Utilization of carbon nanotube and graphene in electrochemical CO$_2$ reduction

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ABSTRACT

The electrochemical reduction of carbon dioxide (ERCO$_2$) driven by renewable energy to produce low-carbon fuels and value-added chemicals has been well known as a way capable of simultaneously solving energy exhaustion and global warming issue. Catalysts play a vital role in low temperature ERCO$_2$, and those well used are single metals, metal oxides and alloys. Due to the characteristics of nanometer size, low resistance, high surface area, chemical stability, special mechanical and electronic properties, some novel carbon nanomaterials (e.g. carbon nanotubes (CNTs) and graphene) show excellent properties in ERCO$_2$ as catalysts or supports which can improve the electrochemical performance: activity, selectivity, and stability. Actually, they mostly act as support materials and little directly as catalysts. The specific surface area and the active sites of loaded catalysts can be increased, then the performance is significantly improved. In this work, we will make a review on the progress as to CNTs and graphene as catalysts and supports in ERCO$_2$ in recent years and give the future prospects.

Keywords: CO$_2$; electrochemical reduction; carbon nanotube; graphene; catalyst; support.

1. INTRODUCTION

The gas emission caused by fossil fuel combustion has resulted in a remarkable carbon dioxide (CO$_2$) accumulation in atmosphere (as shown in Figure 1), followed by global warming. Meanwhile, fossil fuel reserves also fall rapidly. Therefore, it is vital to look for clean and sustainable energy such as solar, wind and tide energy. Conversion of CO$_2$ to valuable chemicals and low-carbon fuels is considered to be a promising way, because it may promote not only waste (gas) utilization but also energy storage. Among many available approaches (e.g. chemical, photocatalytic and electrocatalytic methods) to recycle the CO$_2$, the electrochemical reduction of CO$_2$ (ERCO$_2$) has a considerable potential economy and technical feasibility since it can convert those renewable energies and idle nuclear/water electrical energy into useful products.

As to the ERCO$_2$, there are basically three types of technologies which are applied at low temperature and high temperature [1-5]. In this work, we only focus on the low temperature process, which has obvious advantages over high temperature ones. However, low temperature ERCO$_2$ does require efficient and robust catalysts to lower the reaction overpotential and to minimize energy consumption. The lack of cost-effective catalysts with high activity, selectivity and stability is one of the biggest obstacles for realizing efficient ERCO$_2$ [2,3].

The carbon-based nanomaterials such as carbon nanotubes (CNTs) and graphene have been well used for energy storage and conversion as support material due to their nanometer size, low electric resistance, high surface area and chemical stability, and special mechanical and electronic properties [6]. After being loaded, the specific surface area and active sites of catalysts can be increased and the electrochemical performance is popularly improved [7-17]. Besides, both of them may also directly act as the catalysts of ERCO$_2$ [18-21]. Compared to other carbon materials like carbon fibers [22], boron-doped diamond [23-25], nanoporous carbon [26] and even graphene dots [27-29], which are also used for the catalysis of ERCO$_2$, CNTs [30-34] and graphene [35-39] have attracted more attention recently, both of which, therefore, will be discussed here.

2. CARBON NANOTUBES AS CATALYST SUPPORTING MATERIALS IN ERCO$_2$

CNTs are an allotrope of carbon with a cylindrical nanostructure and classified as single-walled carbon nanotube (SWCNT) and multi-walled carbon nanotube (MWCNT). As described above, CNTs can act as catalyst support owing to their extraordinary thermal conductivity, mechanical and electrical properties, and high surface area [6]. Moreover, the resistance to acid/basic media, the capability of controlling the porosity and surface chemistry within certain limits, nanometer size, low
Mn is an abundant and inexpensive mental element and its metal complexes are found to be effective in the application to ERCO reduction in aqueous solution. To improve the catalytic performance, Walsh et al. [43] developed a simple membrane supported manganese catalyst (e.g. Nafion/MWCNT/[Mn(bpy)(CO)Br]). The results showed that addition of MWCNTs led to a ~10 fold current enhancement (3 mA/cm² vs. 0.3 mA/cm²) at −1.4 V_Ag/AgCl at pH7. Recently, Sato et al. [51] prepared a new Mn complex electrocatalyst, [Mn(4,4’-di(1H-pyrrolyl)-3-propylcarbonate)-2,2’ -bipyridine)(CO)MeCN][PF_6] (Mn-MeCN) loaded on conductive MWCNTs ([Mn-MeCN]/MWCNT). Compared with the bare [Mn-MeCN] cathode which was unhelpful to CO₂ reduction reaction in an aqueous solution, the [Mn-MeCN]/MWCNT electrode could efficiently promote the ERCO to CO. The results showed that Mn-complex/MWCNT electrode could produce CO at a constant rate for 48 h and had a high current density of 2.0 mA/cm² at −0.39 V_RHE. Meanwhile the as-prepared electrode had a very low overpotential (ca. 100 mV) than previous reported Mn-complex catalysts.

