ion exchange resin was the best ion exchange resin for the purification of organic acids.

Membrane technologies have also proven for the advanced recovery of organic acids. Compared to other conventional technologies, pressure-driven membranes offer the advantages of process continuity and high selectivity. For small molecules such as organic acids and salts, nanofiltration (NF) membranes can be used to separate them efficiently. NF membranes consist of a dense, ultrathin skin layer on microporous polymeric supports, and mostly with charged groups on the membrane surface.<sup>66</sup> The rejection mechanism in NF membranes is a combination of size sieving, solution-diffusion, and Donnan exclusion.<sup>67</sup> NF has been used for the recovery of lactic acid by

several researchers.68-71 These studies showed that NF was able to recover highly purified lactic acid by removing 85% of mineral ions such as  $Na^+$ ,  $K^+$ ,  $Mg^{2+}$ , and  $Ca^{2+}$  and 90% of residual carbohydrates. Several researchers also tried to combine NF with other processes such as microfiltration and ultrafiltration to improve the recovery efficiency.72-75 In addition, membrane distillation (MD) has also been studied to recover organic acids. The organic acid is separated from impurities by heating and evaporating the liquid and allowing the vapor to pass through a microporous hydrophobic membrane. Then the vapors are allowed to condense into liquid by cooling on the other side of the membranes.<sup>76</sup> Ban et al.<sup>77</sup> used this method to recover various types of organic acids, including glycolic acid, glyoxylic acid lactic acid, pyruvic acid, malonic acid, and glutaric acid. Their results showed that the rejection rate for all the abovementioned organic acids except pyruvic acid was above 97.0%. Generally, MD is able to recover organic acids with a high recovery rate; however, its thermal energy efficiency is very low.

Table 1 shows the recovery yield of organic acids using various conventional technologies. It can be seen that most of them were successfully utilized to recover organic acids with recovery yields greater than 60%. However, each technology has advantages and disadvantages, as summarized in Table 2. Calcium precipitation offers the advantage of the ability to be directly used on existing equipment, technology, and infrastructure, and thus has very low technological barriers and risks. However, it has low selectivity and produces  $CaSO_4$  sludge. Liquid–liquid extraction offers some advantages, such as high purity product and low energy consumption, but the use of hazardous solvents leads to environmental problems. Moreover, adsorption using solid sorbents or ion exchange resins is a reliable technology, but it requires large amounts of chemicals for the regeneration of the ion exchange resins and adjustment

of the feed pH to increase the sorption efficiency. Meanwhile, pressure-driven membranes have several advantages, such as process continuity and easy scale-up, compared to conventional technologies; however fouling is still the main limitation in membrane separation processes.

# Electro-membrane processes for organic acid recovery

The improvement of technologies for organic acids recovery is very important since it is associated with fermentation process effectiveness and production costs. The conventional technologies generally need further concentration steps, produce dangerous waste, require hazardous solvents, and consume high energy. To solve those problems, organic acid recovery has shifted to electro-membrane processes. In electro-membrane processes, organic acids are separated from water and other impurities based on the electro-migration of ions through ionexchange membranes. These processes are able to recover various types of organic acids from fermentation broth with high purity and yield without requiring the use of solvent.

#### Electrodialysis (ED)

One of the promising technologies to obtain high purity organic acids without the requirement of any solvent is electrodialysis or ED. ED is an electrochemical separation process with the driving force of electrical potential difference to separate ionic species through ion exchange membranes. It has been applied for the production of table salt, organic acid recovery, heavy metal recovery, and sugar demineralization.<sup>98-101</sup> ED mainly consists of three compartments, including diluate, concentrate, and electrode compartments. The compartments are separated

Acid	Method	Recovery yield (%)	Ref.
Acetic acid	Extraction	66–97	35, 45, 46, 48 and 78
	Adsorption	94	79
	Pressure-driven membrane	88	80
Citric acid	Extraction	90	37
Formic acid	Extraction	87	81
Fumaric acid	Extraction	70.7	22
	Adsorption	85-93	19 and 62
Lactic acid	Extraction	37-97	20, 23, 31, 33, 43 and 82-84
	Adsorption	74–95	15, 17, 65 and 85
	Precipitation	92	27
	Pressure-driven membrane	60-100	74, 75 and 80
Picolinic acid	Extraction	75-96.6	38 and 86
Propionic acid	Extraction	75	21
	Adsorption	64	18
Pyruvic acid	Extraction	40-82	87 and 88
Succinic acid	Extraction	67–95	40 and 89–91
	Precipitation	93.3	30
	Adsorption	96-99	57 and 62
Tartaric acid	Extraction	90	23
	Adsorption	75-99	92 and 93

Table 2	Comparison	of the	technologies	for organic	acid recovery
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Technology	Advantages	Disadvantages	Ref.
Precipitation	Very low technological barriers a	and Resulting $CaSO_4$ sludge (notorious for solid pollution)	12, 14 and
-	risks	Low selectivity	24
Extraction	High purity product	Undesirable distribution coefficients	14, 24 and
	Low energy consumption	Environmental problems due to the use of hazardous solvents Expensive extraction agent and diluent	94
Adsorption	Easy to operate	Short lifetime of adsorbents	12 and 18
•		Low capacity	
Ion exchange	Easy to operate	Consumes a great amount of acid, base, and water to regenerate ion exchange resins	12
Pressure-driven	High selectivity	Fouling formation	95
membrane	Easy to operate	Ũ	
	Easy to scale-up		
Electro-membrane	High purity product	Fouling formation	96 and 97
processes	No need salt introduction or discharge		

by anion and cation exchange membranes arranged between an anode and cathode,<sup>99</sup> as shown in Fig. 2(a).

