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### Introduction

Our way of living is strongly dependent on Earth's natural resources to fulfil our needs for energy and materials. There is growing concern that the emissions of greenhouse gases, most notably carbon dioxide  $(CO_2)$ , associated with the use of fossil resources will lead to climate change. Therefore, we need to substantially increase the share of renewable energy

Department of Chemical Engineering and Chemistry, Eindhoven University of Technology, Eindhoven, The Netherlands. E-mail: m.c.costa.figueiredo@tue.nl resources to cover our immediate energy demands. The production of chemicals is also mostly based on fossil resources, and their manufacture typically requires large amounts of energy. Thus, the chemical industry can also contribute to lower CO<sub>2</sub> emissions by introducing renewable feedstock and energy.1

Many different technologies are being explored to enable the envisioned transition to renewable energy with a focus on renewable electricity derived from solar and wind.<sup>2</sup> In the long term, carbon capture and sequestration and use (CCS and CCU)

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### Electrocatalytic synthesis of organic carbonates

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Organic carbonates are considered environmentally benign alternatives for various fossil-derived compounds used in the chemical industry. Replacing current costly and toxic production methods by greener alternatives offers opportunities to cover the increasing demand for these intermediates in a more sustainable manner. In this feature article, the prospect of electrochemical synthesis of organic carbonates is presented as an approach to use carbon dioxide and green electricity to arrive at such compounds. We explore the strengths and limitations of the different methods by looking into the electrode and electrolyte composition effects and operating conditions with a focus on the synthesis of dimethylcarbonate from methanol and either carbon monoxide or carbon dioxide. The proposed mechanisms are discussed in an effort to understand the underlying steps and their challenges. This review concludes with a perspective on the broader developments needed to turn the basic chemistry into a practical application.

ROYAL SOCIETY OF CHEMISTRY



combined with renewable energy can form the basis for the sustainable production of chemicals.<sup>3</sup> Based on the expected availability and low pricing of renewable electricity, electrochemistry can become a key enabling technology for such a sustainable chemical industry.<sup>4,5</sup> Electrochemical conversion routes are attractive for many reasons. The technology can in principle be scaled up relatively easily, and it provides opportunities to use a variety of feedstock including renewable biobased waste products and even  $CO_2$ , which can in principle be captured from the air. This strategy of turning "waste" into valuable products can reduce the carbon footprint of the chemical industry to zero, if not negative.

Organic carbonates are an important class of chemical intermediates, for which a transition of a fossil-based process to an electrochemical alternative involving renewable energy and the use of CO<sub>2</sub> is considered. Organic carbonates find widespread application as aprotic polar solvents, monomers in the manufacture of plastics and intermediates in the production of pharmaceutical, cosmetics and fine chemicals (e.g., glycols, purines, carbamates). They can also be used as fuel additives<sup>6</sup> and electrolyte solvents in lithium-ion batteries.<sup>7</sup> Moreover, their low toxicity and biodegradability make them useful intermediates for green chemistry applications.<sup>8</sup> Currently, the industrial synthesis of organic carbonates is environmentally stressing due to the use of toxic reactants (e.g. phosgene and carbon monoxide), harsh operative conditions and large amount of waste.9 Thus new approaches to manufacture these chemicals sustainably, such as electrosynthesis, are desired. By using electrosynthesis, not only renewable energy is directly used to create bonds, but also electrons are used instead of highly toxic reactants. Moreover, when designing a novel electrochemical process for organic carbonate synthesis, the use of  $CO_2$  instead of carbon monoxide (CO) can be considered.10,11



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is Marta Costa Figueiredo Assistant Professor of Electrocatalysis at Eindhoven University of Technology since April 2019. She obtained her PhD in electrocatalysis, science and technology in 2012 at the University of Alicante, Spain under the supervision of Prof. Juan Feliu. After that, she was a postdoctoral researcher at different Universities in Europe such as Aalto University (Finland), Leiden University and

University of Copenhagen. Before joining TU/e, Marta worked in the industry as Jr Scientist at Avantium (Amsterdam). In Eindhoven, her research is devoted to electrocatalysis and electro(catalytic) synthesis for sustainable processes and production of high value chemicals. The formation of organic carbonates from  $CO_2$  can be achieved *via* two pathways. The first pathway involves the direct reduction of  $CO_2$ , while the second one first reduces  $CO_2$  to COfollowed by organic carbonate synthesis.<sup>12</sup> Technology for the electrochemical  $CO_2$  reduction to CO is advancing very fast.<sup>13-15</sup> In this review, we will discuss both electrochemical pathways. We finalise with an outlook by discussing the challenges in this area.

#### Electrocatalytic synthesis

As a general introduction, Fig. 1 depicts several electrochemical cells involving three electrodes, used for lab-scale testing. The simplest electrochemical cell configuration is an undivided three-electrode cell, in which a cathode and an anode are immersed into a conductive solution containing the substrate(s). At the cathode, electrons are removed from the surface and transferred to the substrate, which results in its reduction. At the same time at the anode, an electron from the substrate is removed, resulting in its oxidation. The electrochemical reaction is driven by a power supply that applies a potential difference between the cathode and the anode (Fig. 1a). In electrochemical synthesis, typically only one of the reactions, anodic or cathodic, is investigated. For example, the synthesis of dimethyl carbonate (DMC) from CO<sub>2</sub> and methanol (CH<sub>3</sub>OH), which requires reduction, takes place in the cathodic compartment (Fig. 1b). On the other hand, to synthesise DMC from CO and CH<sub>3</sub>OH, electrooxidation is required, which takes place at the anode (Fig. 1c).