**Figure 2.** Covalent immobilization of the iron porphyrin catalyst CAT_pyr on MWCNTs [56].

Fe, which is a more inexpensive metal material compared with Mn, has also been used to prepare the catalysts for ERCO. Hori et al. [52] had an early used Fe electrode to reduce CO₂ in aqueous KHCO₃ solution at −0.91 V_SHE. H₂ was measured to be the major product (FE_H₂=94.8%). After Fe being deposited on CNTs (the CNTs was annealed at 700°C in inert gas, and then treated in HNO₃ to introduce oxygen functionalities on the carbon surface) as the catalyst, it was found that FE_H₂ decreased to 80.6% and FE_CO increased to 18.9% [53]. Zhao et al. [54] prepared the electrodes modified with Fe porphyrin and CNTs (FeP-CNTs) for ERCO. Compared with the electrodes modified only with Fe porphyrin or CNTs, the FeP-CNTs electrode exhibited less negative cathode potential and higher reaction rate. The order of electrocatalysis efficiency was measured to be: FeP-CNTs > FeP > CNT. This high efficiency electrocatalysis of FeP-CNTs mainly came from two factors. First, the interaction between MWCNTs and Fe porphyrin reduced the overpotential to reduce CO₂ to formic acid. Second, the excellent conductivity of MWCNTs promoted electron transfer between CO₂ and the active site. Maurin et al. [55] prepared FeP-CNTs by using a different method: covalent grafting Fe porphyrin on CNTs. The FeP-CNTs had high selectivity and activity for the ERCO in water (pH=7.3) at low overpotential (0.5V). After 1h, the catalytic selectivity was 90% toward CO, and then it slightly decreased to 80% after 3h. Later, they used an improved Fe porphyrin catalyst (pyrene-appended iron triphenyl porphyrin bearing six pendant OH groups on the phenyl rings in all ortho and ortho’ positions, namely CAT_pyr) to prepare the CAT_pyr/MWCNTs catalyst. Figure 2 shows that CAT_pyr was immobilized on CNTs by noncovalent interactions. This new catalyst deposited onto a glassy carbon electrode reduced CO₂ to CO at low overpotential (480mV) in water (pH=7.3) and sustained for hours without loss of activity and selectivity [56]. Iron and nitrogen-doped carbon materials have recently been found to be cheap, stable and active alternatives. Jia et al. [57] prepared the Fe/Fe₃C@NCNT catalyst (Fe/Fe₃C nanoparticles embedded in nitrogen-doped CNTs), the highest FE_CO was 50% at −0.74 V_RHE in 0.5M NaHCO₃. The good performance of the Fe/Fe₃C@NCNT catalyst was due to a large specific surface area and high content existence of N with pyridinic-N and pyrrolic-N bonding.

**Figure 3.** (a) Structure of CoII(Ch) and (b) schematic image of CoII(Ch) on MWCNTs [59].

**Figure 4.** Schematic mechanistic pathway for the electrocatalytic production of formic acid, acetic acid and methanol on Cu10-CNT/GDL [64].
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Figure 5. The structure of the Cu/SnOx-CNT catalyst [65].

Co and Fe are in the same Group(VIII). Like Fe, pure Co was also investigated in ERCO₂ in aqueous solution. However, the study showed that Co macrocycles had high catalytic selectivity for the production of H₂ [58]. To enhance its selectivity of non-hydrogen products, Shoko et al prepared the Co⁴(CH) catalysts by adsorbing Co chlorin complex (Co⁴(CH)) on MWCNTs (Figure 3). The results indicated that the Co⁴(CH) catalysts could reduce CO₂ to CO in H₂O (pH=4.6) with a high F_EC(O) (89%) at −1.1 V_RHE, which may be attributed to the three dimensional assembly of MWCNTs with Co⁴(CH) on the electrode surface and the π-π interaction between MWCNTs and Co⁴(CH) providing a suitable hydrophobic environment for binding of CO₂ [59].

As one of the non-precious metals, pure Cu is also a promising catalyst for the ERCO₂. However, the previous works show that Cu has low product selectivity [60]. For example, using Cu electrode as catalyst in 0.1 M KHCO₃ aqueous solution (18.5±0.5°C) at −1.44V_RHE gave various products, and FE_{H₂} was high up to 20.5% [52,53]. Later, Baturina et al [61] supported Cu NPs on SWCNTs to prepare the Cu-SWCNTs catalyst. Compared with electrodeposited Cu electrode (Cu electrodeposition on Pt disk electrodes), the Cu-SWCNTs were more selective toward CH₄ generation, and had a significant (ca. 200 mV) shift in onset potentials, probably because of the increasing number of low-coordination sites (e.g. corners, edges, and defects) on the electrode surface. Hossain et al [45] used a series of CNT-supported Nano Cu (3-60nm) to catalyze the ERCO₂. The catalysts were prepared by homogeneous deposition precipitation method with urea as precipitating agent. When 20 wt. % copper was loaded on CNTs, the catalysts showed the maximum activity owing to the best balance of the amount of metal active sites and metal particle size. The CO₂ reduction had a high current density at the range of 0−3 V_RHE, and FE_{CH₃OH} was 38.5%.