membranes (CEM) possess negatively charged fixed groups and allow the migration of cations, while anion exchange
 membranes (AEM) possess positively charged fixed groups and allows the migration of anions. When a sufficient electrical field





Fig. 2 Schematic of the stack configuration for: (a) ED, (b) EMT, (c) EIS, (d) EED, (e) EDBM, and (f) EDI to produce organic acids.

is applied on the solution, the mobile ions are replaced by same charged ions (counter ion) from the solution and the membrane matrix will repel ions that have the same charge as the membrane matrix (co-ion). This mechanism allows the selective passage of ions.<sup>103</sup>

To recover organic acids from fermentation broth, the feed solution is circulated on each compartment of the ED system, while a potential difference is applied between the two electrodes. Due to the potential difference, positive ions (cations) migrate to the negative electrode (cathode), while the negative ions (anions) migrate to the positive electrode (anode). After some time, organic acids as ionic species are stripped from the diluate compartment and collected in the concentrate compartment. Uncharged molecules are not affected by this driving force, and hence organic acids can be separated from the uncharged components in the solution.<sup>104</sup>

#### **Electrometathesis (EMT)**

EMT is a modified ED process, which has been used for the recovery of citric acid<sup>105</sup> and lactic acid.<sup>94,106</sup> Generally, an EMT process consists of four compartments formed by two alternately arranged anion exchange membranes and two cation exchange membranes with two types of feed streams and outputs, as shown in Fig. 2(b). EMT can realize double composition reactions that hardly occur in other conditions than that in ED systems. In EMT, inorganic acids such as HCl and  $H_2SO_4$  are necessary for the conversion of organic salts into organic acids.<sup>12</sup> The addition of strong acid such as HCl in the product compartment helps to facilitate the transport of ions.<sup>107</sup>

#### **Electro-ions substitution (EIS)**

EIS is another modified ED technique that consists of two adjacent cation-selective (or anion-selective) membranes, one anion-selective (or cation-selective) membrane, and three compartments, as shown in Fig. 2(c). EIS can realize substitution between ions of the same sign. The  $H^+$  needed for ion substitution is supplied by an additional feed stream using an inorganic acid, such as  $H_2SO_4$ . At the beginning of the EIS process, a current efficiency of more than 100% can be reached due to  $H^+/M^+$  Donnan dialysis. However, more  $M^+$  in the acid stream will compete with  $H^+$ , and hence decrease the current efficiency. To suppress this competition, a cation-selective membrane with a higher  $H^+/M^+$  permselectivity can be added.<sup>12</sup>

#### Electro-electrodialysis (EED)

EED is an alternative methods that can be used to convert organic salt into organic acid due to its effective functional integration of ion exchange membranes and water splitting formed at the cathode and anode by reduction and oxidation.<sup>108</sup> A schematic representation of the EED process is presented in Fig. 2(d). In EED, the anion exchange membrane is essential since the acids are displaced by permselective transport of acid anions through the anion exchange membrane and electrodialysis of protons from the electrochemical reactions that occur at the electrodes.<sup>109,110</sup> This method was mainly used for the production of formic acid.<sup>110,111</sup> Formate anions (HCOO<sup>-</sup>) pass through the anion exchange membrane into the anolyte compartment. In this compartment, formic acid is formed by combining formate anions with the protons produced by anodic water oxidation. Meanwhile,  $H^+$  is retained in the catholyte compartment and reacts with hydroxide ions to form water.

Besides formic acid, EED also has been used for the recovery of lactic acid,<sup>112</sup> acetic acid,<sup>113</sup> salicylic acid,<sup>114</sup> glutamic acid,<sup>115</sup> and propionic acid.<sup>116</sup> Generally, the overall electrical current efficiencies of EED are much higher than that from conventional ED. However, the concentration ratio of the concentrated solutions is low, even less than 1.0.<sup>117</sup> Therefore, some studies developed two-phase electro-electrodialysis (TPEED) to improve the recovery ratio of the produced acid. TPEED has been used to recover citric acid<sup>117</sup> and lactic acid.<sup>112,118</sup> This method successfully increased the acid concentration ratio with half the energy consumption for EED.

#### Electrodialysis with bipolar membranes (EDBM)

To improve the separation performance of ED, several researchers combined the conventional ED and water dissociation of bipolar membranes, exploring a new avenue for the development of conventional ED. A bipolar membrane is a combination of a cation-selective and an anion-selective membrane; however, it has distinctive functions from that mono-polar membranes. Under a reverse potential bias, bipolar membranes can realize the dissociation of solvent molecules.12 EDBM can split and separate water into H<sup>+</sup> and OH<sup>-</sup> ions and the membranes can operate at about 80% of the theoretical thermodynamic efficiency.<sup>119</sup> This technology also enables H<sup>+</sup> to be transported to the acid anion to form the free acid, while the OH<sup>-</sup> ion is transported to the cation compartment to form the free base. Therefore, separation efficiency and acid purity can be increased by the addition of bipolar membrane. Besides, the effluent (glucose, calcium, magnesium, etc.) from EDBM can be reused for fermentation after simple processing. However, the cost of membranes is still the main limitation for the large-scale application of EDBM.120

For application in organic acid recovery, EDBM is mainly operated using a three-compartment configuration, as shown in Fig. 2(e). This configuration consists of brine, base, and acidcontaining streams.<sup>121</sup> The fermentation broth passes through the membrane stack in the brine compartment, which is placed between an anion and a cation exchange membrane. The compartment between the cation membrane and the anion side of the bipolar membrane is filled with the base stream, where hydroxide is generated. Meanwhile, the acid stream is placed between the anion membrane and the cation side of the bipolar membrane, where acid is generated. The anions are transported from the brine compartment to the acid compartment through the anion membrane, while cations are transported from the brine compartment to the base compartment through the cation membrane. In this configuration, hydrochloric acid and Na/K hydroxide can be recovered in the acid and base compartments, respectively.102