# Electrocatalytic synthesis of organic carbonates from CO<sub>2</sub>

Several researchers have reported electrocatalytic synthesis methods to obtain organic carbonates,<sup>16–25</sup> mainly targeting DMC. DMC is an environmentally benign alkyl carbonate, which is suitable for a wide range of applications. For example, it can serve as a methoxycarbonylation and methylation agent, replacing toxic and hazardous compounds such as phosgene, methyl halides, and dimethyl sulphate.<sup>26</sup> DMC can also replace *tert*-butyl methyl ether as an oxygenated fuel additive in gaso-line or diesel. Another application of growing importance is as electrolyte solvent in lithium-ion batteries.<sup>7</sup>

Industrially, DMC is produced *via* various routes, all of which are hazardous and energy-intensive due to the high temperatures and pressures required to drive the reactions. The most important commercial process to manufacture DMC is oxidative carbonylation of CH<sub>3</sub>OH with oxygen (R1).<sup>27</sup>

$$2CH_{3}OH + \frac{1}{2}O_{2} + CO \xrightarrow{Cu \text{ salt}} CH_{3}OCO_{2}CH_{3} + H_{2}O \quad (R1)$$

Given the interest in DMC and the environmental concerns of current production methods, substantial research is directed towards cheaper and less toxic alternatives.<sup>28,29</sup> Electrochemical approaches have been explored for this purpose with most efforts made for the reaction of  $CO_2$  with  $CH_3OH$  (R2).

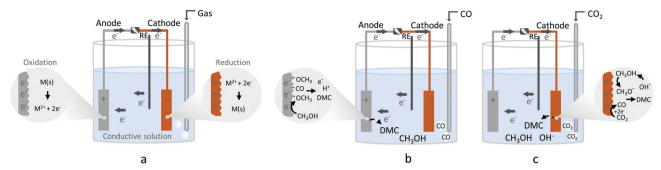


Fig. 1 Electrochemical three-electrode cells. (a) General electrochemical cell, (b) electrocatalytic synthesis of DMC with CO<sub>2</sub>, (c) electrocatalytic synthesis of DMC with CO.

This process involves different electrochemical and chemical reactions. The main electrochemical reactions are the reduction of  $CO_2$  (to CO) and water (R3) and (R4). The OH<sup>-</sup> generated from water reduction will facilitate the deprotonation CH<sub>3</sub>OH to form the methoxy intermediate (R5). DMC is then produced by reaction of the electrochemical formed CO and the methoxy species (R6).

$$CO_2 + 2CH_3OH + e^- \rightarrow CH_3CO_3CH_3 + \frac{1}{2}H_2 + OH^-$$
 (R2)

$$CO_2 + H_2O + 2e^- \rightarrow CO + 2OH^-$$
(R3)

$$2H_2O + 2e^- \rightarrow H_2 + 2OH^-$$
(R4)

$$CH_{3}OH + OH^{-} \leftrightarrow CH_{3}O^{-} + H_{2}O$$
 (R5)

$$CO + 2CH_3O^- \rightarrow CH_3CO_3CH_3 + 2e^-$$
(R6)

The majority of studies involving the direct electrochemical conversion of CO<sub>2</sub> to DMC are performed in imidazolium-based ionic liquids (ILs)<sup>30</sup> like 1-butyl,3-methyl-imidazolium tetra-fluoroborate and 1-ethyl-3-methylimidazolium tetrafluoroborate. These solvents are attractive because of the high CO<sub>2</sub> solubility compared to conventional solvents.<sup>16</sup> The solubility of CO<sub>2</sub>, expressed as the mole fraction of CO<sub>2</sub> in these imidazolium-based ILs are around 0.14 and 0.10, respectively, at ~300 K and 10 bar.<sup>31,32</sup> The corresponding solubilities in acetonitrile (<0.09) and methanol and ethanol (<0.07) are significantly lower under similar conditions.

Literature shows that DMC synthesis in ILs can be carried out with good reaction yields (ratio of the obtained product to the starting amount of alcohol). For example, yields around 75% were reported at In, Cu, and Ag based electrodes (Table 1, entries 9 and 7),<sup>23,24</sup> while a yield of 81% was mentioned for a nanoporous Cu–Pt based catalyst (Table 1, entry 10).<sup>21</sup> Despite these promising DMC yields, these systems use methyl iodide as an alkylating agent, which is a carcinogenic compound. Another study on Pt electrodes reported a high yield of 76%, using (carcinogenic) propylene oxide (Table 1, entry 8).<sup>22</sup> Various efforts to exclude the use of such harmful compounds and reduce the production steps have been reported,<sup>10,25,33–36</sup> which however negatively affect the DMC yield in general. For example, the synthesis of DMC using Pt or graphite working electrodes reported low yields, below 5%.<sup>33</sup> The authors proposed the reaction mechanism shown in Fig. 2a and suggested that the electrochemical synthesis of DMC involves activation of  $CH_3OH$  on the electrode surface as an initial step leading to the methoxide anion. Methoxide then reacts with  $CO_2$  to form DMC. These findings are contradictory to other reports that consider that  $CO_2$  is adsorbed on the electrode surface and interacts with  $CH_3OH$ .<sup>21,22</sup> Even though no additives were used in this study,<sup>33</sup> benzyl chloride is used to prepare the IL solvent, which is also classified as a carcinogenic compound and brings unnecessary risks and waste to the process.

Yuan et al. reported the electrochemical synthesis of DMC on Pt electrodes in a dialkyl imidazolium IL-potassium methoxide (CH<sub>3</sub>OK)-CH<sub>3</sub>OH system, where CH<sub>3</sub>OK acted as a cocatalyst.25 Amongst the basic compounds investigated (CH<sub>3</sub>ONa, NaOH, KOH, K<sub>2</sub>CO<sub>3</sub>, CH<sub>3</sub>OK), CH<sub>3</sub>OK led to the highest DMC yield (3.9%) (Table 1, entry 6). The authors suggested that the K<sup>+</sup> ion interacts with adsorbed CO<sub>2</sub> stabilising the  $CO_2^{-}$  anion (Fig. 2b). This system was further studied in a filter-press electrochemical reactor with a divided anodic and cathodic compartment, resulting in an increased DMC yield of 12.5% (Table 1, entry 2).<sup>34,35</sup> By applying a higher potential, the authors achieved the highest efficiency for the electrosynthesis of DMC from CO<sub>2</sub> without the use of harmful compounds (Table 1, entry 3). Another study by Lu et al. reported a yield of 2.5% with 94.5% selectivity on a graphite electrode using an amino-functionalised, imidazolium-based IL (Table 1, entry 4).<sup>36</sup> The basicity of the amino-functionalised IL is beneficial for capturing and activating the CO<sub>2</sub> molecule. These findings are in agreement with another work<sup>25</sup> and support that basic groups are advantageous for CO2 activation. Even though the use of carcinogenic compounds can be avoided, the use of ILs has some limitations. These include, besides the low yield, their high cost and viscosity.<sup>37</sup> Hence, a more cost-efficient and feasible system for the electrochemical conversion of CO<sub>2</sub> to DMC needs to be developed. To achieve this, understanding the underlying mechanism and identifying the reaction intermediates is essential.