Koo et al [62] electrochemically deposited Cu on aligned CNT sheets to prepare the oxygen-plasma-treated Cu-CNTs (O-CNT/Cu) electrocatalysts. The products were CO and methane, and the yields were 178 µmol cm⁻² mA·h and 346 µmol cm⁻² mA·h, respectively, and higher than CNT/Cu and CNT sheets. Marepally et al [63] used the Cu nanowires (NWs) as the catalyst precursor for the deposition Cu NPs onto CNTs to prepare the Cu NPs/CNTs catalyst by different methods, e.g. (1) the conventional impregnation route (ImR) to prepare the CuCNT-ImR catalyst; (2) use of Cu NWs as a copper precursor to prepare the CuCNT-NW catalyst. The main products were formic acid and CH₃COOH. Compared with the o-CNT (preliminarily activated with HNO₃ to create oxygen functional groups on their external surface) and CuCNT-ImR, the production rate toward the formic acid of the CuCNT-NW catalyst was 3 times higher than them. Later, Genovese et al [64] also deposited CNTs and Cu NPs on a gas diffusion layer (GDL) to prepare the Cu-CNT/GDL catalyst, and studied the mechanism. The Cu-CNT/GDL electrode could reduce CO₂ to CH₃COOH with a high FE_{CH₃COOH} (56%) at room temperature and atmospheric pressure, and the only other products of reaction detected were formic acid and methanol (the latter in some cases), besides H₂. The mechanistic was described in Figure 4. After that, the researchers began to deposited Cu metal with other metals on CNT. Genovese et al [53] found that the FeCu-CNTs electrode could double the productivity to C₁-C₃ hydrocarbons/organics with respect to Fe monometallic electrocatalysts. Also, the deposition of Cu metal with other metal oxides loaded on CNT was researched. Huo et al [65] supported Cu and SnOx on the CNTs to prepare the Cu/SnOx-CNT catalyst (Figure 5). When the ratio of Cu and SnOx was different, and the ERCO₂ products formed were also different. That was when the catalyst contained 6.2% SnOx, CO₂ could be reduced with a high FE_{CO} (89%) and the current density was 11.3 mA/cm² at −0.99 V_RHE, when it contained 30.2% SnOx, CO₂ could be reduced with a high FE_{HCOOH} (77%) and the current density was 4.0 mA/cm² at −0.99 V_RHE.

Sn is one another of the widely-studied catalysts for the ERCO₂. In the early works, the Sn bulk was used to reduce CO₂ to produce formate. With the development of electrocatalysts, the researchers introduce the concept of the porous electrocatalysts on high surface area conductive support to make the performance of catalyst better. Chen et al [66] reported a 3D hierarchical porous structured CNT aerogel-supported Sn spheroidal particles (Sn/CNT-Ags) on carbon cloth (Sn/CNT-Ags/CC) electrode. The Sn/CNT-Ags electrochemically reduced the CO₂ to formate with high selectivity. The highest FE_{formate} was 82.7% at a low overpotential of 361 mV, and the current density was 32.9 mA/cm². Compared with the Sn/CNT catalyst (CNT-supported Sn spheroidal particles), the Sn/CNT-Ags catalytic performance was greatly enhanced. The reason was that Sn/CNT-Ags catalyst had a high specific surface area, superior conductivity and excellent 3D hierarchical structure. Figure 6 exhibits typical SEM images of Sn/CNT-Ags (a and b). Sn metal oxides supported on CNTs are also investigated as catalyst for electrochemical CO₂ reduction. Bashir et al [46] reported the SnO₂/MWCNT electrocatalysts for CO₂ reduction in an aqueous electrolyte solution. It was found that when the content of SnO₂ was 20%, the catalysts showed the best performance, and could reduce CO₂ with a high FE_{formate} (27.2%) as well as a high current density of 80 mA/cm² at −1.7 V_RHE. The good performance was attributed to high electrical conductivity and high surface area of the MWCNT, which contributed to obtain more active sites and disperse evenly at the electrode surface.

Re and its complexes have received wide attention due to its good catalytic properties. Rezaei et al [67] prepared the [ReCl(CO)₅(tpzH)] electrocatalyst (where tpz-H = 2,4,6-tri(pyridine-2-yl)-2H-1,3,5-triazine-1-ide) for homogeneous and heterogeneous reduction of CO₂ in solution and on a carboxylated MWCNT modified pencil graphite electrode (PGE), respectively, to study the electrochemical performance. Compared to the homogeneous catalysis, the heterogeneous one showed a higher cathodic current and a lower over potential (about 650 mV_RHE). A direct reason for the performance difference in the two situations came from the presence or absence of MWCNT capable of increasing the surface area of an electrode and fast electron transfer kinetics for the CO₂ reduction process. The Re(Bu-bpy)(CO)₃Cl is another complex that has been widely studied as a homogeneous catalyst in organic solvents. When the Re(Bu-
Bi is a very promising material for the selective reduction of CO₂ to formic acid in 0.5M KHCO₃ aqueous solution with the property of nontoxicity, negligible environmental impacts, and inexpensive price [69-71]. When combining bismuth with CNTs to prepare the catalyst, the catalytic performance can be greatly improved than pure Bi. Recently, Li et al [72] prepared the Bi-MWCNT-COOH/Cu catalyst by constant current electrodeposition Bi-MWCNT-COOH composite on Cu foil substrate. The as-prepared electrode showed excellent activity with high FE\textsubscript{formate} reaching a value 91.7% at −0.76 V\textsubscript{RHE} better than Bi/Cu electrode without Bi-MWCNT-COOH. In addition, the catalyst showed good stability. After 12h continuous electrolysis, the FE\textsubscript{formate} and current density were basically unchanged. These good performances could be attributed to the special properties of CNTs, which increased the specific surface area of the catalyst, the number of reactive sites and the conductivity of the catalyst.