#### **Electrodeionization (EDI)**

EDI is a modified ED process with the presence of ion exchange resins between the anion and cation exchange membranes, as

illustrated in Fig. 2(f). The presence of ion exchange resins enhances the ionic conductivity of a dilute solution by absorbing and concentrating the ions on the resin beads. Besides, the water splitting reaction on the resins surface enables in situ regeneration of the ion exchange resins and aids in the transport of ionic species across the ion exchange membranes.7 The water splitting reaction produces protons and hydroxyl ions at the resin-resin and resin-membrane interfaces. These ions continuously replace the adsorbed ions on the ion exchange resins surface; thus, resulting in more conductive ion exchange resins pathway.<sup>122</sup> Thus, EDI offers a continuous process where ion exchange resins can be continuously regenerated by a direct current electric field.<sup>123</sup> Due to its ability to produce high purity products, EDI is mainly applied for the production of ultrapure water.<sup>124-126</sup> However, several studies on the recovery of organic acids using EDI also have been established, such as the recovery of citric acid,<sup>7</sup> amino acid,<sup>127</sup> butyric acid,128,129 and lactic acid.123

# Performance of organic acid recovery using electro-membrane processes

For high productivity, fermentation-based processes require maintenance of near neutral pH and addition of alkali in most cases for the production of organic acids. The addition of alkali leads to the formation of organic salt instead of organic acid. To overcome this problem, electro-membrane processes have attracted the attention of several researchers since they do not require the addition of acid or alkali.

Table 3 summarizes the reported results for various electromembrane processes for the recovery of organic acids. Among the various organic acids, many researchers have focused on the recovery of lactic acid due to its multifunctional applications as a preservative in the food, pharmaceutical, leather, and textile industries, and a chemical feedstock.<sup>130</sup> The recovery of lactic acid has been conducted using ED, EIS, EDBM, and EDI. The results showed that these electro-membrane processes were able to recover more than 90% of lactic acid with a final concentration of up to 185 g L<sup>-1</sup>.

The electro-membrane process also gave good results for the recovery of other organic acids. ED and EDBM have been used for almost all organic acids. ED was able to recover 63–65% amino acid<sup>133</sup> and tartaric acid,<sup>152</sup> 85% of butyric acid, and more than 90% of glyceric acid<sup>140</sup> and lactic acid.<sup>3,16,143</sup> Meanwhile, EDBM has been used for the recovery of acetic acid,<sup>132</sup> amino acid,<sup>134,135</sup> citric acid,<sup>137,138</sup> fumaric acid,<sup>139</sup> lactic acid,<sup>142</sup> malic acid,<sup>145</sup> propionic acid,<sup>116</sup> salicylic acid,<sup>121,146</sup> and succinic acid.<sup>149</sup> Most of the studies on EDBM mainly focused on the current efficiency, where the results showed that the current efficiency for organic acid recovery was affected by the EDBM stack configuration and feed characteristics.

Besides ED and EDBM, several researchers used EED for the recovery of acetic acid, formic acid, and propionic acid. Koter<sup>113</sup> used EED for the separation of acetic acid, which resulted in a high acetic acid retention efficiency (>90%) when the process was conducted at current densities lower than the limiting

current density. Akgemci *et al.*<sup>110</sup> and Luo *et al.*<sup>111</sup> investigated the current efficiency of EED for formic acid recovery and obtained a current efficiency of more than 100%. The current efficiency of 90–99% was obtained for the recovery of propionic acid by Boyaval *et al.*<sup>116</sup> Meanwhile, EDI has been used for the recovery of amino acid,<sup>127</sup> butyric acid,<sup>128,129</sup> citric acid,<sup>7</sup> and lactic acid.<sup>123</sup> The results showed that EDI is a potential technology for the recovery of organic acids with a high current efficiency and recovery rate.

In general, electro-membrane processes are feasible to recover organic acids with a high acid recovery ratio and low energy consumption. However, each electro-membrane process has advantages and disadvantages, as presented in Table 4. ED and EED are simple processes without the need for acid; however, their recovery rate and product concentration are relatively lower than other electro-membrane processes. EMT and EIS can realize double composition reactions but require strong acid to facilitate the transport of ions. Meanwhile, EDBM and EDI can obtain high acid purity. However, EDBM has not been industrialized widely due to the high price of bipolar membranes, while EDI requires pretreatment.

## Development of electro-membrane process designs for organic acid recovery

The stack configuration plays an important role in the efficiency and recovery yield of electro-membrane processes. Several modifications of the electro-membrane configuration have been conducted to improve the recovery yield of organic acids for both the laboratory and industrial scales. Besides, the combined process of organic acid production and separation, which is mostly called the *in situ* separation process, also has been proposed to avoid product inhibition and degradation.

#### Modification of electro-membrane configuration

In his patent, Datta<sup>153</sup> designed an efficient and potentially economical process for lactic acid production. This process was configured to have desalting ED, water splitting electrodialysis (WSED), and ion-exchange purification steps. It was able to produce concentrated lactic acid with less than 0.1% of proteinaceous impurities from carbohydrate fermentation. In this process, there was no by-product gypsum and only a small amount of by-product salt was obtained from the ion-exchange regeneration. In addition, this process could be operated continuously, and thus can be scaled up for large-volume production.