A recent investigation of the synthesis of organic carbonates from  $CO_2$  and  $CH_3OH$  in acetonitrile provides better insight into the reaction mechanism and the associated intermediates (Table 1, entry 5).<sup>10</sup> The proposed reaction pathway (Fig. 3b)

Entry	Entry Catalyst	Electrochemical cell	l Electrolyte	Temperature (K)	ıre Potential	Current	Time	DMC yield <sup>a</sup>	DMC selectivity	DMC selectivity By-products
$1^{33}$	Graphite	Divided electrolytic cell	CH <sub>3</sub> OH & BzmimCl CH <sub>3</sub> OH & BmimCl CH <sub>3</sub> OH & BmimOH CH <sub>3</sub> OH & BmimBr CH <sub>3</sub> OH & BmimBr <sub>4</sub> CH <sub>3</sub> OH & EmimBr <sub>4</sub> CH <sub>3</sub> OH & EmimBr <sub>4</sub>	293	4.0 V (cell voltage)	164 to 68 mA cm <sup>-2</sup>	18.5 to 80 h	3.8% 3.7% 2.1% 3.0% 1.3% 6.0%	51.8% 12.9% 39.7% 15.5% 37.6% 24.3%	Dimethoxymethane (DMM), 2,2-dimethoxylpropane, 1,1,2,2-tetramethoxylethane, methyl 2-methoxylpropionate
	Pt Glassy carbon Cu		CH <sub>3</sub> OH & BzmimCl			No detection		1.9%	8.8%	
2 <sup>34</sup>	Pt/Nb plates	Divided electrochemical flow cell	CH <sub>3</sub> OK & BmimBr	279	3.5 V (cell voltage)	I	48 h	12.5% (based on CO <sub>2</sub> )	I	Dimethyl ether (DME), DMM and tetramethyl orthocarbo- nate (OC)
$3^{35}$		Undivided electrochemical flow cell			5.5 V (cell voltage)			$70.7\%$ (based on $CO_2$ )	I	OC, DMM, bromomethane
$4^{36}$	Graphite Pt Glassy carbon		CH <sub>3</sub> OH & BmimOH CH <sub>3</sub> OH & ApmimBr CH <sub>3</sub> OH & BmimBr CH <sub>3</sub> OH & BmimBr <sub>4</sub> CH <sub>3</sub> OH & ApmimBr	293	3.5 V (cell voltage)	I	40 h	0.9% 1.1% 0.4% 0.2% 0.3%	76.7% 94.3% 67.3% 53.1% 84.7% 90.7%	DMM
$5^{10}$	Cu Cu disc Pt disc Dd disc	Undivided electrolytic cell	Tetraethylammonium 298 tetrafluoroborate &	298	-1.2 and -2.4 V νs. Ag/AgCl	Ι	I	No detection No detection	Ι	Alkylammonium methyl carbonate
$6^{25}$	Pt Dt	Undivided electrolytic cell	CH <sub>3</sub> OH, CH <sub>3</sub> OK & hmimBr	303	5.5 V (cell voltage)		48 h	3.9%	88.4%	Trimethoxy methane, OC, H <sub>2</sub> ,
7 <sup>24</sup>	Cu Ni Ag Stainless steel		BmimBF <sub>4</sub> , CH <sub>3</sub> I, (alcohols added after electrolysis)	328	-1.8 V vs. Ag/AgI	I	Stirring for 6 h after electrolysis	74% 18% 73% 67%	100% 100% 100% 100%	No by-products
8 <sup>22</sup>	Ti Pt Glassy carbon Granhite	Undivided electrolytic cell	BmimBr, CH <sub>3</sub> OH & PO	303	-2.5 V vs. Ag/AgI 5.5 V (cell voltage)	I	48 h	58% 75.5% 30.3%	100% 77.8% 53.1% 37.1%	Not reported
$9^{23}$	In	Undivided	BmimBF <sub>4</sub> , CH <sub>3</sub> OH &	313	–1.7 V vs. Ag	Ι	DMC detected 6 h	76%		Not reported
$10^{21}$	Cu skeletons/ Pt shells Pt		BmimBF <sub>4</sub> , CH <sub>3</sub> OH & CH <sub>3</sub> I	298	—2.2 V уя. Ag	3.2 mA cm <sup><math>-2</math></sup> 1.7 mA cm <sup><math>-2</math></sup>		81% 64%	I	Not reported

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Feature Article

 $^{a}$  The reported yield is based on the starting alcohol unless stated otherwise.

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involves the electrochemical reduction of CO2 to CO and the simultaneous activation of CH<sub>3</sub>OH. The CH<sub>3</sub>OH is activated via dehydrogenation/deprotonation by OH- species derived from residual water oxidation. The CH<sub>3</sub>OH activation leads to the formation of methoxy species, which react with CO and form DMC and monomethyl carbonate (Fig. 1c). Under the applied conditions, the researchers did not detect DMC by Fourier transformed infrared spectroscopy (FTIR), as it quickly decomposes below -1.4 V vs. Ag/Ag<sup>+</sup> to the monomethyl carbonate as shown experimentally.<sup>10</sup> Fig. 3a shows the FTIR spectrum for a Cu electrode in the presence of CH<sub>3</sub>OH and CO<sub>2</sub>. The authors explained that the bands at 1673 and 1281 cm<sup>-1</sup> can be attributed to C=O and C-OCH<sub>3</sub> stretching vibrations of monomethyl carbonate, respectively, and not to functional groups of DMC. The formation of this alkyl carbonyl intermediate was observed on different electrodes like Cu, Pt and Pb. The results suggest that the overall reaction requires methoxy species and the ability of the electrode to reduce  $CO_2$  to CO or  $CO_2^-$ . Further experiments with isotopically labelled compounds showed that CO is responsible for the carbonylation step and not the CO<sub>2</sub> anion radical.<sup>25,36</sup>