Unlike the above non-precious metals, Pd is a precious metal element. But, due to the special outer-electron structure and the excellent hydrogen absorption capacity, Pd NP and its nanocomposite may be highly effective electrocatalysts for CO₂ reduction. A few noticeable studies have proved that. For example, Pd-MWCNTs catalysts prepared by Lu et al [40] reduced the CO₂ to formic acid and CH₃COOH with a high FE\textsubscript{formate}=34.5%, FE\textsubscript{CH₃COOH}=52.3% at −4 V\textsubscript{Ag/AgCl} in KHCO₃ electrolyte. Zhao et al [41] fixed Pd NPs on polyamine-covered MWCNT surfaces to prepare the Pd-polyamine/CNT (Pd-PANI/CNT) catalysts by using an in situ method in an ambient alenvironment. The Pd-PANI/CNT catalysts reduced the CO₂ into formate with a high FE\textsubscript{formate} (83%) at −0.8 V\textsubscript{SCE}. Proposed mechanism for CO₂ electrocatalytic conversion to formate on Pd-PANI/CNT catalysts could be seen in Figure 7. The excellent catalytic performance of two catalysts mentioned above had a very important relationship to that the Pd particles were highly dispersed in carbon nanotubes with amorphous structure thereby increasing reactive sites.

Ir is a precious metal material, and has high activity for ERCO₂. It is a commendable candidate to reduce CO₂ to the formate with a high selectivity. Researchers prepared the electrode by immobilizing Ir pincer catalyst on CNTs. They found that the electrode showed notable efficiency, selectivity, and longevity. The current density was 15 mA/cm² in water (0.1M NaHCO₃, 0.5M LiClO₃, 1% MeCN v/v), and the highest FE\textsubscript{formate} could reach as high as 96% [73].

Pt, which is also a precious metal, has inferior catalytic performance for CO₂. Hori et al [52] found that Pt was a good catalyst for H₂ evolution reaction (FE\textsubscript{H₂}=95.7%). However, it was found that after Pt been deposited on oxygen functionalized CNT, hydrogen evolution was inhibited slightly (FE\textsubscript{H₂}=89.8%). Based on this finding, Jimenez et al [42] prepared the Pt-CNT catalyst depositing Pt on CNT by using the supercritical media (supercritical CO₂). When electrochemically reducing CO₂ in gas phase, the products were formic acid (59~89%), methane (2~33%), CO (3~11%), methanol (0~1.9%), and small amounts of acetone, isopropanol, and methyl acetate. Researchers have further found that low temperature contributed to the production of methane (33%), and the high temperature was conducive to the formation of formic acid (89%).

Ag can electrochemically convert CO₂ to CO with high selectivity and current density. The development of Ag catalysts has been carried out around organometallic Ag catalysts, Ag particles with different particle sizes and so on [74]. The electrode most commonly used in the study of the catalysts is gas diffusion electrode (GDE). Because of MWCNT unique properties, it is potentially suitable as support materials for electrocatalysts. Ma et al [75] prepared two kinds of Ag-MWCNT catalysts by using one-step method (1) the Ag NP catalyst and MWCNTs were homogeneously distributed (“mixed” catalyst layer), (2) a layer of MWCNTs was covered with a layer of Ag catalyst (“layered” catalyst layer). Two kinds of catalysts both could reduce CO₂ into CO with a good performance, but the “mixed” layer performed best. The “mixed” layer could reduce CO₂ into CO at −3 V\textsubscript{RHE} with a high current density (350 mA/cm²), FE\textsubscript{CO₂} (>95%), and energy efficiency (45%). Compared with catalyst without MWCNTs, the catalysts prepared above both had a better catalytic performance in the ERCO₂. And a lower charge transfer resistance was responsible for the more superior electrode performance of the “mixed” catalyst layer Ag-MWCNT catalysts.

Au as same as Ag has a decent performance for ERCO₂ to CO. However, on account of aggregation, Au catalyst lacks stability. To reduce the aggregation of the Au NPs and to enhance
the use of precious metals, Au NPs are often loaded on carbon materials (CMs), including CNTs, carbon black and so on, which have a high specific surface area. Feng et al [76] used vapor deposition to prepare Au/CNT catalyst, they found that the prepared catalyst showed high activity for CO production. After a 12h electrolysis in 0.5M NaHCO₃, the FE₇CO was 94% at −0.50 V_RHE. Huan et al [77] anchored Au on CNTs by a layer-by-layer method, so only a few Au were used. Compared with the monodisperse Au electrode, the mass activity of the Au/CNTs was 33 times higher at −0.55 V_RHE, and the FE₇CO reached 70%. Moreover, surface functionalized CNTs are also used in Au-based catalysts and exhibit excellent performance. For example, Ma et al [78] artfully prepared a hybrid catalyst. Au NPs supported on pyridine functionalized oxidized CNTs (Au/Py-CNTs-O). And the Au/Py-CNTs-O catalyst showed good mass activity ten times than Au/CNTs-O, high FE₇CO as high as 93% and enduring stability with little changed FE₇CO and slight decrease in current density in the process of 10h test.

3. GRAPHENE AS CATALYST SUPPORTING MATERIALS IN ERCO₂

Since Andre Geim and Konstantin Novoselov successfully separated the sp² hybrid graphene from graphite in 2004 for the first time, graphene has been widely studied and applied. It is a 2D crystal material with a honeycomb structure formed by a compact arrangement of carbon atoms in a single layer. The special structure endows graphene excellent electrical, thermal, mechanical, and optical properties [80,81]. In recent years, researchers found that graphene has huge value in the field of electrochemical CO₂ reduction. The applications of graphene and derivatives as catalyst and support material in ERCO₂ are introduced below.