In 1997, an EDBM plant for the recovery of organic acids was constructed in France.<sup>154</sup> This plant produces 2600 ton organic acids per year with 98% purity. The process requires 0.88 kW h kg<sup>-1</sup> of energy to produce acids. Generally, EDBM has the main problem of intolerance to multivalent cations such as  $Ca^{2+}$  and  $Mg^{2+}$ , which form insoluble hydroxides at the critical interface of the bipolar membrane where the ions separate. The concentration of divalent cations is usually limited to about 1 ppm, while fermentation broths mostly contain multivalent

 Table 3
 Performance of electro-membrane processes for the recovery of organic acids

Organic acid	Technology	Operation condition	Results	Ref.
Acetic acid	ED	Membrane effective area: 10 cm <sup>2</sup> Voltage: 5, 10, 15 V	Current efficiency: 80–98% Recovery rate: 24.05–40.82% (configuration 1) and 91–394%	131
		-	(configuration 2)	
	EED	Membrane effective area: 32 cm <sup>2</sup>	Removal efficiency: >90%	113
	EDBM	Membrane effective area: 98 cm <sup>2</sup>	Acid recovery: up to 70%	132
		Acid concentration: 0.2%	Current efficiency: 40%	
Amino acid	ED	Membrane effective area: 36 cm <sup>2</sup>	Recovery rate: up to 63%	133
		pH: 12.5	Current efficiency: up to 83%	
		Acid concentration: 25, 50, 75, 100 mM	Energy consumption: 3 kW h kg $^{-1}$	4.2.4
	EDBM	Lysine concentration: 146.19 M	Acid concentration increase 35–50 times	134
		Histidine concentration: 174.21 M		
		Membrane effective area: 20 $\text{cm}^2$		
		Current density: $2-10 \text{ mA cm}^{-2}$		
	EDBM	Methionine concentration: 24.65 g $L^{-1}$	Acid purity: 99.98%	135
	LDDM	Membrane effective area: $945 \text{ cm}^2$	Current efficiency: 75.10%	100
		Current density: $150-300 \text{ mA cm}^{-2}$	Energy consumption: 2.156–3.265 kW h kg $^{-1}$	
	EDI	Membrane effective area: $50 \text{ cm}^2$	Current efficiency: 60–85%	127
		Voltage: 20 V	Energy consumption: 1–1.6 kW h kg <sup>-1</sup>	
		Resin filling: 0, 5, 10, 20 mL		
Butyric acid	ED	Current: 0.05 A	Acid purity: 85%	129
			Current efficiency: 52%	
	EDI	Membrane effective area: 10 cm <sup>2</sup>	Recovery rate: 82–89%	128
		Voltage: 3.6–4.7 V	Current efficiency: 81–85%	
		pH: 2-4.5	Energy consumption: 0.197–0.204 kW h kg $^{-1}$	
	EDI	Membrane effective area: 0.001 m <sup>2</sup>	Acid purity: 92%	129
		Current: 0.05 A	Current efficiency: 59%	
Citric acid	ED	Membrane effective area: 3900 cm <sup>2</sup>	Optimum p: 7.5	136
		Voltage: 10, 15, 20 V	Optimum voltage: 20 V	
		Flow rate: 4, 8, 12 mL min $^{-1}$	Optimum flow rate: 4 mL min <sup>-1</sup>	
	EDDM	pH: 2, 4.5, 7.5 Mombrane officiative error $50 \text{ cm}^2$	Final said concentration, up to 0.05 mol $dm^{-3}$	107
	EDBM	Acid concentration: 0.1 mol $dm^{-3}$	Final acid concentration: up to 0.65 mol dm	137
		Actu concentration: 0.1 mor din Current density 52, 78, 104 mA $\text{cm}^{-2}$		
	FDPM	Membrane effective area: 220 cm <sup>2</sup>	Current efficiency 72 7-100%	120
	EDBM	Current density: 30, 40, 50 mA cm $^{-2}$	Energy consumption: $4-8$ kW h kg <sup>-1</sup>	130
	FDI	Membrane effective area: 50 $\text{cm}^2$	Current efficiency: 40-96%	7
	EDI	Acid concentration: 500–10 000 ppm	Energy consumption: 1.16 kW h $kg^{-1}$	/
		Flow rate: $1-4 \text{ L} \text{ h}^{-1}$	Energy consumption. 1.10 kw if kg	
Formic acid	EED	Temperature: 30 °C	Concentration ratio: 1.2–1.5	111
i oiiiiie ueiu		Current density: $8-29 \text{ mA cm}^{-2}$	Current efficiency: 70–140%	
	EED	Membrane effective area: 7.07 $\text{cm}^2$	Current efficiency: >100%	110
		Current density: 3, 5, 10, 15 mA $cm^2$	Formic acid transport through the membranes: 0.004–0.120 M	
		Temperature: 20–40 °C	1 0	
		Acid concentration: 0.05–0.5 M		
		Anolyte and catholyte concentration: 0.1 M		
Fumaric	EDBM	Membrane effective area: 0.0064 m <sup>2</sup>	Recovery ratio: up to 75%	139
acid		Current density: 90, 120, 150 mA $cm^{-2}$	Current efficiency: 80–90%	
		Acid concentration: $1.45$ – $2.90 \text{ g L}^{-1}$	Energy consumption: 5–13 kW h kg <sup><math>-1</math></sup>	
		Circulation flow rate: 6.2 L $h^{-1}$		
Glyceric acid	ED	Membrane effective area: 550 cm <sup>2</sup>	Acid recovery: 75–95%	140
		Acid concentration: $32.3-130.2 \text{ g L}^{-1}$	Current efficiency: 87.2–100%	
		pH: 7	Energy consumption: 0.19–0.31 kW h kg $^{-1}$	
Lactic acid	ED	Membrane effective area: 58 and 180 $\text{cm}^2$	Current efficiency: 45–83%	141
		Voltage: 1.5 V	Final acid concentration: up to $157 \text{ g L}^{-1}$	
		Current: 1.4 A	Energy consumption: 0.26–0.87 kW h kg <sup>-1</sup>	
	ED	Membrane effective area: $200 \text{ cm}^2$	Acid recovery: 96.4–98.7%	3
		Acid concentration: 80–100 g dm <sup>-3</sup>	Current efficiency: 77.3–83.0%	
	ED	Current: 8–12 A	Energy consumption: $1.2-3.8$	07
	ED	Feed: 0.1 N lactic acid and 0.1 N sodium sulfate	e Final acid concentration: up to $4 \text{ g L}^{-1}$	97
	ED	Flow rate: 150 mL min	Ourset officiency (C. 0404	4.10
	ED	Memorane effective area: 5/.6 and 180 cm <sup>2</sup>	Current emciency: 66–84%	142

Table 3 (Contd.)