So far, the studies discussed target the formation of DMC using  $CO_2$  directly as feedstock. As mentioned, higher yields were achieved with the use of ILs.<sup>21–24</sup> Investigations into avoiding the use of such media have succeeded in producing DMC, but with low yields.<sup>25,34–36</sup> A study from Figueiredo *et al.*<sup>10</sup> examined the formation of DMC in organic solvents. Despite the mechanistic insights obtained, DMC was not detected under the applied reaction conditions due to its low stability at negative potentials. As DMC can also be obtained under oxidative conditions using CO as a reactant, this approach will be discussed in the next section.

## Electrochemical synthesis of organic carbonates from CO

An alternative route to synthesise DMC electrochemically is by using CO under oxidative conditions. Through this route, CH<sub>3</sub>OH and CO are converted to DMC (R7), similar to the current industrial process. Other products often reported in the electrochemical equivalent of this process are dimethyl oxalate (DMO), dimethyl ether (DME), dimethoxymethane (DMM), and methyl formate (MF).  $CO_2$  has also been observed, which points to the undesired direct oxidation of  $CH_3OH$ .<sup>38</sup>

$$2CH_3OH + CO \leftrightarrow CH_3OCO_2CH_3 + 2H^+ + 2e^-$$
 (R7)

An interesting approach of CH<sub>3</sub>OH carbonylation with CO involved a solution-phase  $Br_2/Br^-$  redox couple in an indirect electrochemical system (Table 2, entry 2).<sup>39</sup>  $Br^-$  is oxidised at the Pd anode to  $Br_2$  and  $Br_2$  acts as an oxidant for the oxidative carbonylation of CH<sub>3</sub>OH and CO. This does not constitute an electrochemical reaction. Even though the system is very selective, it is not scalable due to the low yield (7.2%) and the use of  $Br_2$ , a highly toxic compound.

Otsuka et al. tested the activity of PdCl<sub>2</sub> and CuCl<sub>2</sub> anodes for the electrochemical carbonylation of CH<sub>3</sub>OH to DMC in a gas phase system (Table 2, entries 3 and 4).40,41 A higher applied voltage was required for CuCl<sub>2</sub> in comparison with PdCl<sub>2</sub>. This was explained by the involvement of Cu(II) in the carbonylation of CH<sub>3</sub>OH. Subsequent studies of the electrosynthesis of DMC over a PdCl<sub>2</sub>/vapor-grown carbon fibre (VGCF) anode and a CuCl<sub>2</sub>/VGCF anode were done by Yamanaka et al.<sup>42</sup> in a three-phase-boundary system (gas phase: CO, liquid phase CH<sub>3</sub>OH and solid phase: anode material). The results suggested that reduced Pd<sup>0</sup> species are responsible for catalysing CH<sub>3</sub>OH carbonylation from CO, whereas oxidised Pd species catalyse direct CH<sub>3</sub>OH electrochemical carbonylation to the DMO by-product. The CuCl<sub>2</sub>/VGCF anode displayed no activity when tested in the same system. The authors could not further explain this difference between Pd and Cu. The three-phase-boundary system method showed promising results with the PdCl<sub>2</sub>/VGCF anode with a selectivity of 82% and 67% current efficiency (Table 2, entry 5), obtained in CH<sub>3</sub>OH by applying a potential of +1.8 V vs. Ag/AgCl and feeding CO at 101 kPa.

Metal-ligand complexes were also studied for this reaction. CuCl-2,2'-bipyridine (bipy) complex was found to be the most efficient for the electrochemical carbonylation of CH<sub>3</sub>OH to DMC at room temperature and atmospheric pressure (Table 2, entries 6 and 7).<sup>43,44</sup> The drawbacks of this system are that the CuCl(bipy) complex has a low solubility in CH<sub>3</sub>OH and is unstable in air. Other Cu-based materials such as CuCl<sub>2</sub>(bipy)<sup>45</sup> and Cu-based metal-organic frameworks (MOF)<sup>46</sup> (e.g., Cu-2,2bipyridine-trimesic acid) have also been studied in a twocompartment electrolytic cell (Table 2, entries 8 and 9). Based on the results, the  $Cu^{2+}/Cu^{+}$  couple is thought to be solely responsible for catalysing the carbonylation reaction of CH<sub>3</sub>OH and 2,2'-bipyridine is only serving as a non-participating ligand. According to the proposed reaction mechanism ((R8)-(R11)), DMC is formed in solution, and the OH<sup>-</sup> groups from the electrolyte serve as co-catalyst by dissociating CH<sub>3</sub>OH.

Anode:

$$2\mathrm{Cu}^+ \to 2\mathrm{Cu}^{2+} + 2\mathrm{e}^- \tag{R8}$$

Cathode:

$$2CH_3OH + 2e^- \rightarrow 2CH_3O^- + H_2$$
 (R9)

Anolyte:

$$2CH_{3}OH + 2Cu^{2+} + CO \rightarrow CH_{3}OCO_{2}CH_{3} + 2Cu^{+} + 2H^{+}$$
(R10)

$$2CH_3O^- + 2H^+ \rightarrow 2CH_3OH$$
(R11)

This approach showed very high selectivity towards DMC without any detected by-products. Moreover, the results were obtained at low applied potential (+0.3 V *vs.* Ag/Ag<sub>2</sub>O) which makes this process interesting for further development (low energy required). According to the proposed reaction mechanism, the applied potential is essential for forming

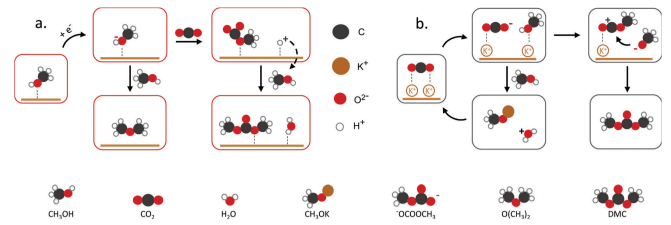


Fig. 2 Proposed mechanisms towards DMC. (a) DMC formation through methoxy and CO<sub>2</sub>, (b) K<sup>+</sup> interaction with CO<sub>2</sub> to DMC. Figures adapted from ref. 25 and 33.