As described before, Sn has the highest chemical reactivity and selectivity in ERCO₂. However, it is less stable under the ambient conditions. In order to protect Sn from the oxidation, the researchers put forward a sandwich-like structure model (ultrathin metal layers confined in graphene). For example, Lei et al [16] synthesized the Sn-graphene catalyst (Sn quantum sheets confined in few-layered graphene) with the same structure and found that the Sn-graphene catalyst had more active sites to adsorb CO₂ (9 times higher than the bulk Sn). The Sn-graphene could reduce CO₂ to formate with a high current density of 21.1 mA/cm² at −1.8V_SCE, being 2, 2.5 and 13 times larger than the 15nm Sn NPs mixed with graphene, 15nm Sn NPs and bulk Sn, respectively. The FE_formate was 85%, during the 50h electrolysis. Several advantages of ultrathin metal layers confined in graphene for the ERCO₂ to hydrocarbon fuels are depicted in Figure 8. Considering the nature advantages of Sn foil, which could be massively produced, directly used without complicated synthetic processes, and easily controlled to alter the size and thickness, Huang et al [82] prepared a single layer nitrogen-doped graphene (NG) coating Sn foil (SL-NG@Sn) catalyst. The experiment results showed that SL-NG@Sn foil had good flexibility and could effectively reduce CO₂ to formate with outstanding FE (92.0%) at −1.0 V_RHE in 0.5M KHCO₃ and high formate partial current density of 21.3 mA/cm², superior to other reported Sn NPs-based catalysts, which was attributed to the synergistic effect between SL-NG and Sn foil.

Pd nanocatalysts themselves also can catalyze the ERCO₂ to produce CO [83-87], formate [88-90] and even CH₃OH [90]. To reduce costs, Pd and Cu mono- and bi-metal NPs loaded graphene catalyst were prepared using co-reduction methods by Liu et al [91]. The results showed that the 1 wt% Pd-2 wt% Cu/graphene exhibited the best performance. The Pd/Cu-graphene catalyst had a relative positive peak potential and reduction current and the values were −1.3 V_{Ag/AgCl} and −2.8 mA/cm², respectively, in CO₂ saturated KHCO₃ solution. Tao et al [87] recently reported a Pd/Te/FLG catalyst (FLG is few-layer graphene) for the ERCO₂ to produce CO, which doping Pd with Te enhanced the performance of the catalyst. When the ratio of Pd and Te was 1:0.05, FE₇CO displayed the maximum 90% at −0.8 V_RHE and CO partial current density were 4.4 mA/cm².

There are many problems that hinder the development of the Cu-based catalyst for ERCO₂, such as, instability, poor selectivity and large reaction overpotentials. Considering the particular nature of graphene as mentioned above, it has been used as support for Cu-based catalyst. Cu is often used with rGO for the CO₂ reduction. Alves et al [92] fabricated the Cu/rGO catalysts for CO₂ reduction. The result showed that compared with other Cu-based electrode, the Cu/rGO had a higher current density, a lower overpotential and better stability: at −1.2 V_RHE the current density observed in the Cu-NPs/rGO electrode was −0.24 mA/cm², while the same current density was reached at −1.54 V for Cu thin film. The current density obtained in the Cu-NPs/rGO at −1.54 V_RHE was −0.97 mA/cm², four fold higher than the Cu thin film. The Cu-NPs on rGO also exhibited better stability, preserving their catalytic activity without degradation for several hours. Recently, Hossain et al [93] reported a unique nanocomposite consisting of Cu NPs and rGO supported on a Cu substrate with high catalytic activity for CO₂ reduction. The optimized nanocomposite could effectively reduce CO₂ to CO, formic acid and methane with a FE of 76.6% at −0.4 V_RHE in a CO₂
saturated NaHCO$_3$ solution. The superior electrocatalytic activity and stability of the Cu-rGO nanocomposite achieved could be attributed to the uniformly distributed small Cu NPs on the rGO and the synergistic coupling effect of the formed nanocomposite. The electron transfer between the rGO and Cu NPs may increase localized electron concentrations, resulting in significant enhancement of the catalytic activities of the nanocomposite for the electrochemical reduction of CO$_2$.

Sb is rarely used for ERCO$_2$ perhaps due to that the catalytically active sites are hidden in the material. However researches find that Sb may be a suitable catalyst for the ERCO$_2$. Li et al [94] recently prepared a few-layer Sb nanosheet-graphene (SbNS-G) composite to reduce CO$_2$. They experimentally found that there was a strong electronic interaction between graphene and Sb, and that the SbNS-G had the capability to catalyze CO$_2$ to formate with high selectively and efficiency. The maximum FE$_{\text{formate}}$ reached up to 88.5% at $-0.96$ V$_{\text{RHE}}$ in 0.5M NaHCO$_3$ solutions. At the same condition, the SbNS electrode reduced CO$_2$ at $-1.06$ V$_{\text{RHE}}$ with a FE$_{\text{formate}}$ being 84%. Besides, the current density of the SbNS-G electrode was 1.5 and 1.6 folds higher than SbNSs and bulk Sb.

The same as Sb, Au is also rarely used in ERCO$_2$. One reason is that Au is a precious metal with expensive cost price. And the other reasons as described above, Au catalyst is lack of stability. Graphene, like carbon nanotubes, can improve the catalytic performance of the Au catalyst. Rogers et al [95] prepared a GNRAuNP catalyst (gold nanoparticles embedded in a bottom-up synthesized graphene nanoribbon (GNR) matrix). The electrochemical studies showed that due to the structural and electronic properties of the GNR, the electrochemically active surface area was increased, and the overpotential of the CO$_2$ reduction was decreased. The onset potential of the GNR-AuNP catalyst was $-0.2$ V$_{\text{RHE}}$, the FE$_{\text{CO}}$ was 90%, and the stability was more than 24 h. Compared with the traditional amorphous carbon AuNPs supports, the catalytic output was increased by 100 times. Saquib et al [96] synthesized Au NPs on the rGO support (Au-rGO) and studied electrochemical performance of it. Through analyzing the values of electrochemical and impedance measurements, they found that Au-rGO had a better catalyst performance than bare Au for CO$_2$ reduction. The experiment also proved that the content of Au in the Au-rGO catalyst would affect the catalytic performance of the catalyst. When the Au content was less than 48.56%, the increase of content, onset potential for CO$_2$ reduction was decreased. When the content was more than 48.56%, onset potential for CO$_2$ reduction had almost no change. For example, when the content of the Au was 35.51%, the current density was 50.95 mA/cm$^2$ and was the 1.65 times higher than that of rGO Au (13.72 wt%). In this work, they found that the defect sites of the rGO played a vitally significant role in CO$_2$ reduction by reducing the overpotential.