Organic aci	d Technolo	ogy Operation condition	Results	Ref.
		Voltage: 1.5 V	Final acid concentration: up to 173 g $L^{-1}$	
		Current density: 7.8 mA cm <sup>2</sup>	Energy consumption: $0.24-0.32 \text{ kW h kg}^{-1}$	
	ED	Voltage: 0–15 V	Acid recovery: 40–100%	16
		Acid concentration: 40 g $L^{-1}$	Energy consumption: 0.163–0.910 kW h kg <sup><math>-1</math></sup>	
	ED	Membrane effective area: 100 cm <sup>2</sup>	Acid recovery: up to 97%	143
		Acid concentration: 0, 1, 5 g $L^{-1}$	Energy consumption: 0.25 kW h kg <sup><math>-1</math></sup>	
		Voltage: 10, 15, 20		
		Temperature: 32 °C		
	ED	Voltage: 7 V	Mineral removal: 90%	144
		Current density: 10–300 A m <sup>-2</sup>	Energy consumption: 0.004–0.014 kW h kg <sup><math>-1</math></sup>	
	EIS	Membrane effective area: 25 cm <sup>2</sup>	Final acid concentration: up to 9 g $L^{-1}$	97
		Feed: 0.1 N lactic acid and 0.1 N sodium sulfate	e	
		Flow rate: 150 mL min <sup>-1</sup>		
	EDBM	Membrane effective area: 57.6 and 180 cm <sup>2</sup>	Current efficiency: 61–92%	142
		Voltage: 12 V	Final acid concentration: up to 173 g $L^{-1}$	
		Current density: 67.7 mA cm <sup>-2</sup>	Energy consumption: 0.84–1.38 kW h kg <sup>-1</sup>	
	EDI	Membrane effective area: 90 $\text{cm}^2$	Final acid concentration: up to 185 g $L^{-1}$	123
		Voltage: 0–70 V		
		Current density: $0-45 \text{ mA cm}^{-2}$		
		Acid concentration: 0–80 g L		
Malic acid	EDBM	Membrane effective area: $0.02 \text{ m}^2$	Current efficiency: 30%	145
		Acid concentration: 24.4 g L $^{-1}$	Energy consumption: 1.15–1.27 kW h kg	
D	FFD	Circulation flow rate: 300 L h		44.5
Propionic	EED	Membrane effective area: 20 cm <sup>-1</sup>	Current emclency: $90.5-99.2\%$	116
acid		Acid concentration: 40 g L Current density 70 mA $\text{cm}^{-2}$	Final acid concentration: 150 g L	
	EDDM	Current density: 70 mA cm Mombrane officiative area: 20 $\text{cm}^2$	Current officiency 05 0 1000/	110
	EDBM	Acid concentration: 40 ct $I^{-1}$	Einel acid concentration: $145 \text{ cm} \text{ s}^{-1}$	110
		Actu concentration: 40 g L Current density 70 mA $cm^{-2}$	Final acid concentration: 145 g L	
Salicylic aci	d fidem	Feed acid concentration: 1 M	Current efficiency: 90-00%	101
Sancyne aei		Free action concentration: 1 M Flow rate: 00 L $b^{-1}$	Final acid concentration: up to 45 M	121
		Circulation rate: 4.6 cm s <sup><math>-1</math></sup>	Fnerov consumption: $14-38$ W h m <sup>2</sup> kg <sup>-1</sup>	
		Current density: 30, 50, 75 mA cm $^{-2}$	Energy consumption. 14 56 within kg	
	EDBM	Membrane effective area: 7.07 $\text{cm}^2$	Current efficiency: 99.6%	146
	LDDM	Acid concentration: $0.05-0.4$ mol L <sup>-1</sup>	Energy consumption: 2.1 W h $m^2$ kg	110
		Current density: $14-50 \text{ mA cm}^{-2}$	Energy consumption. 2.1 () if in fig	
Succinic	ED	Membrane effective area: $178 \text{ cm}^2$	Final acid concentration: 63–77.6 g $L^{-1}$	147
acid		Acid concentration: 51.5 g $L^{-1}$	Current efficiency: 76.2–78.9%	
	ED	Membrane effective area: 64 cm <sup>2</sup>	Current efficiency: 15–25%	148
		Voltage: 0–20 V	Total carboxylate basis: 50–60%	
	EDBM	Membrane effective area: 80 cm <sup>2</sup>	Final acid concentration: 0.25–0.60 M	149
		Acid concentration: 0.05 M	Current efficiency: 90%	
		Circulation rate: 15 L $h^{-1}$	Energy consumption: 1–4 kW h kg $^{-1}$	
		Current density: 12.5, 25, 37.5 mA $cm^{-2}$		
	EDBM	Membrane effective area: 80 cm <sup>2</sup>	Current efficiency: 96.8%	149
		Current density: 12.5–37.5 mA $cm^{-2}$	Energy consumption: <4 kW h kg <sup>-1</sup>	
	EDBM	Membrane effective area: 207 cm <sup>2</sup>	Current efficiency: 75.4%	150
		Current density: 90 and 120 A $m^{-2}$	Energy consumption: 1.5–3.2 kW h kg $^{-1}$	
		Acid concentration: 43, 100, and 200 g $L^{-1}$		
	EDBM	Current density: 90 and 120 mA cm $^{-2}$	Current efficiency: 14.3–19%	151
		Acid concentration: 15 and 16.9 g $dm^{-3}$	Final acid concentration: 13 and 15.7 g $dm^{-3}$	
		Feed flow rate: 100 $dm^2 h^{-1}$		
Tartaric aci	d ED	Temperature: 25–40 °C	Current efficiency: 33–65%	152
		Feed: 10 kg m <sup><math>-3</math></sup> tartaric acid and 60 kg m <sup><math>-3</math></sup>	Final acid concentration: 170–300 kg m <sup><math>-3</math></sup>	
		glucose	Energy consumption: 5.103–12.103 kJ kg $^{-1}$	