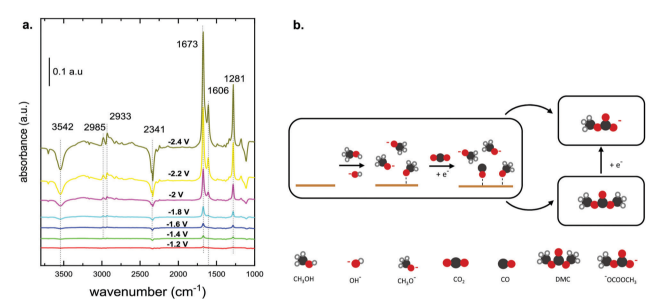


Fig. 3 (a) FTIR spectrum of  $CO_2$  reduction in  $CH_3OH$  on a Cu electrode at potentials between -1.2 and -2.4 V vs. Ag/AgCl.<sup>10</sup> (Reprinted with permission from American Chemical Society.) (b) Proposed reaction mechanism that involves the electrochemical reduction of  $CO_2$  to CO with a simultaneous activation of  $CH_3OH$ . Scheme adapted from ref. 10.

DMC, but its formation proceeds through an indirect electrochemical pathway.

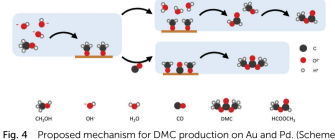
A corollary of the work mentioned above is that DMC selectivity depends strongly on both the reaction conditions and the presence of additives. Funakawa *et al.*<sup>47,48</sup> studied CH<sub>3</sub>OH carbonylation on a Au anode, in the form of Au/ activated carbon (AC) with vapor-grown carbon fibre (VGCF) and reported a strong dependence of selectivity on the applied potential (Table 2, entries 11 and 12). The reaction products were mainly DMO and DMC with DMM and MF as by-products. According to the authors, the DMO and DMC selectivity is controlled by the oxidation state of Au and hence the applied potential. However, the formation of DMC is through indirect electrochemical carbonylation of CH<sub>3</sub>OH, mediated by the Au<sup>3+</sup>/Au<sup>0</sup> (or Au<sup>+</sup>) redox reaction on AC at high potential

(>+1.3 V vs. Ag/AgCl). On the other hand, Au<sup>0</sup> was found to be the active species for the formation of DMO by direct electrochemical carbonylation at low potential (<+1.2 V vs.Ag/AgCl). Both suggested reaction pathways proceed *via* methoxycarbonyl intermediates. These observations agree with results of a density functional theory (DFT) study that showed that DMO is formed at lower potentials due to the presence of MF at the catalyst surface, whereas with increasing potential methoxy starts to adsorb favouring the formation of DMC.<sup>49</sup>

Figueiredo *et al.* reported for different electrode materials, including Au, Pd, Pt, and Ag, that DMC can be formed without any additives or oxidised metals (*i.e.*, formed at potentials below +0.4 V *vs.* Ag/AgCl) (Table 2, entry 14).<sup>50</sup> *In situ* FTIR was applied to detect intermediate products formed at different electrode potentials. The results revealed that DMC was only