Mo is earth-abundant composition and has high d-electron density near the Fermi level, which is suitable active sites for CO$_2$ reduction. Based on this found, scarce single Mo atom loaded ultrathin NG (Mo@NG) catalyst was conducted by Huang et al [97]. The as-prepared electrode proved to be effective to reduction CO$_2$ toward formate in 4 mol% ionic liquids (ILs) with formate yield of 747 mmol/(g$_{\text{cat}}$/h), which was superior to conventional NG. The one of effects of NG in the catalyst was to offered proper anchor sites for stabilizing the single atom and relative endurance for a broader negative potential region thereby improving the catalyst performances for CO$_2$ reduction.

Graphene materials are also used as bi-metal based catalyst carriers. For example, He et al [98] prepared the metal/3D-rGO electrode for CO$_2$ reduction. The results showed that Pd$_{0.5}$–In$_{0.5}$/3D-rGO catalyst had the best performance. The highest FE$_{\text{formate}}$ was 85.3% at $-1.6$ V$_{\text{Ag/AgCl}}$ in 0.5M KHCO$_3$ higher than the Pd$_{0.5}$/3D-rGO (78.6%) and In$_{0.5}$/3D- rGO (63.1%) electrode. The FE$_{\text{formate}}$ could be kept for 24h without obvious attenuation.

Graphene is also doped with heteroatoms to alter the surface properties and used as a catalyst carrier for CO$_2$ reduction. For example, the researchers prepared a copper NP/NG electrode consisting of Cu NPs loaded on the NG (it presents a surface of intense folds and spikes, so it called the carbon nanospikes or CNS) to reduce CO$_2$ to CH$_3$CH$_2$OH. The FE$_{\text{CH}_3\text{CH}_2\text{OH}}$ was 63% at $-1.2$ V$_{\text{RHE}}$ in 0.1M KOH. The current density of the CNS/Cu was 5 times higher than the bare CNS and 3 times higher than the Cu [99], which was attributed to the synergistic interaction between Cu and neighboring CNS. Li et al [100] assembled 7nm Cu NPs on the p-NG (pyridinic-N rich graphene) at the mass ratio of 1:1 to prepare the p-NG/Cu catalyst. The p-NG itself could catalyze the CO$_2$ reduction to formate, but in the composite p-NG-Cu structure, the pyridinic-N functioned as a CO$_2$ and proton absorber, facilitating hydrogenation and carbon–carbon coupling reactions on Cu for the formation of C$_2$H$_4$. The p-NG–Cu catalyzed the reduction of CO$_2$ to formate (62% FE, and 97% hydrocarbon selectivity) at $-0.8$ V$_{\text{RHE}}$ but to CH$_3$H and CH$_4$ (3% FE, 2.9A/gCu mass activity, and 79% hydrocarbon selectivity over other hydrocarbons) at $-0.9$V$_{\text{RHE}}$, achieving much superior selectivity towards C$_2$H$_4$ to any other Cu-catalysts reported previously. The data showed that the combination of p-N in the p-NG and Cu NPs of a proper size (7 nm in this work) facilitated CO$_2$ reduction, hydrogenation and C–C coupling for the formation of C$_2$H$_4$.

CO$_2$-based catalysts are often used in water oxidation, O$_2$ reduction reaction, H$_2$ evolution reaction. Recent studies showed that ultrathin layers of CoO$_x$ had high CO$_2$ reduction activity and selectivity. But there were disadvantages such as poor electron conductivity, limited surface area, and inefficient electron transfer and so on. Sekar et al [101] prepared a nanohybrid catalyst made up of loading of CoO$_x$ supported on NG (NG-CoO$_x$). The NG-CoO$_x$
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electrode showed a higher catalytic activity compared with undoped graphene and the difference mainly came from the significant effect of NG able to obtain a controlled morphology and to facilitate a topotactic transformation during electrocatalytic conditions to CoO which resulted to be the true active phase. By the way, some cobalt complexes are also an effective composition that can be used for ERCO₂. For example, Wang et al [102] prepared a heterogeneous catalyst consisting of planar Co(II)-2,3-naphthalocyanine (NapCo) complexes loaded on doped graphenes including graphitic sulfide dopant and carboxyl dopant. The experimental results showed that NapCo supported doped graphene matrices had better good catalyst performance for CO₂ reduction to CO than Pristine NapCo without graphene supports under the identical conditions in aqueous solution. And the sulfide dopant was superior to carboxyl dopant with a FE for CO production up to 97%, coming from the reason that the sulfide dopant could further improve the electron communication between NapCo and graphene. Su et al [103] synthesized novel Ni-N-modified graphene (Ni-N-Gr) by the short-duration heat treatment of a Ni-N organo-metallic complex in the presence of GO. The HER activity of Ni-N-Gr was much lower than that of Ni-metal electrode, which was likely responsible for the high efficiency of CO formation. Compared with N-Gr and the Ni metal electrode, Ni-N-Gr exhibited a higher activity for CO formation with a high FE (90%) at −0.7 to −0.9 V_RHE. The structure of Ni-N-Gr can be seen in Figure 9.