cations with concentrations of up to 1000 ppm. Therefore, Datta and Henry<sup>11</sup> used desalting ED to remove the multivalent cations and concentrate the lactate salt, followed by treatment in a WSED unit with bipolar membranes to produce concentrated lactic acid and alkali for recycling. WSED is a general purpose unit operation for converting water-soluble salts to their corresponding acids and bases.<sup>155</sup> The results showed that 98–99% of divalent ions were rejected with high recovery yield (>95%) and low power consumption, giving approximately 0.33 kW h kg<sup>-1</sup> of lactate. However, when the feed solution contains

Technology Advantages		Disadvantages	
ED	Simple process, low energy consumption	Relatively low recovery rate	
EMT	Can realize double composition reactions	Acid requirement	
EIS	Can realize double composition reactions	Low current efficiency, acid requirement	
EED	High current efficiency, can be operated continuously	Low product concentration, low membrane stability and selectivity	
EDBM	High separation efficiency, high acid purity	Expensive bipolar membrane price	
EDI	High recovery rate, self-regeneration, can be operated continu	uously Requires pretreatment	

metal ions, such as calcium and magnesium, fouling occurs faster in WSED due to the precipitation of metal ions in the cation-exchange membrane.<sup>97</sup>

Another attempt was also investigated to increase the recovery efficiency by molding ion exchange resins into a porous resin wafer (RW) and inserting it into the electro-membrane stack.122,156,157 This modification is usually called resin wafer electrodeionization (RW-EDI). Compared to conventional EDI, RW-EDI provides simpler assembly and efficient operation. RW-EDI demonstrates a very stable removal rate due to the rigidity of the porous wafer. Besides, the presence of a resin wafer enhances the electrical conductivity in the solutions and allows thinner EDI systems at reduced energy consumption, and regeneration of the resin wafers can occur within the cell through water splitting.<sup>158</sup> Therefore, the use of resin wafer broadens the applications of EDI, such as for esterification,<sup>159</sup> enzyme-based conversion and recovery of organic acids,156 and capture of carbon dioxide from flue gas.<sup>160</sup> Datta et al.<sup>122</sup> reported that RW-EDI successfully removed greater than 99% and 95% of sulfuric acid and acetic acid, respectively, while sugar retention was greater than 98%.

Furthermore, Lopez *et al.*<sup>158,161</sup> used ionic liquids instead of a resin wafer. They used two types of ionic liquid, including 1ethyl-3-methylimidazolium-trifluoromethanesulfonate ([EMIM] [OTf]) and 1-butyl-3-methylimidazolium acetate ([BMIM][OAc]). By using ionic liquids as the wafer in EDI stacks, they were able to obtain a recovery rate of 99%, while the current efficiencies reached 37–90% with energy consumption rates of approximately 1.25–2.80 kW h kg<sup>-1</sup> acid recovered.

#### In situ separation processes

Many fermentation processes have low productivity and yield due to product inhibition or hydrolysis of the product by further catalytic reactions.<sup>162</sup> This problem can be solved by optimizing technological parameters, for example by keeping the dissolved product concentration as low as possible in the fermentation medium. Thus, the *in situ* separation process is promising for selectively removing the fermentation product from the vicinity of the biocatalyst as soon as it is formed and also increasing the efficiency of recovery processes.<sup>123</sup>

The *in situ* separation process is a combination of product formation and separation of organic acids. This process has been studied by several researchers with various schemes and designs. Arora *et al.*<sup>156</sup> compared two schemes of combined bioreactor and EDI. The first scheme was called side-stream bioreactor, while the second was named immobilized separative bioreactor, as shown in Fig. 3. In the first scheme, the reaction takes place in a traditional bioreactor containing biocatalysts (enzymes/cells), sugars, and the organic acid products all in solution, which is then fed to the EDI separation unit. In the second scheme, the biocatalyst is immobilized directly in the RW-EDI to enable simultaneous reaction and separation. The results showed that immobilized separative bioreactor had a stable reaction rate, while the reaction rate decreased as a function of pH in the side-stream bioreactor.

Gao et al.<sup>163</sup> studied the use of electrodialysis fermentation (EDF) for lactic acid production. In EDF, the fermentation broth is continuously fed to the ED to concentrate the organic acid. The solution with a high concentration of organic acid is collected as the product, while the solution with a low organic acid concentration is returned to the fermentation reactor. Using EDF, the acid productivity was 1.5 times higher than the convention fermentation-ED process. The yield also increased by above 30% and glucose transport decreased from 0.46 to 0.05. Hirata et al.<sup>164</sup> also used an EDF system equipped with a glucose concentration controller (GC controller). The GC controller was added to control the glucose concentration to be stable and low in the fermentation broth. Meanwhile, Danner et al.165 studied integrated continuous cell recycle cultivation using ultra-filtration membrane bioreactor (MBR) systems. The permeate from the MBR was streamed to an ED to recover and concentrate lactic acid. The recovery yield of lactic acid was stable at around 80% with the energy consumption of 0.49 kW h kg<sup>-1</sup> of lactic acid.