 Table 2
 Electrocatalytic synthesis of DMC and other organic carbonates from CO

Futry	Entry Catalvet	Electrochemical cell	Flortrolvto	Temperature (K)	Dotential	Current (mA cm <sup>-2</sup> ) Time	) Time	DMC formation	DMC faradaic/ DMC		Buaroduote
	y valatyst			(v)	rucilual						by-produces
$1^{20}$	Cu	Divided H-cell	NaClO <sub>4</sub> /MeOH	298	0.2 V vs. SCF	4.5	70 days post 24 μM electrolvsis	24 μM	6%	100%	No other products detected
$2^{39}$	Pd/AC + VGCF	Pd/AC + VGCF Divided H-cell	LiBr & MeOH	298	$1.0 V \nu s$ .	9-10	60 min	150 $\mu mol~h^{-1}~cm^{-2}$	80%	95%	DMO, DMM, MF, and
$3^{40}$	CuCl <sub>2</sub> -	Divided flow cell	H <sub>3</sub> PO <sub>4</sub> & MeOH 34	343	Ag/AgCl 1.4 V vs.	3.2	120 min	0.1 $\mu$ mol min <sup>-1</sup> cm <sup>-2</sup> 11%	11%	Not reported	CO <sub>2</sub> Not reported DMM, MF, CO <sub>2</sub>
$A^{41}$	graphite PdCloranhite	graphite DdCloranhite Divided flow cell	HOAM/, Od. H	343	Ag/AgCl	6 6	30 min	0.1 ${ m umol}{ m min}^{-1}{ m cm}^{-2}$ 10%	10%	300%	DMM MF CO.
- 42	- 101 #100				Ag/AgCl	1	-	0 <u>1</u> -1 -2			
2.5	PdCl <sub>2</sub> /VGCF	Divided flow cell	MeOH/NaClO <sub>4</sub> 298 or HClO <sub>4</sub>	298	1.8 V νs. Ag/AgCl	œ	60 min	80 µmol h <sup>-+</sup> cm <sup>-2</sup>	67%	82%	DMO, DMM, MF, $CO_2$
6 <sup>43</sup>	CuCl(bipy)	Divided H-cell	MeOH/NaOCH <sub>3</sub> 293	293	0.3 V vs.	5	I	18 mmol	64%	Not reported	Not reported No other products
7 <sup>44</sup>	CuCl(bipy)	Divided H-cell	MeOH/NaOCH <sub>3</sub> 293	293	ысе 0.3 V VS. ссе	I	I		85%	Not reported	Not reported No other products
	$PdCl_2(bipy)$	Divided H-cell	MeOH/NaOCH <sub>3</sub> 293	293	эсе 0.9 V vs.		I		64%	Not reported	reported
8 <sup>45</sup>	CuCl <sub>2</sub> (bipy)	Divided electrolytic cell MeOH/KOH	MeOH/KOH	298	SCE 0.3 V vs.	2.3	120 min	0.3161 mmol	76%	100%	No other products
9 <sup>46</sup>	Cu(bipy) MOF	Cu(bipy) MOF Divided electrolytic cell MeOH/KOH	MeOH/KOH	298	Ag/Ag <sub>2</sub> O 0.3 V νs.	I	I	110.33 µmol	Ι	Not reported	detected Not reported No other products
$10^{47}$	$PdCl_2 + VGCF$	PdCl <sub>2</sub> + VGCF Divided H-cell	Н	298	Ag/Ag <sub>2</sub> O 1.0 V <i>vs</i> .	4.6	I	58.6 $\mu mol~cm^{-2}~h^{-1}$	68%	Not reported	reported Not reported DMO, DMM, MF, CO <sub>2</sub>
$11^{47}$		Divided H-cell	NaClO <sub>4</sub> /MeOH	298	Ag/AgCl 1.0 V vs.	2.4	I	2.3 $\mu$ mol cm <sup>-2</sup> h <sup>-1</sup>	5%	Not reported	Not reported DMO, DMM, MF, CO <sub>2</sub>
$12^{48}$	VGCF HAuCl <sub>4</sub> /AC	Divided H-cell	NaClO <sub>4</sub> /MeOH	298	Ag/AgCI 1.5 V vs.	8	250 min	300 µmol	20%	32%	DMO, DMM, MF, $CO_2$
$13^{49}$	Au	Divided H-cell	NaClO <sub>4</sub> /MeOH	298	Ag/AgUI 1.2 V vs.	9	20 min	430 µM	30%	Ι	MF, DMM, DME and
$14^{50}$	Au and Pd Ag and Pt	Three electrode spectro- LiClO <sub>4</sub> /MeOH electrochemical cell	- LiClO <sub>4</sub> /MeOH	298	SCE 0 to 1 V vs. Ag/				Not reported Not active	CO2 Not reported MF —	
$15^{53}$	þd	Divided H-cell	NaClO <sub>4</sub> /MeOH 29	298	Agu 1.2 V vs. scr	0.4	30 min	30 µM	15%	I	DMO, DMM, MF, $CO_2$
$16^{55}$	Au	Three electrode spectro- LiClO <sub>4</sub> /PhOH, electrochemical cell MeOH		298	эсь 0.5 to 1 <i>vs.</i> Ag/AgCl	I	I	I	Not reported	Not reported	Not reported DPC, Methyl phenyl carbonate (MPC)
<sup>a</sup> Th	e selectivity rep	$^{a}$ The selectivity reported is based on the conversion of starting alcohols.	version of startin	g alcohols.							



adapted from ref. 50.)

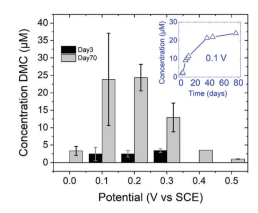
formed on electrodes that are capable of oxidising  $CH_3OH$  to (adsorbed) methoxy groups. Another requirement was that CO does not adsorb too strongly, because it would inhibit any further reaction. Ag and Pt were found to be inactive towards the formation of DMC. Pt suffers from CO poisoning, while the reaction on Ag is inhibited by  $CH_3OH$  adsorption.

On the other hand, Au and Pd, which bind CO weaker than the other metals, allow CO to act as a carbonylation agent to  $CH_3OH$  (Fig. 4). The importance of  $CH_3OH$  decomposition on the surface and the fact that DMC is formed through the methoxy intermediate was supported by FTIR spectroscopy and the formation of MF. The observation of MF as a by-product highlights the involvement of methoxy groups at the electrode surface in the overall reaction.<sup>51</sup>

Due to the technical limitations of *in situ* spectroelectrochemical FTIR, quantitative analysis cannot be easily performed. Thus, techniques such as gas or liquid chromatography are necessary<sup>52</sup> to analyse the product distribution in more detail. However, the application of these analytic techniques is limited by experimental aspects like clogging of separation columns by salts used as electrolytes to enhance conductivity.

Davies *et al.*<sup>53</sup> combined FTIR spectroscopy with headspace (HS) gas chromatography (GC) and mass spectrometry (MS) to achieve a quantitative analysis of the CH<sub>3</sub>OH carbonylation with a Pd electrocatalyst (Table 2, entry 15). This HS-GC-MS approach was applied in order to avoid salts entering the analysis system, as only gas-phase products enter the GC system.<sup>54</sup> The electrolyte solution was collected and analysed offline within 48 hours. The analysis time plays a crucial role in the analysis since it has been previously shown that in some systems DMC can be formed in the solution *via* an indirect electrochemical, homogeneous reaction.<sup>38,45,46,49</sup> Therefore, a shorter analysis time can improve the analysis of the results. The drawback of this analytical technique is that it can only detect products that are volatile below the boiling temperature of the sample matrix.