4. CARBON NANOTUBES AND GRAPHENE FAMILY

In recent years, the good chemical properties and low price of metal free catalysts have attracted the attention of the researchers. Compared with the metal-containing catalyst, the metal-free catalyst can be the next-generation, renewable materials [18].

4.1. CNT-based catalysts.

Pristine CNTs themselves actually cannot catalyze the ERCO₂ only if they are nitrogen-doped.[26,57,79,104-106] Most works have shown that NCNTs can reduce CO₂ to CO. For example, Sharma et al [105] found that the presence of graphitic and pyridinic nitrogen significantly decreased the overpotential (ca. φ0.18 V) and increased the selectivity (ca. 80%) towards the formation of CO. They prepared a series of NCNTs catalysts by using different precursors (e.g. ACN, DMF, and TEA) and various growth temperatures, 750, 850, and 950°C. The NCNTs-ACN-850 (synthesized with ACN precursors at 850°C) catalyst showed the best performance. Xu et al [106] also prepared the NCNTs catalyst by a different method. MWCNTs were firstly oxidized by HNO₃ at 120, 130 or 160°C to get three types of oxidized CNTs (OCNTs). Then the washed and dried OCNTs were treated by adding poly(diallyldimethylammonium chloride) (PDDA) solution (35 wt%) with vigorous stirring, resulting in three composites (e.g. PDDA@OCNT-1, PDDA@OCNT-2 and PDDA@OCNT-3) After being filtered and washed with distilled water and ethanol and then dried for 12 h at 80°C in vacuum, they were treated via the plasma treated NCNTs.[35] By the way, some cobalt complexes which has a good electrochemical performance for CO₂ reduction. The graphene has a similar chemical structure to CNTs, making it equally attractive for catalytic applications. Graphene, because of its planar structure, provides a platform for a systematic independent study of catalytic activity on different N-defect structures inside carbon network.[35] Importantly, nitrogen-doping and curvature can effectively tune the activity and selectivity of graphene/CNT catalysts.[35] Therefore, the researchers began to study the catalytic performance of N-graphene to CO₂ reduction.

Wu et al [35] reported a 3D graphene foam which had nitrogen defects as a metal-free catalyst for CO₂ reduction (Figure 11). The nitrogen-doped 3D graphene foam required low onset overpotential (−0.19 V_RHE), and showed superior activity over Au and Ag. It reduced CO₂ to CO with a high FEₐₐₚₐₜ (85%) at −0.47 V_RHE and long stability for at least 5 h.

4.2. Graphene-based catalysts.

Like NCNTs which has a good electrochemical performance for CO₂ reduction. The graphene has a similar chemical structure to CNTs, making it equally attractive for catalytic applications. Graphene, because of its planar structure, provides a platform for a systematic independent study of catalytic activity on different N-defect structures inside carbon network.[35] Importantly, nitrogen-doping and curvature can effectively tune the activity and selectivity of graphene/CNT catalysts.[35] Therefore, the researchers began to study the catalytic performance of N-graphene to CO₂ reduction.

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Sun et al [107] used NG-like materials (NGM)/CP as electrodes to reduce CO$_2$ to methane in ionic liquids. The metal-free electrodes showed excellent activity and selectivity ($F_{\text{CH}_4}=93.5\%$). Moreover, with the increasing content, the $F_{\text{CH}_4}$ was increased. At the same time, the addition of water in the ionic liquid also had an effect on the reaction. When 3 wt% water was added in ionic liquids, the current density had double (1.42–3.26 mA/cm$^2$). Compared with Cu electrodes, this NGM-CP electrode showed a 6 times higher current density.

Wang et al [36] also synthesized the NG catalyst by the pyrolysis of the graphene oxide (carbon template) and melamine. When the content of the N was 5.5 atom%, the catalyst reduced CO$_2$ to formate in KHCO$_3$ electrolyte with a higher $F_{\text{formate}}$ (73%). When the applied electrode potential was $-0.84$ V$_{\text{RHE}}$, the stability was 12 h, and the current density was 7.5 mA/cm$^2$. The researcher tried another method to improve the performance of the catalyst. Soliman et al [108] described a one-pot, bottom-up, pathway for constructing a novel composite material of a pyrimidine-containing porous organic polymer (PyPOP) atop graphene (PyPOP@G) sheets for application in CO$_2$ capture and electrocatalytic reduction. The PyPOP demonstrated an appreciable affinity toward CO$_2$ capture but was found to be largely insulating, hindering its usage in potential ERCO. However, its composite with graphene was found to be microporous, with a maintained affinity toward CO$_2$ and further demonstrated significant electrochemical activity toward CO$_2$ reduction (5 mA/cm$^2$ at $-1.6$ V), not observed in either of its two components separately. The one-pot, bottom-up, synthesis of the PyPOP@G is described in Figure 12.

**Figure 11.** Schematic of N configuration and CO$_2$ reduction pathway.

Wu et al [27] prepared the nanometre-size NG quantum dots (NGQDs) used for the ERCO$_2$ into value-added chemicals at high FE, high current density and low overpotential. The NGQDs showed a high total FE up to 90% at $-0.75$ V$_{\text{RHE}}$, and had a high selectivity for ethylene and ethanol conversions (45%). The C$_2$ and C$_3$ product distribution and production rate of NGQDs catalyzing reduction of CO$_2$ was similar to that of Cu NP-based electrocatalysts. And the partial current density for production of CO, C$_2$H$_4$ and C$_2$H$_5$OH (23, 46 and 21mA/cm$^2$ at $-0.86$ V$_{\text{RHE}}$) were similar to the commercial Cu.