Kumar et al. used an electro-membrane reactor where ion substitution and separation of acids occurred simultaneously for the recovery of glutamic acid<sup>115</sup> and amino acid.<sup>166</sup> Their studies resulted in a high recovery of glutamic acid and amino acid close to 96% and 85%, respectively. Meanwhile, Liu et al.167 investigated the potential of an integrated EDBM and biochemical process named the microbial electrodialysis and chemical-production cell (MEDCC) to produce malic acid. In general, MEDCC has same operating system with EDF; however, MEDCC uses bipolar membranes, as shown in Fig. 3(d). Liu et al.167 showed that MEDCC could convert 0.3 M malate into 0.23 M malic acid with a low energy consumption of 0.34 kW h  $kg^{-1}$ . This energy consumption was only 10–30% of that in the reported EDBM. The same method was also used by Wang et al.<sup>168,169</sup> for the production of lactic acid. The productivity of lactic acid was 1.76 g  $L^{-1}$  h<sup>-1</sup> and the yield coefficient was 56.77%. In addition, Luo et al.<sup>170</sup> also used MEDCC for the production of citric acid. The maximum citric acid production of 0.443  $\pm$  0.096 M was achieved within 96 h operation. The



Fig. 3 Schematic of (a) a side-stream bioreactor, (b) an immobilized separative bioreactor, (c) the stack configuration for EDF, and (d) the stack configuration of MEDCC.

energy consumption was 0.81  $\pm$  0.03 kW h kg<sup>-1</sup>. Lu *et al.*<sup>171</sup> also used MEDCC to produce formic acid. The minimum electricity consumption to produce 0.34  $\pm$  0.04 kW h kg<sup>-1</sup> of formic acid in the MEDCC with 72 cm of the anode fiber length was only 3.1–18.8% of that in the EDBM.

In addition, Zhu *et al.*<sup>172</sup> developed a new of bioelectrochemical system called a microbial reverse-electrodialysis chemical-production cell (MRCC). This system was developed to produce acid and alkali using energy derived from organic matter (acetate) and salinity gradients (NaCl solutions representative of seawater and river water). A bipolar membrane was placed next to the anole to prevent Cl<sup>-</sup> contamination and acidification of the anolyte, and to produce protons for HCl recovery. The MRCC reactor produced electricity (908 mW m<sup>-2</sup>) as well as concentrated acidic and alkaline solutions, and therefore did not require an external power supply. The results showed that the acid and alkali-production efficiencies based on the generated current were  $58 \pm 3\%$  and  $25 \pm 3\%$ , respectively.

Fig. 4 shows a comparison of the energy consumption for various electro-membrane processes for organic acids, including separated downstream processes and *in situ* separation processes. It can be seen that the *in situ* separations such as EDF and MEDCC require much lower energy consumption, and thus are more promising to be developed on an industrial scale. However, the maximum current density of EDF and MEDCC is much lower than that of ED, which greatly limits their application in practice.

#### Integrated separation processes

Fermentation operates most efficiently and effectively at near neutral pH, and hence produces the acid salt instead of the acid

itself. Besides, the presence of impurities such as water, particles, microbial cells, unconverted carbon sources, and inorganic ions cannot be avoided. Therefore, the pretreatment of feed solution is important before pumping it into the electromembrane stack to ensure stable operation and process performance. Press filtration and UF membrane are the alternative technologies to remove particles or colloids (bacteria, cellulose, proteins, or suspended solid chemicals) and keep the electro-membrane stack free from serious membrane fouling.<sup>142,173</sup> Activated carbon can be used to clear up coloring matter to prevent fouling or the appearance of unfavorable byproducts.<sup>174</sup> Meanwhile, chelating resins are able to



Fig. 4 Comparison of energy consumption for various electromembrane processes for the production of organic acids.

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selectively remove multivalent inorganic ions (especially,  $Ca^{2+}$ ,  $Mg^{2+}$ ,  $Ba^{2+}$ ,  $Mn^{2+}$ ,  $Zn^{2+}$ ,  $Fe^{3+}$ , and  $Sr^{2+}$ ) to prevent scale formation.<sup>142,175,176</sup>

Boyaval *et al.*<sup>177</sup> produced lactic acid in a continuous process comprising three distinct operations in a bioreactor, UF module, and ED cell. UF recycled all or part of the biomass and separated low molecular weight metabolites, such as sodium lactate, which resulted from lactose fermentation. The product of UF was then extracted and concentrated continuously by ED. This process was able to yield lactate solution in a high concentration of up to 130 g L<sup>-1</sup>. Meanwhile, Sikder *et al.*<sup>71</sup> demonstrated that combined NF-EDBM in downstream purification is able to replace multiple purification steps with only two steps, while yielding monomer grade lactic acid from a mixture of unconverted sugars and lactic acid. NF also led to nearly total discoloration of the fermentation broth.