DFT studies on metallic surfaces<sup>38,49</sup> showed that Cu, different from Au and Pd, is an interesting candidate for the electrosynthesis of DMC, as it is predicted to require lower overpotentials. A good catalytic material would bind the methoxy group strongly enough in order to lower the overpotential needed for CH<sub>3</sub>OH oxidation and, at the same time, allow for an optimal coverage of CO. However, the binding of CO and methoxy



**Fig. 5** DMC concentration at 3 and 70 days after electrolysis. (inset) DMC concentration change over time since the day of electrolysis, shown for 0.1 V vs. SCE.<sup>38</sup> Reprinted with permission from American Chemical Society. Copyright@2019, American Chemical Society.

intermediates should not be too strong, otherwise the surface will be poisoned by one of these intermediates. With the help of FTIR, ICP-MS and HS-GS-MS, the authors<sup>38,49</sup> found that Cu does not act as a direct electrocatalyst for CH<sub>3</sub>OH carbonylation to DMC (Table 2, entries 1 and 13). According to spectroelectrochemical measurements, adsorbed methoxy, which is an intermediate for the direct electrochemical synthesis of DMC, was not observed.<sup>50</sup> On the other hand, a Cu carbonyl (Cu-CO) species was observed in the same potential range, in which DMC formation was seen. Besides, analysis of the electrolyte solution after the experiments did not show any DMC. However, after 70 days DMC was observed in the solution and its concentration increased over time. (Fig. 5) The authors proposed that the Cu(I) species act as the initiator of a post-electrolysis homogeneous reaction towards DMC, according to the mechanism ((R12)-(R14)), shown below.

$$\mathrm{Cu} + \mathrm{CO} \xrightarrow{E(0.0-0.4 \mathrm{V}_{\mathrm{SCE}})} [\mathrm{Cu}CO]^+ + \mathrm{e}^- \tag{R12}$$

$$[CuCO]^{+} + 2CH_{3}OH + \frac{1}{2}O_{2}$$
  

$$\rightarrow [Cu(OCH_{3})_{2}CO]^{+} + H_{2}O \qquad (R13)$$

$$[Cu(OCH_3)_2CO]^+ \rightarrow Cu^+ + CH_3OCO_2CH_3 \qquad (R14)$$

Based on the evidence that this reaction does not require an oxidised surface to obtain DMC,<sup>50</sup> the electrosynthesis of diphenyl carbonate (DPC) was investigated on Au under similar conditions (Table 2, entry 16).<sup>55</sup> DPC is also an important organic carbonate, because it has versatile chemical properties and can be used in synthetic chemistry as a phenylation and carbonylation agent.<sup>56</sup> Most importantly it is used for the solvent-free production of polycarbonates. Similar to DMC, DPC manufacturing procedures involve several steps and a large number of reactants.<sup>8,57</sup>

Employing *in situ* FTIR, DPC was identified at low overpotential (+0.5 V vs. Ag/AgCl) on Au electrodes via an indirect

electrochemical path. This pathway involves DMC conversion to DPC through an "electro-transesterification" step.55 The results suggest that DPC formation occurs similarly as DMC formation in which phenoxy groups take the role of the methoxy groups instead of a direct mechanism via phenol (PhOH) carbonylation. DMC is regarded as an electrophile and has three active centres for reaction with a nucleophile, *i.e.*, the carbonyl group and the two methyl groups. According to the HSAB (hard-soft acid-base) principle, hard nucleophiles prefer to react with hard electrophiles and soft nucleophiles with soft electrophiles. The carbonyl group is considered to be the harder electrophile due to the partial positive charge on the carbon atom and its  $sp^2$  hybridisation. On the other hand, the methyl groups are softer electrophiles due to their sp<sup>3</sup> hybridised orbital and saturated carbon atom.<sup>27</sup> Although alkoxides are considered hard nucleophiles, PhOH has a decreased nucleophilic nature due to the conjugation of the oxygen lone pair electrons with the aromatic ring when deprotonated, thereby rendering a softer base/nucleophile and facilitating the reaction with the methyl groups.

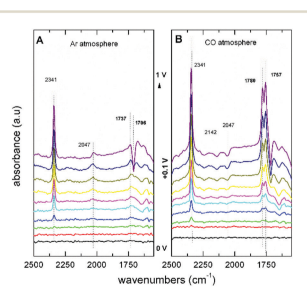
The first step in the proposed mechanism is the formation of DMC *via* carbonylation of the methoxy groups from CH<sub>3</sub>OH and CO, which are then replaced by the phenoxy groups. The application of an electrochemical potential was found to be essential for the nucleophilic attack to occur. FTIR spectra showed that the addition of PhOH does not change the reaction pathway of CH<sub>3</sub>OH oxidation to CO<sub>2</sub>, and no evidence of PhOH decomposition or oxidation products were observed. In Fig. 6A, the spectra under Ar saturation are characterised by bands at 2341 cm<sup>-1</sup> (CO<sub>2</sub>), 2047 cm<sup>-1</sup> (CH<sub>3</sub>OH), and 1737 cm<sup>-1</sup> (MF) related with the adsorption and oxidation of CH<sub>3</sub>OH. In spectra obtained after saturation with CO (Fig. 6B), two new positive bands are observed at potentials higher than +0.4 V vs. Ag/AgCl. These bands at 1757 cm<sup>-1</sup> and 1780 cm<sup>-1</sup> are attributed to DMC and DPC, respectively. These findings are relevant because they show that the direct DPC production by carbonylation of PhOH can be achieved with non Pd-based catalysts.

This research provides evidence that the transesterification of DMC with PhOH can occur under electrochemical conditions, avoiding thermocatalytic processes, and using the same catalyst as used to obtain DMC by electrochemical methods. Yet, more in-depth research into the nature of the "electrotransesterification" step is required.

### Future directions and outlook

A summary of the suggested reactions pathways and intermediates towards DMC and DPC (and the other major products) from  $CH_3OH$  and  $CO/CO_2$  is shown in Fig. 7.

The works cited demonstrate the substantial efforts made by the scientific community to achieve efficient electrosynthesis methods for organic carbonates, in particular those involving the electrochemical conversion of CO and CO<sub>2</sub>. While the understanding of the processes associated with CO<sub>2</sub> reduction reaction (CO<sub>2</sub>RR) has come a long way,<sup>58–62</sup> we need to understand this reaction much better as such and certainly in the context of the formation of organic carbonates. A major drawback remains the low yields of the desired organic carbonates, which requires high energy costs in downstream processing. Garcia-Herrero *et al.*<sup>63</sup> showed that the DMC yield in electrochemical synthesis from CH<sub>3</sub>OH and CO<sub>2</sub> must be at least 20% in order to compete with the commercial process. For these methods to become viable in the long term and to produce



**Fig. 6** FTIR spectra of an Au electrode in 0.1 M LiClO<sub>4</sub> in CH<sub>3</sub>OH with 0.1 M PhOH saturated with (A) Ar and (B) CO.<sup>55</sup> Reprinted with permission from American Chemical Society. Copyright©2019, American Chemical Society.