Sreekanth et al [109] prepared a metal-free B-doped graphene (BG) to reduce CO$_2$ to formate in 0.1 M KHCO$_3$. Figure 13 display that compared with the Bi electrode, the B-doped graphene electrode showed a better catalyst performance for the reduction of CO$_2$ in 0.1 M KHCO$_3$. DFT calculations maybe can account for fact that boron-doping in graphene introduced asymmetric spin density, making it suitable for ERCO$_2$ by adsorbing on BG and not on pristine graphene and thereby undergoing reduction to formate.

**Figure 12.** The one-pot, bottom-up, synthesis of the PyPOP@G. C (orange), N (blue), H (white), G (green hexagons).

**Figure 13.** (a) LSV of BG (red lines) and Bi (dark lines) in 0.1 M KHCO$_3$ in the presence of CO$_2$. (b) SG-TC mode: cyclic voltammetric response of the Pt UME tip probe to the product generated from the B doped graphene (red) and Bi (black) kept at a substrate potential (Es) $-1.4$ V in 0.1 M KHCO$_3$ saturated with CO$_2$; scan rate 0.05 V/s.[109]

5. SUMMARY, CHALLENGE AND PROPOSED RESEARCH DIRECTIONS

This paper introduces the application of CNT and graphene in the electrochemical CO$_2$ reduction in recent years. Because these two kinds of Carbon Materials (CMs) have the characteristics of nanometer size, high surface area and chemical stability, which has aroused the extensive attention of the researchers. Generally speaking, CMs generally have two effects in electrochemical CO$_2$ reduction: (1) supporting material. This takes advantage of the good conductivity of the carbon material and the better dispersion of the metal particles on it. (2) Catalyst. CMs used as catalysts are usually doped with N or B elements. Recently, scientists have found that when carbon nanotubes or graphene are synthesized, the metal (Ni, Fe and Mn) residue on CNT and graphene will affect the catalytic performance of CMs. When the residual metals on the carbon material are washed with ultrapure nitric acid, the activity of the carbon material for the reduction of CO$_2$ will be greatly reduced.

There are still several technical challenges in the application of CMs for electrochemical CO$_2$: (1) Low CO$_2$ solubility. The low solubility of CO$_2$ is an urgent problem, which affects the efficiency...
of the electrochemical CO₂ by reducing the adsorption degree of CO₂ on the surface of the electrode; (2) Low catalyst activity. It can be seen from the above that for every single type of catalyst developed in the literature, the overpotential for CO₂ electroreduction is normally too high, indicating that these catalysts’ activities are still not good enough for practical applications in terms of energy efficiency; (3) Low product selectivity. For the majority of the carbon-based catalysts discussed above, even if the activity is high, the selectivity is not good enough. In addition to the target product, there will be hydrogen and other products. And some carbon-based catalysts which have a high selectivity are suffered from low stability. So, if we want to promote the large-scale application of electrochemical CO₂, we can change the way of application of the product. For instance, if the products are H₂ and CO, we can change the ratio of H₂ and CO and use the product as syngas.

Because of the good performance of CNTs and graphene, they have been widely developed in recent years. A large part of them is not used in the field of electrochemical reduction CO₂. For example, (i) different forms of CMs: the catalytic performance of the catalyst is affected by the morphology. The morphology of the catalyst is different, and the catalytic performance will vary. Fiber-like CNT, direct spun CNT fiber, 3D graphene. (ii) The properties of different elements doped CMs are also different. In addition to the common nitrogen-doping, there are Fe-N-doped CNTs, B-O-dually doped MWCNTs, N-S-coded graphene and so on. (iii) Generally, alloy materials will improve the properties of metal materials. The catalysts which are synthesized with alloy and CMs can improve the catalytic performance of the catalyst. (iii) Recent studies have found that some organic compounds, such as MOF, have high activity for the reduction of CO₂. But the organic compounds itself has no good electrical conductivity. The catalytic performance of organic compounds can be further improved if the catalyst is synthesized with CMs and organic compounds.

6. CONCLUSIONS

ERC0₂ to value-added chemical and low carbon fuels has been recognized as an economic, environment-friendly and easily scale-up technology to efficiently deal with the problems of energy dilemma and greenhouse effect. For this purpose, efficient and robust catalysts will play important roles in getting down the reaction overpotential and reducing energy consumption in the process. In this paper, we review the applications of CNTs and graphene as support materials and even catalysts for ERC0₂, which have attracted wide attentions in the past decades. When these two novel carbon materials acted as catalyst supports, more perfect performances of catalysts, such as higher FE, product selectivity and stability for ERC0₂, were popularly achieved compared with that of simple catalyst materials. It can be attributed to nanostructures and characteristics of CNTs and graphene, which are helpful for increasing the specific surface area and the active sites of loaded catalysts, resulting in enhanced the catalytic performances. When both novel carbon materials are doped with heteroatoms such as nitrogen and sulfur, they can act as the metal-free catalysts with significant electrochemical activity toward CO₂ reduction. Consequently, Both CNTs and graphene are promising materials for boosting the ERC0₂. To make large utilization of carbon materials in ERC0₂, further work is needed for some key problems such as the effects of heteroatom doping and structural defect.

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