Bailly et al.<sup>176</sup> proposed a complete organic acid production using a membrane for clarification, concentration and conversion. The fermentation broth was first clarified by cross-flow MF. Divalent cations were then removed from the clarified broth since they act as a poison in the process. ED was further used prior to EDBM to increase the concentration of ionic species comprising the organic acid salt. Meanwhile, Norddahl<sup>178</sup> combined UF and a two-step ED process. The first step used conventional ED membranes, while the second ED process used a bipolar membrane. A bipolar membrane was applied to separate the salts formed into lactic acid, inorganic acids, and ammonium hydroxide solution. By combining UF and two-step ED, the overall recovery rate of lactic acid was quite high at about 85-90%. Madzingaidzo et al.174 also combined ED and EDBM. ED was used for sodium lactate purification, while the recovery of lactic acid was done using EDBM. Sodium lactate with a feed concentration of 125 g  $L^{-1}$  was concentrated by ED to a maximum of 150 g L<sup>-1</sup>. The lactate solution was then passed through EDBM to obtain free lactic acid. The EDBM unit was able to obtain 160 g L<sup>-1</sup> free lactic acid while color and other chemical impurities were significantly reduced. Later, further development based on this process involved adjustment of the pH to below the  $pK_a$ -value of lactic acid (3.86).<sup>179</sup> As a result, the free lactate ions combined with hydrogen ions to form lactic acid having no net electrical charge. In 2018, Prochaska et al.<sup>151</sup> proposed an integrated system consisting of UF, EDBM, and three steps reactive extraction for the removal of succinic acid. UF acted as pre-clarification to remove high molecular contaminants. Meanwhile, EDBM allowed the acidification of the broth to be eliminated. The succinic acid present in the aqueous stream after EDBM was then removed in a threestep reactive extraction at more than 90% efficiency.

The downstream process on the industrial scale is mostly more complicated than the laboratory scale. As example, lactic acid is separated and substantially purified from fermentation broths by several membrane-based unit operations. MF or UF are used for cell separation and recycle, while NF is essential for separation of the lactic acid from other broth components using low rejection (LR) membranes.<sup>180</sup> Meanwhile, to concentrate the lactate, RO or a combination of high rejection (HR) and low rejection (LR) NF can be used. Further, ED is added for the simultaneous separation and concentration of lactate. The lactate product of ED is still in the salt form, and hence, EDBM is required to form lactic acid in the acid form and allow the recycle of the alkali used for neutralizing the fermentation broth. By adding EDBM to the process, the alkali cost is minimized, while the waste products (*e.g.*, calcium sulfate) generated in conventional downstream processes for organic acids are eliminated.<sup>176</sup>

### Future prospect and challenges

The increasing demand for organic acids in various applications has led to further exploration for their efficient production. The development of recovery processes is one of the crucial approaches to obtain efficient organic acid production with high quality product. The separation methods that have been studied for the recovery of organic acids mainly include precipitation, extraction, adsorption, and nanofiltration. Unfortunately, the existing processes still have many obstacles that need to be addressed. Precipitation and liquid–liquid extraction need a large amount of chemical agents and energy. They produce a large amount of water effluent as well as solid residue and involve phase changes that lead to a degradation in the quality of citric acid. Meanwhile, adsorption has a short lifetime of adsorbents and low capacity.

Electro-membrane processes are the promising technologies to replace the conventional technologies for the organic acid downstream process. Electro-membrane processes such as ED, EMT, EIS, EED, EDBM, and EDI can recover organic acids from their fermentation broth without the introduction of salt or discharge. Besides, their energy consumption is much lower than conventional technologies. However, concentration polarization and fouling are still main issues in electromembrane processes. The concentration polarization is a result of the difference between the transport number of the ions in the membrane and in the feed solution. These phenomena then lead to an accumulation of ions on the membrane surface in concentrate compartment, while ion depletion occurs on the membrane surface in the diluate compartment.<sup>181</sup> As a consequence, the operating currents of electro-membrane processes are restricted by the occurrence of concentration polarization.

In addition, fouling also undeniably occurs during membrane operation. Fouling is associated with the accumulation of substances on the membrane surface or within the membrane pores.<sup>182</sup> Fouling not only leads to a decline in productivity, but also requires an additional energy supply to keep the membrane performance constant.<sup>183-186</sup> In electromembrane processes, fouling occurs due to the accumulation of colloids and organic substances. Colloidal fouling is mainly deposited on the AEM since most of the colloids treated by electro-membrane processes are negatively charged, which leads to interactions with positively charged ion-exchange groups of AEM. Meanwhile, organic substances stick to the surface of all types of membranes and/or lodge themselves inside the membrane. In colloidal fouling, the interaction of the foulant and membrane is only electrostatic in nature, while organic fouling on the IEM surface and inside the IEM may be due to electrostatic and hydrophobic interactions.<sup>187</sup>

Fouling and concentration polarization are inevitable but can be controlled. The usage of suitable strategies allows for a longer membrane life and lower operational costs. Various attempts can be undertaken to control fouling and concentration polarization, such as selection of appropriate pretreatment, modification of the membrane, and optimization of operation conditions by increasing the pH and cross flow velocity, or decreasing the initial flux. Selection of an appropriate pretreatment can be used to inhibit the fouling and concentration polarization by reducing the concentration of impurities in the feed water. Meanwhile, optimization of the operation can minimize the interaction between impurities/foulant and the membrane surface. As an example, an increase in cross flow velocity can reduce fouling due to the disruption of the fouling layer by a strong hydrodynamic shear rate.<sup>188</sup> In addition, the properties of membrane materials plays an important role in fouling formation. Hydrophobic membranes are more prone to fouling formation than hydrophilic membranes. Therefore, in several cases, hydrophilic modification is required. Hydrophilic modification can be conducted by blending the original polymer with hydrophilic materials or the addition of hydrophilic polymer layers on the active surface of the membrane.189-193

Furthermore, there is still a need for further research to develop processes that are simple to carry out and allow the purification of organic acids directly from the fermentation broth. The integrated process of fermentation and product recovery, which is usually called *in situ* separation, is an alternative to solve this problem. The integration of fermentation and the electro-membrane process can increase the recovery yield to above 30%, while decreasing the energy consumption by at least 90%. However, this separation process still has room for improvement to make it industrially feasible, including the development of membrane materials, process design, and optimization of operating conditions.

## Conflicts of interest

There are no conflicts to declare.

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