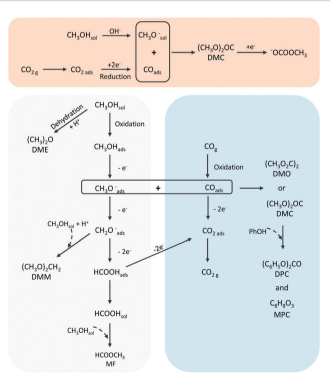


Fig. 7 Suggested reaction pathways and intermediates from CO $_2$  or CO and CH $_3\text{OH}.$ 

#### **Feature Article**

the targeted products on a global scale, different aspects such as thermodynamics, kinetics and infrastructure (reactors and process units), must be understood and optimised. In this review, we focused primarily on the electrocatalysts-solventconditions conducive to organic carbonate formation, which remains a key aspect to be improved before a process can be realized. However, also other aspects need to be addressed, such as stable membranes and electrodes, optimised electrolyser and environmental conditions (local pH, electrolytes). In the following, we briefly touch upon such aspects.

In terms of catalyst development, there are more than a handful of examples that can be obtained from other research areas as a source of inspiration. Catalyst design approaches that include alloying, dopants, tuning oxidation state, ligand and facet control can be used. For example, the addition of ligands has been implemented in previously mentioned works with the use of CuCl-2,2'-bipyridine (bipy) complexes,43-46 producing DMC with high selectivity showing that this strategy can be used for improved results. Catalyst morphology and structure are additional factors that can be adjusted to improve catalytic activity and selectivity towards the desired product.61 So far, to the best of our knowledge, no research has been done on the impact of the particle size, shape, exposed facets and defects of the nanomaterials (e.g. atom vacancies) or catalyst dopants for the electrochemical production of carbonates. One potential method is tuning the catalyst's oxidation state as it affects its local electronic structure. Oxide-derived materials, for example, can be used to tune the oxidation state of the catalyst. Enhanced performance of such electrocatalysts is attributed to the formation of active grain boundaries, edges, and steps that can have significantly different adsorption energies and interactions with the reactants and intermediates.<sup>64</sup> The use of bimetallic electrocatalysts is another viable strategy. Such catalysts have been widely investigated in the CO<sub>2</sub> reduction towards C1 and C2 products.<sup>62,65,66</sup> The surface interaction between two different materials results in altered electronic states and chemical properties, while at the same time, the local atomic arrangement can affect the binding strength of the reaction intermediates.

The lack of in situ analytical and catalyst characterisation studies for the electrochemical production of organic carbonates from CO<sub>2</sub> or CO limits the fundamental understanding of the reaction mechanism. This hinders improving the overall system and optimising the design of the catalyst. Furthermore, there is a lack of fundamental theoretical studies, which can help understand structure-performance relationships and design better catalysts.<sup>38,49,67</sup> For instance, modelling of the electrode-electrolyte interface can provide insight into the reaction kinetics and mass transport phenomena, which are of particular importance for continuous cell applications and porous electrode materials. Porous materials, usually prepared by catalyst deposition on a gas diffusion layer (GDL), have the advantage of supplying enough CO and CO<sub>2</sub> to the reaction zone, while enhancing a rapid electrolyte diffusion rate at the catalyst surface minimising mass transport limitations. Besides, GDLs act as low-resistance transportation media for protons, electrons, and reduction products from the catalyst to the electrolyte.<sup>68</sup> Catalysts assembled with GDLs have been widely used in  $CO_2$  electroreduction to  $C_{2+}$  products, and even though they are linked to high performances, stability is often problematic.<sup>69</sup> The reports of the use of GDLs for electrosynthesis of organic carbonates are, as far as we are aware, absent in the literature despite their proved advantages in reactions involving gaseous reactants. Their applicability, optimisation and stability should be explored in order to accelerate electrosynthesis as an industrial solution for the synthesis of organic carbonates.

Another aspect that should be further explored is the influence of the electrolyte. It is known that the use of ILs and organic solvents provide a higher CO<sub>2</sub> solubility and the absence of protons improves the current density while lowering the overpotential required for the CO<sub>2</sub> conversion.<sup>70</sup> The utilisation of mixed electrolytes, containing both organic and aqueous solvents, allows tuning the proton concentrations and consequently controlling the selectivity.<sup>71</sup> Additionally, mixing ILs with an organic or aqueous solution can reduce their viscosity<sup>37,72</sup> and therefore enhance the CO<sub>2</sub> diffusion rate, leading to better performance and cost-effectiveness. The used concentrations of ILs reported in literature vary from pure ILs to millimolar, but the optimum concentration has not been determined. Another alternative for making the overall process more cost-efficient, while maintaining the advantages of ILs, is the use of deep eutectic solvents (DES).73,74 DES have not been investigated in electrosynthesis of organic carbonates, but they can be used either as additives or as the electrolyte. These solvents are typically not consumed in the reaction, making them easier to recycle, reducing the operational costs of the process. However, their recycling might prove costly due to their viscosity and volatility.

The engineering aspects of the reactions should also be evaluated. Development of electrochemical processes requires significant capital investments, which even affects the scale-up of electrochemical reactions to a larger scale than that of the lab. Continuous flow microfluid cells are an emerging class of electrolysers that allow more effective scale up by numbering.<sup>75</sup> Such reactors can also have a positive impact on the overall efficiency of the reaction by improving mass transfer and reducing cell resistance, resulting in higher current densities.<sup>76</sup> The evaluation and optimisation of electrosynthesis of organic carbonates in continuous flow cells is a very promising research direction in our view.

Overall, this review sketched the basic electrochemistry of organic carbonate synthesis with an emphasis on potential electrocatalysts and electrolytes for obtaining dimethyl carbonate. Clearly, significant improvements are needed in terms of product yield to make a step towards implementing such technology at a larger scale. The main research questions to arrive at this stage were formulated in this work and can hopefully be an inspiration to others working in this field.

### Conflicts of interest

There are no conflicts to declare.

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